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Critical Micelle Concentrations of Aqueous Surfactant Systems

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FOREWORD

The National Standard Reference Data System provides effective access to the quantitative data of physical science, critically evaluated and compiled for convenience, and readily accessible through a variety of distribution channels. The System was established in 1963 by action of the President's Office of Science and Technology and the Federal Council for Science and Technology, with responsibility to administer it assigned to the National Bureau of Standards.

The System now comprises a complex of data centers and other activities, carried on in academic institutions and other laboratories both in and out of government. The independent operational status of existing critical data projects is maintained and encouraged. Data centers that are components of the NSRDS produce compilations of critically evaluated data, critical reviews of the state of quantitative knowledge in specialized areas, and computations of useful functions derived from standard reference data. In addition, the centers and projects establish criteria for evaluation and compilation of data and make recommendations on needed improvements in experimental techniques. They are normally closely associated with active research in the relevant field.

The technical scope of the NSRDS is indicated by the principal categories of data compilation projects now active or being planned: nuclear properties, atomic and molecular properties, solid state properties, thermodynamic and transport properties, chemical kinetics, and colloid and surface properties.

The NSRDS receives advice and planning assistance from the National Research Council of the National Academy of Sciences-National Academy of Engineering. An overall Review Committee considers the program as a whole and makes recommendations on policy, long-term planning, and international collaboration. Advisory Panels, each concerned with a single technical area, meet regularly to examine major portions of the program, assign relative priorities, and identify specific key problems in need of further attention. For selected specific topics, the Advisory Panels sponsor subpanels which make detailed studies of users' needs, the present state of knowledge, and existing data resources as a basis for recommending one or more data compilation activities. This assembly of advisory services contributes greatly to the guidance of NSRDS activities.

The NSRDS-NBS series of publications is intended primarily to include evaluated reference data and critical reviews of long-term interest to the scientific and technical community.

LEWIS M. BRANSCOMB, *Director.*

Critical Micelle Concentrations of Aqueous Surfactant Systems

Pasupati Mukerjee* and Karol J. Mysels**

Critical micelle concentrations (CMC's), have been collected, organized and evaluated. The literature has been scanned for numerical values from 1926 up to and including 1966. In addition, over 800 values, hitherto available only in graphical form or implied in experimental data, have been extracted from the publications and are included. Close to 5,000 entries, based on 333 references, dealing with 720 compounds are tabulated in the main tables. Whenever available, the temperature, any additives present, the method of determination and the literature source are given for each CMC value and an indication of the apparent quality of the preparation and method used are included. A shorter table gives selected values which are believed to be particularly reliable, including highly accurate ones. Among these, concordant values from at least two independent laboratories are emphasized.

Included in the Introduction is a general discussion of the importance and significance of CMC values and of methods for their determination, as well as a summary of the procedures used in the collection, evaluation and presentation of these values in the present work. Extensive indexes are provided.

Key words: Association colloid; bibliography; CMC; colloid; colloidal electrolyte; critical concentration; critical micelle concentration; detergent; hydrophobic bonding; Krafft point; long chain compounds; micelle; paraffin chain salts; selected values; soap; solubilization; standard values; surface active agents; surface chemistry; surface tension; surfactant.

1. Introduction

Critical micelle concentrations are here to stay! This conclusion is evident from figures 1 to 3 which are based on the literature used in this work. They show a continuing growth since the middle thirties in the number of articles appearing each year which contribute new values and in the number of new values reported. The number of new values per article seems to have passed its peak, which suggests more careful and critical work in recent years.

The reason for this growth is that a critical micelle concentration (CMC) is probably the simplest means of characterizing the colloid and surface behavior of a surfactant solute, which in turn determines its industrial usefulness and biological activity, and gives a measure of the structurally interesting solute-solvent and solute-solute interactions. However, these published CMC values are widely scattered through the literature—we have consulted 87 different publications—and vary greatly in quality from clearly erroneous data to highly accurate values.

Furthermore, some of the existing values are clearly tabulated, but others—often the best ones—are hidden in graphs, or even in tabulations of some

measured property such as conductivity. These require considerable effort and judgment to retrieve. Frequently, the quality of the work cannot be judged without consultation of several references and intercomparison with other pertinent publications. Hence, much of the literature is not now readily accessible or useful to those interested in learning what has been established thus far.

The primary purpose of this publication is to provide a list of values in which the user can place high confidence. In the process of obtaining these, we had to make a survey, as complete as possible, of all available values. To present the results of this survey so as to make both the literature and the results contained therein readily available became, therefore, a secondary objective. Perhaps the best evidence for the usefulness of this effort is that nearly two-thirds of the best data reported herein were not previously directly available in the literature but required at least some, and often quite extensive, interpretation of a publication or individual correspondence.

The book itself is divided into four parts:

(1) The Table of Recommended and Selected Values lists the values we believe to be most reliable. They contain further guides to the quality of the data,

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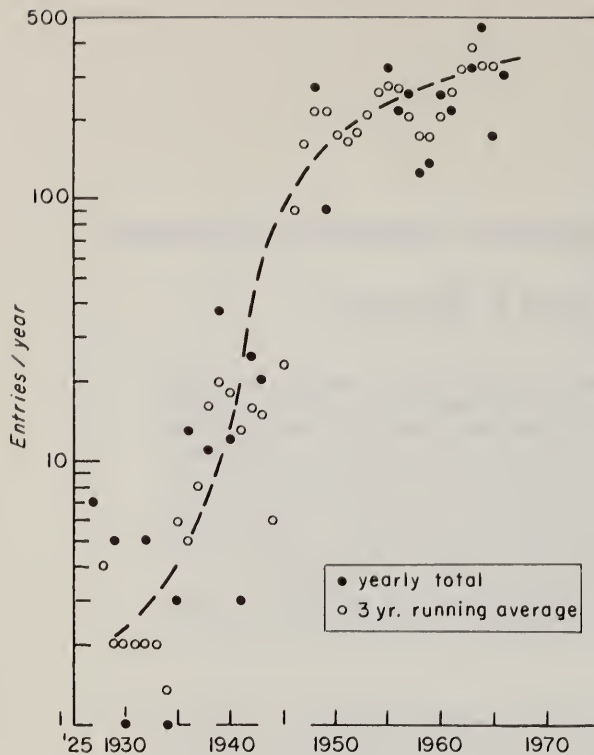


FIGURE 1. Number of entries for the complete tables originating within each year from 1927 to 1966.

There are no entries in 1928, 1931, 1933, 1934, 1937 and 1945.

(2) The Complete Table contains all values found which were published through 1966.

(3) Several indexes and lists, particularly the compound indexes, should permit the reader to find any desired compound or its closest analogs, give him the meaning of any abbreviation or symbol, and also guide him to the pertinent literature.

(4) The Introduction discusses the thoughts that went into the collection, evaluation, and presentation of the data. A glance at "How to Use These Tables" may be helpful before consulting them.

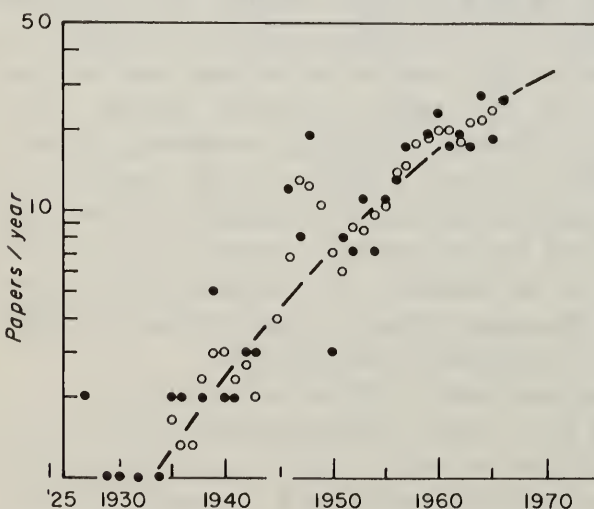


FIGURE 2. Number of papers containing at least one entry for the complete tables originating within a given year.

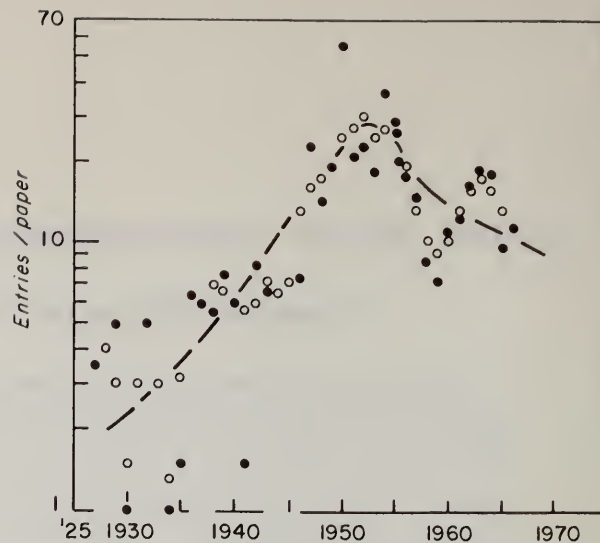


FIGURE 3. Average number of entries per paper containing at least one entry within a given year and within a three year period.

The largest number of entries from a single paper was 167, from reference 55004.

We hope that this work will make us more friends than enemies. We tried to be objective in the evaluation of the data, but some errors and personal prejudices are unavoidable. Our evaluation procedure of the individual data is described in the introduction, which contains also general considerations about the validity and significance of the various methods of determination.

We are grateful to many who have helped us in this work. Close to a hundred authors have responded to our request for reprints and many have provided additional comments, data, and interpretations. Dr. Edward L. Brady was most helpful in getting us started properly in the task of handling this multitude of data. Dr. H. J. White's patience and cooperation are greatly appreciated. The extensive computer handling of the data was made possible by the free availability of the data processing facilities of the R. J. Reynolds Tobacco Co., and the programming skill and understanding of Mr. Bill Donovan. Several secretaries have been involved in the careful verification of the hundreds of thousands of bits of information recorded. Mrs. Jerry Wilson and Miss Judy Tate were particularly involved in the final stages. The work leading to this publication was begun in 1964 at the University of Southern California under contract with the National Bureau of Standards and continued there until September 1966 when the authors transferred to their present connections with the University of Wisconsin, where the support of the National Bureau of Standards continued, and the R. J.

Reynolds Tobacco Co. Clearly, the resources of these three institutions made the completion of our work possible.

2. How to Use the Tables

This section presents a brief guide to the admittedly complicated arrangement of the tables. Space and computer requirements dictated much of this complexity; some is inherent in the dissimilarity of the compounds and the variety of conditions used for CMC determinations. The guide is arranged to answer a series of questions which may be raised by readers.

How do I find the compound I am interested in? In the tables the compounds are arranged in numerical order by arbitrary "Compound Numbers." To find this number you must go through the "Compound Index" in which the listed compounds are arranged by structure. There are five parts to this index (plus an alphabetical one for commercial names) and in each the compounds are listed according to different structural properties. These arrangements are described on the first page of the compound index (p. 23).

What can I do if my compound is not listed? The indexes will lead you to the most closely related compounds that are listed. These should permit you to make a good guess by interpolation and extrapolation.

How do I learn about the effect of an additive? The names of additives are abbreviated (if the abbreviation is not clear, its meaning can be found in the list on p. 222). Surfactant additives are indicated by their "Compound Number." For each compound, CMC values in the presence of additives are listed after the simple (surfactant-water) system in alphabetical order of the abbreviations. This is followed by systems with two additives and then by those with three additives.

What if the additive is not listed with my compound? The additive index shows all the compounds reported for any additive. You may find some useful analogies in this way.

What is the effect of temperature? Within each system (compound-additive(s)) the values are arranged by increasing temperature. By checking the author, or better the reference column, you can locate groups of values that were obtained specifically to show the effect of temperature (which is often small).

Which are the "good" CMC values? The shorter tables beginning on page 51 contain the "Selected" and "Recommended" values (15% of all reported values). Those that carry a "1" in the last column have been independently confirmed and should be highly reliable (to 1.5%, keeping in mind that different methods can give significantly different values—cf. p. 11). Those marked "2" are of the same apparent quality but lack confirmation. Among those marked "D" for each system, there is probably one that is as good as those of the preceding categories, but we do not know which. The many marked "3" do not seem to be in the same class but should be good to 10 percent.

What do I find in the long tables? These tables beginning on page 66 contain all the "Recommended and Selected" values plus all the others that we have located. In a number of cases, indicated by "R" in the last column, we make references to the literature where additional data or calculated values may be found or to warn the reader that the values are duplicates of those already listed or are in error. The bulk of the values carry an "L" in the last column. These may be useful and some may be excellent but we could not "recommend" or "select" them for a variety of reasons. Some clue to these reasons may be found in the "quality" column.

What is the "quality" column? In this column the first letter refers to the material and the second to the measurement. The meaning of the letters may be found on page 6. In general the quality decreases in alphabetical order. It represents our opinion after a careful study of the reference.

Are there more data in the literature? Our search does not cover anything published in 1967 or later (including the 1964 Congress of Surface Activity which did not appear in print until 1968). There are also older references that we may have overlooked. All the references within this field that we have scanned are listed in the literature index starting on page 213 whether they have yielded any entries or not. We would like to be informed of overlooked articles. Some of the literature scanned does contain data which, if properly interpreted, could lead to a CMC value which is not included. We have made such interpretations in many hundreds of cases, but not always. However, if a CMC value was mentioned as such in the article, we have tried to include it in all cases.

What are those various "methods"? The "method" column contains generally an abbreviation of the

method by which the CMC value was obtained. These methods are discussed briefly (and their abbreviations given) on pages 8 to 11. The "methods index" lists the references which have used each. These references should be consulted for details.

Occasionally the methods column contains information about the literature or a cross-reference. This is only the case when the entry does not give a CMC value.

In what units are the CMC's? We have followed the references except for order of magnitude conversions (e.g., from millimoles to moles) and as a result have a large number of units. The meaning of the abbreviations is given on page 222 and in the footnote to the table. For noncommercial compounds, for which a molecular weight is likely to have meaning, we have added a value in moles (per liter or kg of solution or kg of solvent) when the corresponding weight concentration of the compound was given. This was done by the computer on the basis of the molecular weight listed for the compound which in turn was also obtained by the computer from a structural or empirical formula of the compound. The value is printed by itself on a separate line below the value given by the author and is characterized by "M" in the "source" column.

What units are used for additives? The same units and symbols as for CMC's plus a number of others, including such peculiar ones as pH, again following the authors. In addition, we have used the additive columns to record certain special conditions such as pressure. The meaning of the abbreviations is listed on page 222. For additives we have not made any conversions to mole units.

What compound nomenclature is used? We have generally followed the first author whom we encountered dealing with the particular compound in the hope that this will also be the most common and understandable name. In case of ambiguity or some exotic names, we have added an alternative name or a formula in parentheses.

Are there any values for solvents other than water? If the solvent is a mixed one including water, the other components have been considered as additives. Nonaqueous systems have not been included for reasons discussed on page 18 with the exception of D₂O which is treated as an additive at 100 mole percent concentration!

What is the meaning of "source"? This column serves to indicate in what way the pertinent CMC value was obtained by us. The meaning of the abbreviations is listed on page 222. In some cases the

reader can check our listing directly or after carefully reading a graph or replotting some numerical data. In a few cases, however, our listing is based not only on what appears in the article but on correspondence or conversation with the authors. In this case an L in the source column is given. We have not included, however, values made available to us privately which did not have a basis in the published literature.

Where do these CMC's come from? The exact reference may be found in the Reference index starting on page 213 through the number in the "reference" column of each entry. However, much information can be obtained from this number itself since the first two digits give the year of publication and from the "authors" column which carries the first four letters of the name of the author or two of the authors of that publication. Particularly for those familiar with the field, this should often permit identification of the reference.

Are the numbers of digits really significant? Not in the great majority of entries. We have again followed the authors for the sake of the record and it is clear that most authors paid no attention whatsoever to the rules pertaining to significant figures. A better idea of the precision of the values is given by our "quality" rating of the method (second letter). See page 6 for the approximate meaning of these letters. When the value quoted is obtained by ourselves from published graphs, etc., the significant figures refer to how well these graphs could be read or interpreted without digging further into the uncertainties of the experiment.

3. Usefulness of CMC Value

The expression critical micelle concentration (CMC), as will be discussed later, is slightly misleading because of the use of the singular form of the noun "concentration." The formation of micelles from the constituent monomers involves a rapid, dynamic, association-dissociation equilibrium. Experimentally, it is found, in accord with the expectations from such equilibria, that micelles are undetectable in dilute solutions of the monomers, and become detectable over a narrow range of concentrations as the total concentration of solute is increased, above which nearly all additional solute material forms micelles. The concentration at which the micelles become first detectable depends on the sensitivity of the experimental probe used. The concentration range over which

the fraction of additional solute which forms micelles changes from nearly zero to nearly unity depends on such factors as the number of monomers in the micelle, the chain length of the monomer, the properties of counterions and other details affecting the monomer-micelle equilibrium. An approximate rule is that the higher the CMC value, the broader is the concentration range over which this transition takes place, in absolute value as well as in relative value in comparison to the CMC. Since different experimental methods may reflect this transition to different extents, some systematic variations in operationally defined CMC's are expected, as discussed in more detail later (p. 11).

Nevertheless, in spite of these various sources of uncertainty in defining and pinpointing the CMC exactly, the range of uncertainty is often no more than ± 1 to 2 percent of the CMC value. Thus, the CMC is a quantity which can be, and often is, determined experimentally to a much higher precision and accuracy than nearly any other property which is characteristic of solutions of surface-active agents, a point we would like to emphasize strongly. For comparative purposes, in careful work, the precision is often within ± 1 percent.

The usefulness of CMC values in various qualitative and quantitative investigations involving surfactant solutions arises basically from the fact that the surface and interfacial activity of the amphipathic (polar-nonpolar) monomers is closely reflected in the value of the CMC. The tendency to form micelles arises mainly from the presence of a hydrophobic part in the amphipathic monomers. The role of the hydrophilic part, nonionic, zwitter-ionic, or ionic (with associated counterions), which is essential for conferring enough of a solubility to the hydrocarbon chain so that CMC values can be reached or exceeded, is essentially a negative one as far as the stability of micelles is concerned. The same factors are involved qualitatively in the surface activity of the monomers, irrespective of whether the surface is an air-water interface, oil-water interface, or a nonpolar solid-water interface. There is thus an excellent correspondence between the adsorbability of the monomers, their ability to reduce surface and interfacial tensions, and the value of the CMC [1, 2].¹ The more surface active the monomer is, the higher is the tendency to form micelles and the lower the CMC value. Since above the CMC the monomer activity rises only very

slowly, the CMC is also a measure of the concentration at which the thermodynamic activity of the monomer and, therefore, its net surface activity and adsorbability to various substrates, level off to nearly constant values [1-3]. In closely comparable systems, particularly if the hydrophilic moiety of the monomer is kept the same and the hydrophobic part is varied, there is a considerable similarity in the amount of adsorption to air-water and oil-water interfaces at concentrations close to the CMC. It is thus often possible to obtain rough estimates of equilibrium monolayer concentrations from the CMC values in homologous systems [1-3].

Since adsorption from surfactant solutions is involved in widely ranging systems of technical importance such as foams, froths, emulsions, suspensions, and surface coatings, CMC values are important in a wide variety of industrial operations [4, 5].

In striking contrast to monomers, the micelles, which have a hydrophilic exterior, are not surface-active. As a result, above the CMC, excepting in some cases where small micelles form and the monomer activity increases appreciably, the surface and interfacial tensions decrease very little [1, 2, 6]. The CMC, therefore, indicates the concentration at which surface and interfacial tensions reach, approximately, their lowest values. Characteristic values at room temperature are often about 35 dyn/cm for surface tensions and 5 dyn/cm for interfacial tensions.

The CMC, of course, is the concentration at which the micelles make their first appearance. Micelles provide in many ways one of the most convenient systems available to study in depth the properties of colloids. As the properties of micelles depend on micelle-medium interactions and also micelle-micelle interactions, to understand the former without the latter complication, it is necessary to extrapolate properties of micelles to a point where micelle-micelle interactions become negligible. The corresponding extrapolation of preformed colloidal systems, such as polymers or proteins, which do not dissociate on extensive dilution, is made to "infinite dilution." For micellar properties, the CMC serves as a convenient point for extrapolation, i.e., "infinite dilution" for micelles. Just as binary protein-protein interactions (i.e., those involved in second virial coefficients) are experimentally determined from the slopes of curves as they approach infinite dilution, so in micellar

¹ Italicized figures in brackets indicate the literature references on page 20.

systems, the corresponding concentration range is the one just above the CMC [7-10].

In systems involving solubilization of an additional component or its distribution between the bulk solution and the micelle, the CMC again is a measure of the concentration at which such phenomena become first apparent. It will be discussed later that the addition of the third component may modify the CMC itself to some extent and, therefore, the CMC of the system in presence of the third component is the value to be used. The change in the CMC, however, is often small.

In situations where a quantitative estimate of the amount or concentration of micelles is desired, for example, in estimating solubilizing powers, or the effect of micelle concentrations on the chemical reactivities of constituent monomers or solubilized species, an area of research which is of considerable current interest [11-15], the CMC again serves the purpose of giving a rough estimate of the monomer concentration in the solution. The micelle concentration in equivalents, therefore, can be closely approximated as the total concentration minus the CMC.

For the quantitative study of the thermodynamics of the interactions involved in the monomer-micelle equilibrium, the CMC is of paramount importance [16-21]. Although considerable uncertainties still exist with regard to the proper means of estimating the charge effects in ionic micelles, for uncharged systems the CMC itself gives an approximate quantitative measure of the standard free energy of formation of micelles. These free energies and other derived thermodynamic quantities are of great potential and actual use in understanding hydrophobic interactions in general [22-24]. Such interactions are involved in a wide variety of biochemical phenomena, e.g., the stability, structure, conformation, and activity of proteins, enzymes, and membranes. With ionic micelles, as mentioned before, the calculation of thermodynamic quantities characterizing the various interactions is not on sure ground as yet. For comparison of related systems, however, e.g., in noting the effect of varying the chainlength, salt concentrations, or counterions, the CMC provides quite a good quantitative measure of the changes as they affect the monomer-micelle equilibrium [8, 18, 25].

4. Evaluation

An important part of this work is the evaluation of the data presented. We hope to guide the reader to those data that are most useful and reliable in our considered judgment and also to show him other values that exist in the literature so that he may make more easily his own evaluation. We also tried to indicate the relevant literature and data that we have considered but not used in final listings.

We divided our evaluation into two steps: one, which we may call the individual or preliminary evaluation; the other, the comparative evaluation. The former represents our opinion on the basis of the individual paper (and its references or related papers of the same author); the latter is based on intercomparison of all the available data for a given compound under the same, or closely related, conditions. The former was done as the work progressed over a period of three years, 1965-68, and, therefore, is subject to the drift and development of our ideas and skills during that time. The latter was done in a short period of time after all the data had been collected and sorted by the computer.

Individual evaluation. A preliminary separation involved the question whether a given CMC value should be reported in detail or not. Values which are indicated in the article as being duplicates of other published values are omitted completely. Others, however, which are clearly duplicates but not explicitly indicated as such by the authors, are mentioned as "VALUES FROM REF IN CMC," with the article from which they are taken listed in the column in which the CMC is normally found.

Values which could not be retrieved profitably, e.g., those in the form of small-scale graphs or summarizing equations, are indicated as "GRAPH DATA NOT RETRIEVED" and "SUMMARIZING EQN ONLY" for the reader who wishes to examine them himself. There are 41 entries in the former category and 22 in the latter.

Once a value was included explicitly, we attempted to evaluate the purity of the materials and the accuracy and precision of the method used. These were noted separately as reported in the "Quality" Rating columns. The meaning of the symbols is as follows:

MATERIAL

A Highest purity—not likely to be significantly improved in the future

METHOD

precise to about 1% accurate to 1.5%

B	Very pure but may still contain significant traces of impurities	precise to about 2% accurate to 3%
C	Purified but likely to contain significant impurities	precise to about 3% accurate to 5%
D	Compounds purified without special precautions against surfactant impurities	precise to about 10% accurate to 10%
E	Good quality mixture of head groups	order of magnitude
F	Good quality mixture of homologues	wrong
G	Not purified	likely presence of systematic errors of uncertain absolute and relative magnitude
H	Commercial surfactant material	
P	High purity claimed without supporting data	
Q		theoretically calculated
X	No information given	

These ratings should be taken with certain reservations, both because of the possibility of personal bias and of the frequent lack of all the information required to form a definite opinion, and the unavoidable lack of consistency over the four-year span during which they were being assigned. All that can be said is that they represent our best opinion at the time of a detailed reading of the article cited. Because the evaluation was individual, it is not surprising that upon later comparative evaluation, some values which were initially ranked high appeared more questionable or vice versa. Hence, the ratings are at best meaningful to plus or minus one category. We may mention, however, that during the final evaluation we frequently referred to the original papers, and only in two cases did we feel the need to change the preliminary ratings.

Comparative Evaluation. This evaluation was performed after all the data were collected and classified by the computer into the form of the "Complete CMC Listing" of this report. This permitted easy intercomparison of values reported for the same or similar systems. Each value was then assigned to one of a set of categories. This assignment also formed the basis for obtaining the "con-

firmed," "recommended," "disputed," and "selected" values for a separate listing.

The following categories were used:

1-Confirmed.—Values from at least two independent laboratories which are of good quality both with respect to materials and methods and which agree within the expected limit for a given method or between methods. These may be used with high confidence.

2-Recommended.—Values of the same quality as above but lacking independent confirmation.

D-Disputed.—These are values which seem as accurate as the preceding ones, but disagree by more than would permit them to be classified as "confirmed." It is likely that one of the values is correct but we did not have a firm basis for deciding which. In some cases correlation with values for other systems indicated a definite choice and the preferred value was assigned a "2," i.e., "recommended" rating, and remaining values were assigned the "L" or "literature" rating.

3-Selected.—These are values which do not appear to have the accuracy of the above categories but should still be of considerable utility. We feel that they are probably within 10 percent of the "true" value, as measured by the same approach. Their total number is less than 10 percent of all the entries, and they represent the next most reliable group after (1) and (2).

L—Literature.—This category includes the great majority of the values. They are provided for bibliographical completeness and because some readers may prefer any value—no matter how unreliable—to no value at all. Some of these values may be accurate, but there is insufficient information to certify that this is the case. Some will turn out to be off by orders of magnitude. In some cases where the literature provided *prima facie* evidence that the CMC value was erroneously assigned, we have indicated this by the statement "QUESTIONABLE CRITERION" in the "method" column.

P—Preferred.—Occasionally, widely differing values have been reported for what should be a single CMC. The reader's choice can generally be guided by the quality rank assignment to the compound and method. In a few cases we thought it advisable to indicate the preferred value by a P. These data did not qualify for one of the "selected" or "recommended" categories, but seem to be clearly preferable to the others.

R—Indicates a literature reference in which the reader may find a value not tabulated by us. A

statement in the "methods" column indicates whether the reason for omission is that we considered it a duplication (without explicit reference) of a value from another publication, or a value which could not or should not be listed.

X—Indicates a cross-reference within our tabulation and is used for mixed systems of surfactants which are reported only once under one compound but cross-referenced under the others.

In deciding about the probable accuracy of a CMC value we considered not only the details of the particular measurement but also the general validity and limitation of the method used. It may, therefore, be appropriate now to review briefly the multitude of these methods and then to consider some factors which guided us in judging their intrinsic accuracies.

5. Methods of Determining CMC's

Although abrupt changes in the concentration dependence of several properties of several surfactant solutions had been observed before the 1930's [26-28] and the concept of micelles in such solutions had been developed by McBain and co-workers [29, 30], the existence of a narrow concentration range, called "the critical concentration for micelles" [31], below which the solution contains negligible amounts of micelles and above which practically all additional surfactant is found in the form of additional micelles, was established by Bury and his coworkers [31, 32] and Hartley and his coworkers [33] in the early 1930's. Since that time, CMC determinations have multiplied and the results have been used in a variety of ways. The importance of a definite CMC value to which micellar properties could be extrapolated so as to give the infinite dilution behavior of micelles was emphasized by Debye soon after World War II [34].

In the process of collecting the present data we have distinguished 71 methods of determining the CMC, ranging from a few widely used ones to a few reported only once. These may be grouped as follows, with the numbers in parentheses giving the number of CMC's in this report for each method or group of methods. Experimental details of these methods may be found by following the references given for each in the index to methods.

The entry "METHOD NOT CITED" (128) in most cases refers to just that, when a CMC value appears with no further indication. In a few cases,

however, it results from the fact that the article is not clear as to which of two or three well-defined methods was used in determining each individual value. One or two of such cases deserved, in our opinion, enough confidence to be included among "selected" values.

Also in a special category is the THEORETICALLY ESTIMATED (5) entry which we included for completeness.

The other methods can be divided into two broad classes depending on whether another material is added to the system specifically for the purpose of the measurement (as opposed to an additive whose effect is being studied). They are at present all based on the study of a property of the system as some function of concentration and detection of a particularity, such as a change of slope or a discontinuity, at the CMC.

Methods Requiring No Additive

1. SURFACE TENSION (940). This method of increasing popularity involves the measurement of surface tensions of solutions by a variety of methods which we did not attempt to classify (such as the du Nouy ring detachment method, the Wilhelmy plate equilibrium or detachment, drop weight or volume or shape). The data are generally plotted against the logarithm of concentration—LOG PLOT (843)—as the abscissa and the transition between a descending line (often assumed to be straight) and another one close to the horizontal is taken as the CMC. The data can also be plotted directly against the concentration—LINEAR PLOT (18)—in which case the curvature of the descending portion is much more marked and the transition less sharp. A MINIMUM (15) in the curve is now known to be due to the presence of a third component (contamination or products of hydrolysis) which is more surface active but removed from the surface by solubilization in the micelles above the CMC. It is sometimes reported as the CMC of the system. We have also encountered the UNSPEC (64) where no details are given. Interfacial tension methods are considered later among methods involving an additive.

A related method is based on the FOAMING POWER (2) of the solution. It is not clear at present how the changes in this property are related to the association of monomers into micelles.

2. Electric conductivity—COND or COND-CTNCE (953)—is based on the measurement of the A-C electric resistances of the solutions. These can then be interpreted in terms of the specific

conductivity—SPECFC (386)—and plotted—GRAPH (343)—against concentration to give two almost straight lines whose intersection is the CMC or the data corresponding to each straight line can be summarized by equations—EQUATNS (43)—and the CMC obtained analytically. The data can also be converted to equivalent conductivity—EQUIV (352)—and the CMC obtained graphically—GRAPH (319)—usually plotting against the square root of concentration. Occasionally linear or cube root plots have been encountered. In much early work it was the first detected deviation—1ST DEVIATION (18)—that was reported as the CMC. As the CMC thus assigned depended clearly on the sensitivity of the method, this approach seems to have been abandoned. A few surfactants having low CMC's show a maximum of equivalent conductance in the CMC region. The origin of this maximum is not yet clear [34a] and in some cases the beginning of the rise—MAX BEGINING (10)—in others the MAXIMUM (5)—itself is reported as the CMC.

Some authors report averages—AVER (110)—of two of the above methods, namely specific and equivalent—SP EQUIV (104)—conductivities or of the beginning of the rise to the maximum and of specific conductance—COND BEGINNING MAXIM (6). Some report results based on conductance without further details—UNSPEC (105).

The variation of electric conductance with frequency at high frequencies, the so-called WIEN EFFECT (1), is different for micelles and monomers and has been used to determine the CMC.

3. There is a variety of methods to investigate optical and spectroscopic properties of a solution, and they may be classified as follows:

(a) Measurements of scattered light—LITE SCATR or LIGHT SCATTER (317) depends on the measurement of the intensity of light at an angle, generally 90°, to the incident beam.

A plot of this intensity or of the turbidity of the solution (which is proportional to it) shows a low slope for dilute solutions and a steeper one above the CMC. The intersection of the two parts on a TURBIDITY PLT (291) gives the CMC. Debye has pointed out that the concentration above the CMC divided by the turbidity (or excess turbidity above that at the CMC) is close to a straight line. Conversely the CMC may be obtained by selecting the value which gives the best straight line on such a DEBYE PLT (25). In light scattering the exact

procedure was not indicated only once—UNSPEC (1).

(b) The CMC can be obtained from the change of slope of the REFRACTIVE INDEX (134) when plotted against concentration.

(c) The absorption spectrum of some surfactants is different when they are in micellar and in free form. Hence, plotting the absorbancy at a suitable wavelength can give a change of slope corresponding to the CMC. This MICELLAR SPECTRAL CHANGE (30) method should be distinguished sharply from the multitude of other spectral change methods in that it does not require any additive.

(d) Others. A CMC has been obtained by the change in the X-RAY DIFFRACTION (1) pattern but this method has a very low sensitivity and, therefore, precision. Another method using a spectroscopic technique which has been introduced after the closing of this survey involves shifts of nmr peaks.

4. Calorimetric methods used to obtain CMC's are those of SPECIFIC HEAT (4) and HEAT OF DILUTION (24).

5. Two colligative properties have been used. FREEZING POINT (14) lowering which gives the CMC at a single temperature, generally close to zero, determined by the nature of the solute and VAPOR PRESSURE LOWERING (38) generally measured with a so-called vapor pressure osmometer based on the temperature comparison of two droplets, one of solution, the other of solvent.

6. The abrupt increase of solubility with increasing temperature which occurs once the solubility reaches the CMC region is the basis of the KRAFT POINT SOLUBILITY (21) method. This change is sharpest when the logarithm of the solubility is plotted against the inverse absolute temperature. Often other plots or even single point experiments are used which have little or no value.

7. Several of what may be called transport properties have been used. They include:

(a) Measurements of the DIFFUSION COEFFICIENT (2) of the surfactant which gives, of course, an average of the very different mobilities of micelles and monomers and leads to a change of slope in a D versus C plot to give the CMC.

(b) The VISCOSITY (9) of the solution plotted as the specific viscosity $[(\eta - \eta_0)/\eta_0]$ or as the reduced viscosity (η_{sp}/c) shows a change of slope at the CMC. A related method depends on an observed VISCOSITY MINIMUM (14) in the CMC region. This minimum probably results from a combination

of surface tension, contact angle, and viscosity effects in the capillary instrument used, so that its relation to the micelle formation is not clear.

(c) Measurements of STREAMING CURRENT (6) have been also used for determining the CMC and should perhaps be classified among methods requiring an additive since these electrokinetic phenomena depend on the adsorption on the solid involved which is specific to the surface. They also depend, though only in a secondary way, on changes occurring in the bulk of the solution which involves the CMC.

(d) Changes in the concentration of the filtrate in ULTRAFILTRATION (9) and in the sedimentation coefficient in ULTRACENTRIFUGATION (1) also lead to CMC values.

8. Potentiometric measurements use several approaches:

(a) ELECTROMOTIVE FORCE (32) methods involve measurements of potential of either specific ion electrodes other than pH ones or of an insoluble mercury-surfactant salt electrode against a salt bridge-reference electrode. It may be noted that the presence of the salt bridge often causes local precipitation of the surfactant and that insoluble salts can be solubilized by micelles. These complications are often overlooked but should, perhaps, cause these methods to be considered with those involving additives.

(b) EMF ALONG CONC GRADIENT (1) uses two closely spaced identical electrodes moved through a solution having a known concentration gradient. Hence, it depends on a change of the slope of the emf—versus—concentration curve.

(c) PH AND HYDROLYSIS (9) method involve pH measurements interpreted either directly or after conversion to a degree of hydrolysis. They are grouped together since they often involve problems of carbon dioxide contamination as well as salt bridges.

9. Other bulk properties that have been used to determine CMC's are density (23) based generally on magnetic float methods plotted directly as DENSITY (17) or after conversion to PARTIAL VOLUME (6) and ultrasonic VELOCITY OF SOUND (10).

Methods Involving the Use of an Additive in the Bulk of the Solution

1. Spectral change—SPCTR CHNGE (1602)—methods.

This family of methods whose limitation (which we

consider quite severe) will be discussed below (p. 12) is by far the most fertile one as far as supplying CMC values. It is based on the fact that the spectra of many dyes added in very small amounts to a surfactant solution are very different in the region below and in that above the CMC. Pinacyanole—PNCN (1176)—is also by far the most popular among these dyes. The CMC may be determined, for example, by titration of a solution above the CMC by one below the CMC, both containing the same concentration of dye. Some definite shade is chosen as the end point corresponding to the CMC. Alternatively, solutions having concentrations bracketing the CMC are prepared containing the same concentration of the dye and their color examined visually, or their spectrum or their absorbancy at a specific wavelength measured. In any case, the concentration at which some sharp change occurs is taken as the CMC. In some cases the values are extrapolated to zero dye concentration. For one dye Rhodamine 6G—RHD6 (74) in addition to the spectral change, the change in fluorescence FLUOR CHNGE (19) was used and is based on the same principles.

We have noted whether the technique used was VISUAL (1277) photometric—FOTOMTR (305)—or UNSPEC (20)—as this is related to the precision expected.

Other dyes which have been used in this method are Fluorescein—FL (2), Erythrosin—ERTS (7), dichlorofluorescein—DCLF (9), Benzopurpurine 4B—BZP4 (11), Bromphenol Blue—BRPB (21), Sky-blue FF—SKYB (43), Eosin—EOSN (51), Indophenol—INPX (129) and, included because it performs the same function although not a dye in the strict sense, Iodine—12 (52). There is also unspecified VISUAL SPECTR CHNGE (27).

2. The fact that many water-insoluble substances dissolve significantly in the presence of micelles, i.e., are solubilized, has been used in the determination of CMC's by solubilization—SOLUBLZTN (293)—methods. The solubility of dyes has always been determined photometrically—FOTOMTR (274)—unless it is UNSPEC (12), that of TOLUENE (4) has been determined volumetrically, and the limit of solubility of lauryl alcohol has been established turbidimetrically—TURBIDMTR LOH (3)—. The most popular dye is the so-called orange OT—OROT (158)—which is always 1-o-tolyl-azo-2-naphthol, CI Solvent Orange 2, m.p. 128–9°, and not the complicated pigment which was once available under that name which is CI Pigment Orange 13,

m.p. 332°. Other solubilizates used are paradimethylaminoazobenzene—PDMAB (59), Sudan 4—SDN4 (46), azobenzene—AZBA (11), Yellow OB—YLOB (9), 2-nitro-diphenylamine—2NPA (2) and dimethyl yellow—DMYL (1).

3. Methods based on liquid-liquid interface phenomena.

(a) Measurements of INTERFACIAL TNSN or TENSION (52) between an aqueous solution of a surfactant and an immiscible liquid have been used in the determination of the CMC. Most values were obtained from semilogarithmic plots, LOGM (46) but we also found UNSPEC (6). These methods are classified among those requiring an additive because of the always finite solubility of the other liquid in the aqueous phase especially above the CMC. A frequently present source of error in this method is the solution of some of the surfactant in the nonaqueous phase which can radically change its real concentration.

(b) We are placing in the same category methods based on the suppression of the POLAROGRAPHIC MAXIMUM (45) since this suppression is related to an increase in surface (dilational) viscosity of the mercury-solution interface which reduces the convection currents responsible for the maximum in the "diffusion" current. These surface viscosity changes are in turn related to the adsorption of the surfactant and thus depend on its activity. However, it is not clear at present how the final polarographic criterion is related to the CMC. These measurements require also the presence of a very high concentration of an inert electrolyte and that of an electrochemically active indicator ion. This makes the composition of the solutions rather unique and prevents comparison with other data. Although we are quite skeptical about the significance of data obtained by this method, we have reported them in view of lack of any definite evidence of their invalidity. A further disturbing fact is that sometimes two CMC's are reported for a given system. One of them is, therefore, certainly wrong. However, for the sake of completeness we had to record both.

4. Other methods in this class include a result obtained from changes in the FLOCCULATION RATE (1) of a suspension and those from changes in the (reaction) REACTN RATE OF A SOLUBILIZATE (2) as it becomes solubilized above the CMC.

6. Reasons for Methodical Differences Between CMC Determinations

As shown by the above review of methods used, the determination of a CMC involves a series of measurements of some property of solutions of surfactant, alone or with an indicator, as a function of concentration followed by the detection of some characteristic point which is called the CMC. Methodical differences may originate from the choice of the characteristic point, the kind of plot on which this point is chosen, the kind of data which are plotted, and the effect of the indicator, if any, upon these data. We use here the term "plot" in a general sense to include any serial representation and even titration, although in the great majority of cases it is a real graphical plot that is involved.

If the CMC were a sharply defined point, such as a melting point, above which some properties were qualitatively different from those below it, methodical differences would be nonexistent or greatly reduced. In fact, however, all properties of a solution in the CMC region vary in a continuous manner and so do all their derivatives. There is, nevertheless, a relatively narrow region of concentration in which these changes are most marked. This is illustrated by precise measurements on pure systems such as those of figure 4 (and is, of course, exaggerated on impure ones) but is perhaps best

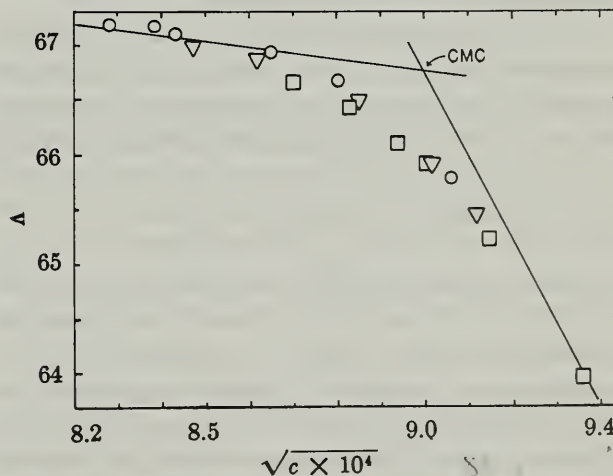


FIGURE 4. Detail of the CMC region for sodium dodecyl sulfate.

The straight lines are based on many points beyond the limits of the figure. The points were obtained by two investigators on two different preparations of the compound. From reference 58013, reprinted by permission of the Journal of Physical Chemistry.

grounded on broad theoretical considerations. One is that a micelle is by definition a reversible aggregate of a large but not infinite number of monomers and that the micelle-forming reaction must obey the laws of chemical equilibrium. As such, the

concentration dependence of the degree of micellization has to change gradually although it may change rapidly [32, 33, 35]. A truly abrupt, discontinuous, transition is excluded. Consequently, all properties of the solution must show similarly rapid but gradual changes. Another argument which is less self-evident but seems to be well supported is that there is not a unique number of monomers which can form a micelle but a range with relatively wide limits. Hence, micelles are polydisperse. This is in agreement with the fact that (the average) micellar size varies continuously with conditions such as temperature, concentration of surfactant, concentration of counterion or other additives, nature of counterion, chain length and structure, and also that micelles are good solvents both for other materials as shown by solubilization, and for each other as shown by the frequent continuous formation of mixed micelles. Reasonable assumptions about forces determining the size of the micelle also lead to the conclusion that they must be polydisperse [35a]. Hence, in the CMC region not one kind of micelle, but many kinds appear, each with a slightly different concentration dependence which further spreads and complicates the changes of bulk properties occurring in this region.

In light of these considerations, we shall now review in more detail the sources of methodical discrepancies in CMC determinations and their effect upon our critical evaluation.

1. *Choice of the characteristic point.* As already mentioned, in some early work, it was the first detectable deviation from monomeric properties that was taken as the CMC. This depended greatly on the sensitivity of the method and has generally been replaced by an extrapolation from below and from above to a point of intersection. If the two lines are straight and differ markedly in slope, this is a simple procedure. Unfortunately, such is seldom the case. Some of the procedures, therefore, involve a treatment of the data to obtain a straight line. Thus, equivalent conductivity of strong electrolytes is plotted against the square root of the concentration in accordance with the Onsager limiting slope.

It also seems true that those properties which give the best straight lines, because they are rather insensitive to interparticle interactions, also give small differences of slope for the same reason. Density and refractive index or specific conductance in the presence of high salt concentrations are good examples. Hence, they require precise meas-

urements to give useful results. This implies keeping constant all factors other than the concentration of surfactant, such as the temperature, and also the concentration of other components such as added salts. The latter often requires extreme precautions against evaporation. In the interpretation of such precise measurements, difference plots (i.e., experimental value minus a straight line value) are useful. A few methods, however, give good straight lines of very different slopes, e.g., conductance in the absence of high salt concentrations, solubilization of some dyes, or micellar spectral changes of some pyridinium compounds.

Closely related to the extrapolation method is that of the point of maximum rate of change of slope (i.e., zero value of third derivative) which is useful in theoretical calculations [16] but is so difficult to apply experimentally that it does not seem to have been used.

In the spectral change methods using indicator dyes there is generally a change of absorbancy at any given wavelength from one characteristic value below the CMC to another above the CMC. Some authors take the beginning, some the end, and most the midpoint of that change. The latter is by far the most objectively defined but, as we shall see later, is a function of concentration of the indicator. In visual methods there is much subjectivity and the result can depend markedly on local illumination because of the very complicated absorption spectrum of certain dyes, particularly the most frequently used one, pinacyanole.

Clearly, depending on which point is defined as the CMC, values covering a considerable range may be obtained. We have given weight only to those methods which defined a point in the middle part of the range, particularly those using an extrapolation procedure, and much of the following discussion will be restricted to those.

2. *The kind of plot chosen.* A point which is not often appreciated is that the same experimental data can give different values for the CMC by extrapolation procedures depending on how they are plotted. The best known example of this is the difference between CMC's obtained by plotting conductance versus concentration and the corresponding equivalent conductance versus the square root of concentration. Figure 5 shows an example of the same set of experimental data plotted in these two ways. Very reasonable linear extrapolations give a well-defined CMC in each case, but the two values

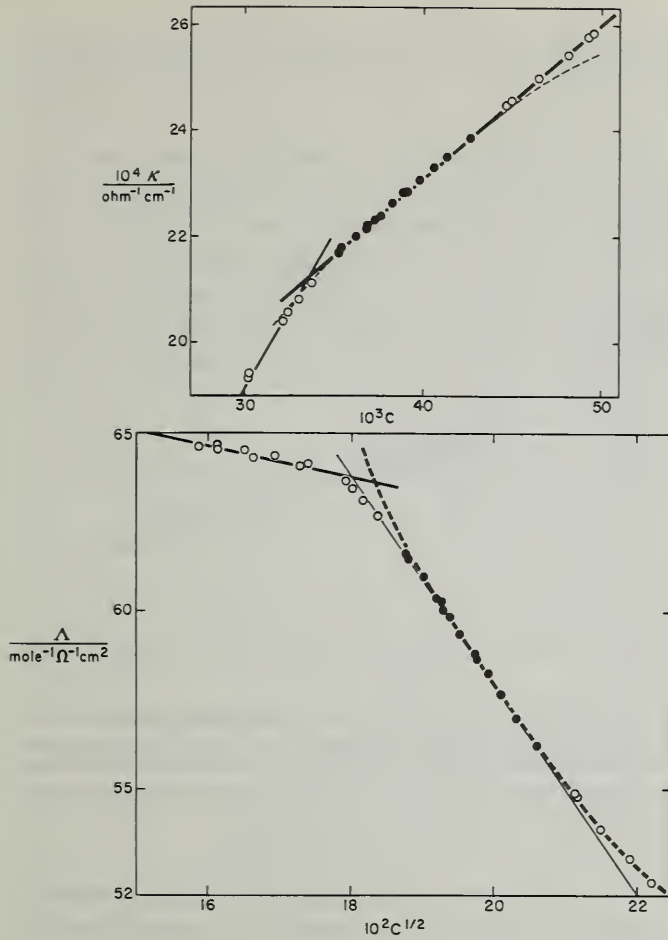


FIGURE 5. The effect of the way of plotting upon the value of the CMC obtained from the same data.

Above: conductivity versus concentration; below: equivalent conductivity versus square root of concentration. The dotted curves on each graph correspond to the straight line of the other graph. The filled points are common to both lines. Data for sodium decyl sulfate obtained by Dr. R. J. Otter. From the 14th Technical Report, Office of Naval Research, Contract Nonr-274(00) by K. J. Mysels, R. J. Otter and P. Kapauan, University of Southern California, October, 1960. (Cf. Ref. 61017).

differ by about 3 percent, which is much more than the uncertainty in each. The discrepancy is due in large part to the fact that, a straight line on one plot is a curve on the other so that different points are chosen as the basis for extrapolation and the intersection is shifted.

3. *The kind of data.* If micelles were monodisperse then, for CMC determinations, the molecular weight or charge dependence of a property would not matter as long as it was linear with some function of the concentration. However, if one accepts the premise that polydispersity is possible, it is clear that different physical properties will give differently weighted measures of micellar concentrations and, therefore, different extrapolated values of zero micelle concentration, i.e., different CMC's [18]. The most accessible experimental variable is the total concentration of surfactant conveniently ex-

pressed in equivalents. We will estimate the kind of average that is plotted against this variable in some popular methods without attempting to discuss this subject fully.

Solubilization. As a first approximation we may assume that the surfactant chains have a constant solvent power independent of their configuration, i.e., independent of the micellar size. The quantity measured—the concentration of solubilize—is then directly proportional to the equivalent concentration of micellized surfactant and independent of polydispersity. More formally, we call n_i the number of monomers per micelle of a given size, M_i the molar concentration of these micelles, and C_i their equivalent concentration. Each group of micelles then dissolves an amount of solubilize proportional to $M_i n_i$ and the measured total give us

$$\sum M_i n_i = \sum C_i = C_m$$

where C_m is the total equivalent concentration of micelles. Thus, the result is unweighted.

Colligative measurements. These give the sum of molar concentrations of micelles (for the simple nonionic case). Hence, we get

$$\sum M_i = \sum C_i / n_i$$

the inverse-size weighted equivalent concentration of micelles.

Turbidity. The total turbidity is the sum of the turbidities of the individual micelles and these in turn are proportional to the square of their mass (again neglecting charge effects and other interactions). Hence, we measure

$$\sum \tau_i = \sum M_i n_i^2 = \sum C_i n_i$$

or the size weighted equivalent concentration of micelles.

Thus, it is clear that these different methods must yield different results. Yet, when the polydispersity is small, the micelles large, and the transition region narrow, the differences may be small, often smaller than experimental uncertainties. Thus, depending on the precision involved and the particular system under consideration, the term CMC may have a definite meaning or may have to be tempered by a specification of the experimental method used and also of the way in which the results are interpreted. Unfortunately, there is little definitive information about the final effect that these factors have on the CMC values determined by various methods.

In the final evaluation of the CMC values, we had to face the question whether two numbers obtained under allegedly identical conditions and by apparently reliable measurements did agree sufficiently to confirm each other. In making the decision, we tried to take into account the differences expected between methods in addition to differences due to the unavoidable experimental uncertainty on which we placed an arbitrary limit of 1.5 percent. Hence, two CMC values differing by 4 percent may be taken as confirming each other, if the methods used would be expected to differ by 3 percent (in the proper direction), as is indeed the case for conductance and equivalent conductivity measurements.

4. The effect of the concentration range used. Extrapolation generally uses straight lines. Few physical properties vary, however, exactly linearly. Generally, some curvature is present in reality. Hence, the line drawn tends to be a chord and its direction and position are a function of the portion of the curve that is being approximated. This remains true whether a line is drawn by inspection or after a least square calculation. In terms of CMC determinations, it means that the value obtained depends generally on the range of concentrations above and below the CMC over which the extrapolation is taken and, therefore, the number and spacing of points. There is no general agreement as to what these ranges should be and herein lies another source of discrepancies. Figure 6 shows the effect of the range of data used upon the CMC.

In dilute solutions there are some theoretical guides to the expected curvature which suggest the kind of graph which is likely to make the experimental points fall close to a straight line. The square root for conductivity, the logarithm for surface tension are good concentration scales for monomers, and micellar concentration divided by turbidity is a good one for light scattering. In more concentrated solutions the systems depart more and more from ideality and all plots become nonlinear. Extrapolation then becomes more and more a question of judgment and of the range and spacing of available data, and the CMC value becomes highly subjective. Since compounds with shorter chain lengths tend to have higher CMC's, this is an important factor in reducing the accuracy of CMC determinations as the chain length decreases below 10 carbon atoms.

The CMC region is itself a region of curvature of any measured property. This means that experi-

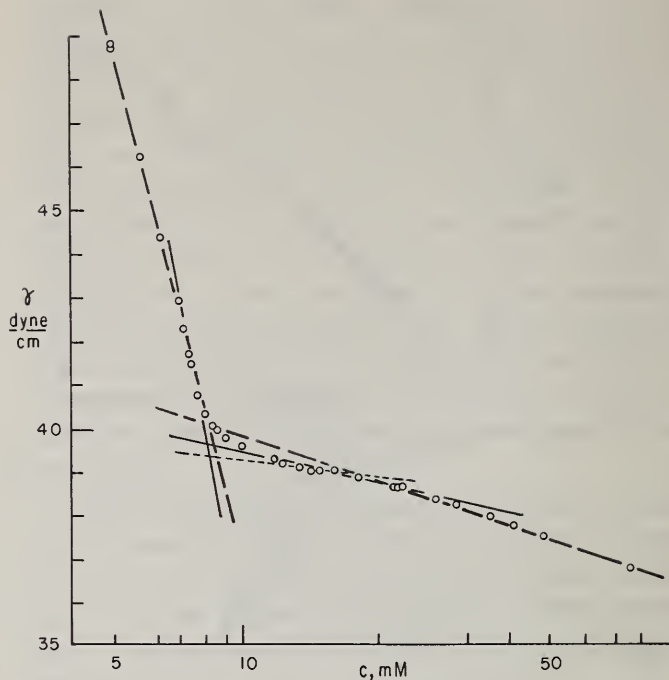


FIGURE 6. *The effect of the range of values used upon the CMC.*

Surface tension data for highly purified sodium dodecyl sulfate from reference 66007. The various straight lines represent extrapolation based on points within a certain range of concentrations. If points outside this range were not available, each of these extrapolations could be considered as quite reasonable.

mental points very close to the CMC cannot be used for its determination because they deviate from both lines used in the extrapolation. The sharper the curvature the smaller the unusable concentration span and the simpler the extrapolation. Theoretical calculations show clearly [33, 35] and it is easily seen physically, that the larger the micelle the more cooperative is micelle formation and the sharper the curvature. Polydispersity of micelles can only smear out the transition region. Hence, pure systems show much better defined and more easily measured CMC's because of reduced polydispersity. For space-filling reasons, long-chain monomers can form larger spherical micelles, and do form larger micelles in general according to available experimental evidence [8]. Short-chain compounds, therefore, present an additional difficulty because of the spread of their CMC region not only in absolute terms but also in relative ones.

The combined effects of curvature at higher concentrations and of the relative width of the transition region account for the fact that we have not been able to include any C-8 compounds among those for which confirmed or recommended values are available. This is despite the fact that these compounds are intrinsically easier to prepare in a state of high purity, that the measurement of the physical properties of their solutions can be made

more precisely and that any impurities in the solvent are less disturbing just because the CMC's are higher.

Long-chain compounds present different but no less serious difficulties because their CMC's are so low. Here the actual measurement of the physical property becomes difficult because it often differs little from that of the solvent, the control of concentration may become a problem because surfactants tend to adsorb at interfaces, and trace impurities in the solvent play a larger role. Furthermore, the compounds themselves are intrinsically less easily purified because certain operations, such as distillation of intermediates, become more difficult, and also because there is an increased number of isomers and because properties of homologues and isomers become more similar. In our collection only one value for a C-16 compound became recommended and an inordinate fraction is marked "D" for disputed, reflecting the difficulties involved.

Thus, the bulk of reliable CMC data deals with compounds in the C-10 to C-14 range. It is clear that better experimental methods are needed to extend the upper limit and better interpretation to overcome the lower one.

5. *The effect of indicators.* A very large fraction of all reported CMC values was obtained by methods involving the deliberate addition of an indicator, generally a dye which is solubilized and presents an easily measurable difference in the solubilized and not-solubilized state. This requires a discussion of the role of these indicators and of the reasons which led us to accept certain of these methods as valid and to reject the great majority as likely to be biased by significant methodical errors.

Solubilization is the marked increase of solubility associated with the formation of micelles. It is of special interest for substances which are sparingly soluble in water but is not limited to these. To a first approximation the excess solubility is directly proportional to the concentration of micelles [36] showing that these act as a solvent having a limited solvent power for the solubilizate. We can consider the saturated solution as an equilibrium system in which both the true solvent (water) and the micelles are saturated with the solubilizate. In undersaturated systems there are good reasons to believe [36, 37] that a distribution equilibrium between water and micelles exists.

It is a well-known rule of physical chemistry that the activity of the solvent is lower in a solution than

in the pure state. Hence, in the macroscopic world, the liquid phase forms at a lower vapor pressure in the presence of a solute than in its absence. In fact, at equilibrium the first drop always forms at the vapor pressure of the saturated solution. In the microscopic world of the micelle considered as a solvent, the analogy is clear: the activity of the surfactant in the micelle is lower in the presence of a solubilizate. Hence, micelles form at a lower concentration of monomers in the presence of the solubilizate. In fact, the first micelles exist at the equilibrium monomer concentration corresponding to a micelle saturated with the solubilizate under the conditions of the experiment. Two conclusions can, therefore, be drawn: the presence of a solubilized indicator always lowers the CMC, and this lowering reaches a maximum if the micelle is saturated with the indicator during the determination.

If the solubilizate is assumed to form an ideal solution in the micelle and related simplifying assumptions are made, it has been shown by Shinoda [38] that the CMC is changed by a factor $(1 - 1.36x)$ for an ionic surfactant where x is the mole fraction of additive in the micelle. For the nonionic case, the factor becomes simply $(1 - x)$. The value of x depends, of course, on the concentration of micelles at the point taken as the CMC, the total available amount of indicator, and its solubility in water and in the micelle.

An additional and often neglected complication occurs when the indicator is a high molecular weight organic ion such as most of the dyes used in the spectral change methods, in particular pinacyanole. If the charge of the dye is the opposite of that of the surfactant, an insoluble salt may form. In fact, such a salt generally does form [39-41] as is evident from the color change produced by sub-CMC amounts of surfactant. This precipitate often remains finely dispersed by an excess of the surfactant and escapes detection. It is this precipitate which then becomes solubilized by micelles when the concentration of surfactant is further increased, causing a second color change that is generally reported as the CMC. The indicators are mostly used in small concentrations ($10^{-4} - 10^{-5} M$) so that the amount of surfactant consumed by precipitation is often negligible compared to the CMC but can become significant for low CMC's.

The presence of this water-insoluble precipitate has, however, a more profound effect on the determination of the CMC: it maintains a saturated solution until the last of the precipitate is dissolved. This also

corresponds to the end of at least one absorbancy change, the disappearance of the color of the precipitate (the γ band for pinacyanole) [39]. Hence, it has the effect of producing the maximum lowering of the CMC through the actual determination range in many of these methods. Lowering the dye concentration reduces the fraction of surfactant used up in precipitating the dye but does not affect significantly the degree of saturation of the micelles at the color change.

Saturation by the indicator in case of *oppositely charged dyes* is particularly objectionable because of their frequently high solubility in the micelles. Thus, the solubility of pinacyanole was estimated at 15 to 20 mole percent, in the first mixed micelles that formed [39].

There are other methods of CMC determinations in which the situation is more favorable, either because saturation of the micelle is avoided or because the mole fraction at saturation is much lower. In the former category falls the use of *similarly charged dyes*. The formation of a water-insoluble, micelle-soluble salt is then avoided; the dye remains in aqueous undersaturated solution and its mole fraction in the micelle is only a corresponding fraction of saturation. Unfortunately, no methodical studies on this approach are known to us. In the same class are dye or iodine methods used with nonionic surfactants.

Some of the solubilization methods use nonionic, water-insoluble dyes whose solubility in the micelles is known to be low. "Orange OT" is the most popular of these, and here it is known [42-44] that the mole fraction at saturation is about 0.01, thus producing a generally negligible error. Some of the other solubilized indicators such as lauryl alcohol or toluene are likely to lead to mole fractions, and hence CMC changes, of the order of 10 percent or more.

The above discussion makes clear our objection to accepting values, obtained in the presence of easily solubilized indicators under conditions favoring saturation, as valid CMC measurements for the surfactant itself. They are classified as G as far as "Method Quality" is concerned, indicating that a substantial methodical error is likely to be present. On the other hand, such data could be, in principle, excellent values for the mixed system (surfactant + indicator + water). In the great majority of the cases this is not so because the conditions are not sufficiently specified and the proportion of additive often unstated. In some cases, however, we

were able to treat the data in just this way and some of the indicators may be found among the additives.

6. *The effect of impurities.* What has been said above about the effect of indicators applies to the effect of any impurities with the important provision that indicators are added consciously and generally specified in an article, whereas impurities are included inadvertently and their nature is uncertain. We have always scanned an article for clues to the presence and nature of impurities. Such clues can be found in details of the method of preparation, of analyses, and of other physical constants or properties reported. The results are included in our "quality rating" of the compound. In general we have been more skeptical of the older measurements performed before concern about the effect of impurities became widespread and modern analytical techniques were developed. We have given great weight to foam purification at a concentration below the CMC and to chromatographic methods.

The effect of impurities upon the CMC value depends both on the nature of the impurities and on the method used in the determination. We can learn something about the effects of different classes of impurities from their effects upon the CMC when they are added deliberately as "additives." Some have little effect unless present in concentrations too high ever to be reached by impurities. Hydrophilic organic compounds such as sugar or ethanol are in this class as are "inert" salts. On the other hand, oleophilic impurities and salts containing ions forming water-insoluble easily solubilized precipitates can have large effects as we have seen in connection with indicators because they tend to saturate the few micelles present at the CMC and lower the activity of surfactant in them. Higher homologues or incompletely reacted organic intermediates such as alcohols are among objectionable impurities likely to be encountered.

The case of true soaps, i.e., salts of higher carboxylic acids, merits a special mention. Here the acid itself, highly water-insoluble and easily solubilized, is the most important impurity. Some of it is normally formed by hydrolysis and must be considered as a normal constituent of the (water-surfactant) system. Additional substantial amounts are readily formed unless the presence of atmospheric carbon dioxide is carefully excluded. If precautions to this effect were not explicitly mentioned, we assumed that the system was impure and gave it a "D" rating. The effect of CO₂ can be largely neutralized by operating in the presence of a slight

excess of base. The CMC is not likely to differ substantially from that of the pure system but we have always reported such experiments among systems with additives, specifying the nature and concentration of the base used whenever possible.

The effect of an impurity upon different methods of determining the CMC is largely unknown. Some of our unpublished experiments suggest that traces have a larger effect on values obtained by plotting the equivalent conductivity than those obtained by plotting the specific conductance for the same systems. Dye indicator methods seem to give particularly low results in the presence of higher homologues or intermediates because both tend to concentrate in the first mixed micelles formed. It is likely that pinacyanole may change color by dissolving in droplets of emulsified higher alcohol far below the CMC [45].

The best studied effect is that of a surface active impurity upon the surface tension method. It is now well understood that such impurities may lower the surface tension significantly below the CMC and then become solubilized sufficiently in micelles to be more or less completely removed from the surface [6, 46-48]. This leads then to a higher surface tension after an intermediate minimum. As already mentioned, such minima have been often used as CMC values but we now take them as *prima facie* evidence of the presence of easily solubilized—and therefore significant—impurities or hydrolysis products.

An impurity leading to a minimum must have a surface activity at least comparable with that of the surfactant. If the surfactant is very surface active, the same impurity may remain unnoticed. This is shown by experiments of Harrold [49] who found that addition of salt (which increases the surface activity of the ionic surfactant much more than that of the alcohol) leads to a disappearance of the minimum along with a lowering of surface tension and of the CMC. Hence, the more surface active the surfactant, the less significant is the absence of a minimum as a criterion of purity.

When a minimum is present it becomes difficult if not impossible to obtain an accurate value for the CMC from surface tension measurements. However, even when a minimum is absent it seems that the surface tension method is particularly strongly affected by traces of impurities. This is a point which became apparent gradually in the course of our evaluations. Initially, we rated careful surface tension measurements showing no minima as BB. Later,

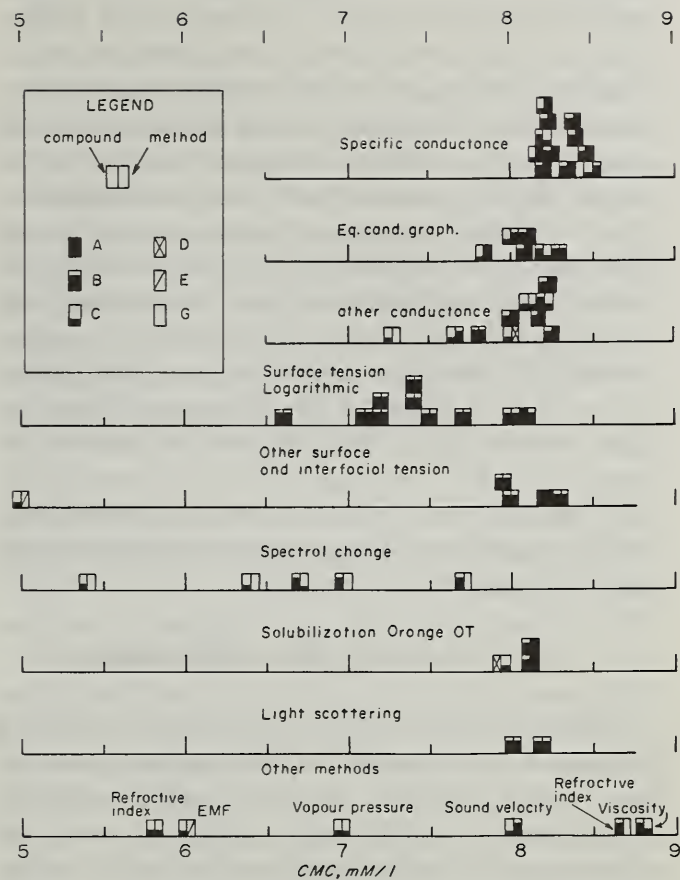


FIGURE 7. All the critical micelle concentrations for sodium dodecyl sulfate at temperatures between 20 and 30 °C, carrying a quality rating of C-G or better, obtained by various experimental methods.

Each square represents a determination and as shown in detail in the legend, the amount of black increases with the quality of the determination. It may be seen that CMC's determined by surface tension from a logarithmic plot scatter much more than other high-quality data obtained by a single method.

we found that this was often an overestimate of the accuracy of the method, although the details of the source of error remain obscure. Figure 7 illustrates our case. It shows all the measurements rated CC or better, and CG for sodium dodecyl sulfate at $25 \pm 5^\circ$. Values obtained by different methods are shown on separate lines and the shading of the points indicates their individual quality rating. It may be seen that for conductance methods and for the miscellaneous ones the better values cluster together and the methodical difference mentioned earlier between equivalent and specific conductance plots is visible. The spectral change methods give, as would be expected, widely scattered results. However, the surface tension values rated BB, although not nearly as bad as the spectral change ones, cover a surprisingly wide range and show no clustering near the CMC expected for the pure compound. Part of this may be due to differences in extrapolation, but measurements on an especially

carefully purified sample (foamed *in situ*) yielded much more acceptable values using different ranges of points. This suggests that residual impurities are to blame. It also suggests that for surfactants having greater surface activity the effect may well be reduced, but there are no data to test this hypothesis.

As a result of these considerations we did not depend on surface tension values alone for any "confirmed" result although we have used them occasionally to confirm results obtained by another method. We have also used them more liberally for the nonionic compounds both because these are more surface active and because alternatives are fewer.

For the sake of consistency, we have maintained the BB rating for this type of data throughout in the individual quality evaluations.

7. Scope of this Collection

Micelle-forming systems cover a wide variety of molecular species which have in common one feature often described as amphipathy, a built-in asymmetry in the molecule which is composed of a nonpolar, hydrophobic portion, and a polar, hydrophilic part. The systems we have covered include monomers containing straight or branched aliphatic chains, sometimes partly or completely halogenated, and sometimes partly aromatic, with a wide variety of head groups, nonionic, zwitterionic, or ionic, including a variety of counterions in the last case. The solvents are aqueous, partially or completely; in a few systems the medium is heavy water. The variables which are explicitly cataloged, are temperature, pressure (except when it is atmospheric), the nature and the concentration of additives to the medium, the experimental method used, and the type and source of the CMC data themselves.

Certain types of systems were excluded intentionally. The reasons for such exclusions are given.

The association of monomers to produce oligomers and multimers in solution is a frequently observed phenomenon. The existence of a critical micelle concentration, however, requires fairly large multimers (containing roughly 20 or more monomers) in preference to small oligomers. Thus, for example, if monomers associate to produce dimers, trimers, and higher multimers with approximately equal stepwise association constants (the association constant describing the equilibrium between the N th mer, the $(N+1)$ th mer and the monomer),

the degree of aggregation is a mild function of the total concentration; the aggregates are very polydisperse, and the system does not show any pronounced critical concentration.

The difference between systems which exhibit fairly sharp critical concentrations and those which do not is obviously a matter of degree. It seems, however, that the requirement of relatively greater stability of large multimers as compared to small oligomers is met mainly by monomeric systems which contain some flexible aliphatic chains. Those monomers which contain fused aromatic ring systems, such as dyes and other flat organic molecules, or fused alicyclic systems such as bile-salts, are expected to have, and seem to have in fact, rather diffuse and extended concentration ranges over which the degree of association increases from low to high values. For many such systems, CMC values have been reported in the literature [50]. Because of the relatively greater uncertainty of defining and determining the CMC values in such systems, particularly when the average degree of association is small, we have not extended our compilation to such systems. We have, however, recorded CMC values for many systems where the monomer is partly aromatic and partly aliphatic.

The phenomenon of micelle formation in aqueous systems is primarily a result of hydrophobic interactions. Hydrophobic interactions are a net result of a number of factors involving structural rearrangements of water molecules and solute-solvent, solvent-solvent, and solute-solute van der Waal's interactions. It is, therefore, not unlikely that in solvents other than water, similar interactions, perhaps on a reduced scale, can occur to produce other kinds of "solvophobic" bonding. In our compilation, we have many examples of mixed aqueous-nonaqueous systems including such media as 96 percent sulfuric acid. It is expected that future research will provide examples of micellization in many other completely nonaqueous systems where the micelles will have essentially the same kind of structure as in aqueous media, namely a hydrocarbonoid core and a polar interface.

When the medium is nonpolar, however, e.g., benzene or other hydrocarbon-like solvents, it is relatively improbable that the hydrophobic moiety of the monomers will have any great tendency to aggregate. For such systems, any aggregation that occurs is more likely to be due to the association of the hydrophilic groups of the monomers, which are now "solvophobic," to produce micelles of the

opposite kind of structure, i.e., "invert micelles," having a core composed of the hydrophilic groups of the monomers with the hydrocarbon groups remaining outside. Although such associations have been studied, and CMC values have been quoted [51-54], considerable uncertainty remains about their significance. We have, therefore, excluded such systems from our compilation.

Well above the CMC, in aqueous media, there occur extensive interactions between micelles, which are often difficult to unravel, particularly if the micelles are ionic. In such concentrated solutions, many physical properties of solutions exhibit breaks or kinks somewhat similar to the ones that occur at the CMC where micelles first form. The concentrations where such breaks or kinks occur are often described as "the second CMC" [55, 56]. The second CMC is not well understood, but it clearly involves changes in inter-micellar interactions as also monomer-micelle interactions. These "higher CMC" values have also been excluded from our compilation.

While the above restrictions imposed on our compilation are voluntary, there are some involuntary omissions and probable errors which must be mentioned. Of foremost concern, of course, is that many publications must have been missed altogether. A balance has had to be struck between completeness of coverage (up to December 1966) and inordinate delays in producing this report. It is hoped that not many high quality data on well-characterized systems have been omitted. It is probably unwise to hazard a guess as to what percentage of available CMC values in the literature has been missed, but we hope that it is below 10 percent. It is possible, also, that some CMC values in the publications we have examined have been unintentionally overlooked. We hope that readers of this report who note either of such omissions will draw them to our attention.

8. Methods and Techniques in the Collection of Data

This section deals with some of the problems we encountered in the retrieval of the literature pertaining to CMC measurements and in obtaining numerical values of the CMC.

Because of the wide-spread importance of and interest in surfactants in solution, CMC values appear in a wide variety of journals. As the values themselves are often of auxilliary and secondary in-

terest to the main purposes of the investigation, neither the expression "critical micelle concentration" nor the word "micelle" may appear in the title of the articles. For the standard methods of literature search using Chemical Abstracts and other such publications, we, therefore, had to use a variety of key words for searching purposes. Use was made whenever possible of published books and review articles for the entries they provide into the literature. It was found necessary, however, to make extensive use of personal appeals to various investigators in the field of surfactant solutions to provide references to published work.

To obtain numerical values of the CMC, several procedures had to be used, as indicated by the source symbols. All numerical values quoted in the publications were given the source symbol T. The majority of these values were found in tables, but a substantial number were found dispersed in the written part of the text of the papers. It was necessary, therefore, to scrutinize the whole paper carefully. Of about 4700 numerical values reported in this compilation, 3207, or 68 percent, were obtained from this "direct" source. Numerical data were thus not available in the published literature for about 1500 CMC values which had to be obtained from various "indirect" sources described below.

In many publications, CMC values themselves are presented in graphical form, or some measured physical property of surfactant solutions is plotted as some function of the concentration, so that the CMC can be obtained from the "breaks" in the curves. A large number of numerical values of this type, i.e., values which are not given in the papers, were obtained by indirect means. A total of 598 values, given the source symbol L, involve, in addition to the published data, a private communication from an author. 515 values were "read" from published graphs and are given the source symbol G. 122 values used combined sources, GL. 90 CMC values were obtained from the kinks of published graphs of some physical property vs. concentration and are given the source symbol K.

In many publications, numerical values of some physical property measured, e.g., conductance, are tabulated as a function of concentration. When such measurements seemed to be of high quality, we obtained CMC values by making plots ourselves of the physical property measured. These CMC values, 96 in all, were given the source symbol P.

A small number of values, 21, were obtained by solving published equations relating numerical

values of some physical property, particularly conductance, with concentration, below and above the CMC. These have the source symbol E.

43 CMC values have combined source symbols, e.g., TL, KL, indicating dual sources of the values. These include 13 containing the symbol A, which denotes values which have been corrected for obvious misprints.

A substantial number of entries in the compilation do not report numerical CMC values because the entries either give a cross-reference to mixed surfactant systems, involve references to published graphical data for which retrieval was uncertain, or relate to numerical values obtained using criteria which we consider questionable.

The private correspondence from several investigators contained numerical data and references to unpublished work. We have not used these data in our present compilation.

A partial justification of the use of the various "indirect" sources referred to above is that their contributions to the categories of all selected data are high. Out of 620 data included in the categories 1, 2, 3, and D, only 270 were obtained from tabulated sources, T. Thus, 56 percent of all selected data are from "indirect" sources. Similarly, of the 106 confirmed (1) and recommended (2) CMC values, 67 are from "indirect" sources, not readily available from the literature.

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STRUCTURAL INDEXES OF COMPOUNDS WITH KEYS TO COMPOUND NUMBERS

PART 1. *GROUPED BY CHARGE* (Arranged by number of carbon atoms in longest hydrophobic tail and by heads)

- A. Anionics
- B. Cationics
- C. Nonionics

PART 2. *GROUPED BY CHARGE AND HEAD GROUPS* (Arranged by number of carbon atoms in longest hydrophobic tail and by heads)

- A. Anionics
- B. Cationics
- C. Nonionics

PART 3. *GROUPED BY STRUCTURE OF HYDROPHOBIC PART* (Arranged by number of carbon atoms in longest hydrophobic tail)

- A. Alkanes by Branching and Number of Carbon Atoms
- B. Alkyl Aryl
- C. Unsaturated and Substituted
- D. Heterocyclic

PART 4. *POLYOXYETHYLENES GROUPED BY DISTRIBUTION OF HEADS* (Arranged by number of carbon atoms in longest hydrophobic tail and by heads)

- A. Homogeneous Head Groups
- B. Reduced Polydispersity of Head Groups
- C. Natural Distribution of Head Groups

PART 5. *IONICS GROUPED BY COUNTERION* (Arranged by number of carbon atoms in longest hydrophobic tail and by heads)

- A. Anionics
- B. Cationics

PART 6. *COMMERCIAL NAMES AND ILL DEFINED STRUCTURES* (Arranged alphabetically)

Structural Indexes of Compounds with Keys to Compound Numbers

PART 1. Grouped by charge and arranged by number of carbon atoms in longest hydrophobic tail and by heads.

1A. ANIONICS BY NUMBER OF CARBON ATOMS

C-1	428 PERFLUORO ACETIC ACID	605 SODIUM AMYL ALPHAPHOSPHONO PELARGONATE
C-2	429 PERFLUORO PROPIONIC ACID	260 SODIUM DI-N-OCTYL SULFOSUCCINATE
C-3	43 BUTYRIC ACID 430 PERFLUORO BUTYRIC ACID 452 3-4-DICHLOROPERFLUORO BUTYRIC/KEL-F/ACID 699 SODIUM BUTYRATE 272 TRI-ISOPROPYL BENZENE SULFONIC ACID 271 SODIUM TRI-ISO-PROPYL BENZENE SULFONATE	262 SODIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE 286 AMMONIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE 2 SODIUM OCTYL 1 SULFATE 66 SODIUM OCTYL 2 SULFATE 347 OCTYL TRIMETHYLLAMMONIUM OCTYL SULFATE 643 DODECYL TRIMETHYLLAMMONIUM OCTANE SULFATE 181 SODIUM OCTYL 1-SULFONATE 340 MAGNESIUM OCTANE SULFONATE 287 OCTYL TRIMETHYLLAMMONIUM OCTANE SULFONATE 503 SODIUM OCTYL BENZENE SULFONATE 49 SODIUM P OCTYL BENZENE SULFONATE 172 SODIUM 2-N-OCTYL BENZENE SULFONATE 510 SODIUM 2-ETHYL-HEXYL BENZENE SULFONATE 676 SODIUM 1,1,3,3-TETRAMETHYL BUTYL BENZENE SULFONATE 447 SODIUM PARA-BIS-/N-OCTYL/ BENZENE SULFONATE 677 SODIUM DI-/1,1,3,3-TETRAMETHYL BUTYL/BENZENESULFONATE
C-4	484 SODIUM PENTANOATE/VALERATE/ 257 SODIUM DI-N-BUTYL SULFOSUCCINATE 261 SODIUM DI-ISOBUTYL SULFOSUCCINATE 681 SODIUM DIBUTYL BENZENE SULFONATE 445 SODIUM PARA-BIS-/N-BUTYL/ BENZENE SULFONATE 682 SODIUM DIBUTYL NAPHTHALENE SULFONATE /NEKAL/	C-9 530 DECANOIC ACID 455 PERFLUORO DECANOIC ACID 299 SODIUM DECANOATE 90 POTASSIUM DECANOATE 702 POTASSIUM PERFLUORODECANOATE 668 DIPOTASSIUM OCTYL MALONATE 29 SODIUM NYNOL ALPHA SULFOPELARGONATE 612 ALPHAPHOSPHONO DECANOIC ACID 616 MONOSODIUM ALPHAPHOSPHONO DECANOATE 606 SODIUM AMYL ALPHAPHOSPHONO CAPRATE 136 SODIUM ALPHA DIMETHYL AMINO CAPRATE 295 SODIUM NYNOL 1-SULFATE 536 NYNOL SULFONIC ACID 504 SODIUM NYNOL BENZENE SULFONATE 493 SODIUM P-NONYL BENZENE SULFONATE 138 SODIUM NYNOL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN
C-5	700 HEXANOIC ACID 416 PERFLUORO HEXANOIC ACID 453 3-5-6-TRICHLOROPERFLUORO HEXANOIC/KEL-F/ACID 188 POTASSIUM HEXANOATE 701 POTASSIUM PERFLUOROHEXANOATE 485 SODIUM HEXANOATE/CAPROATE/ 343 SODIUM PENTANE SULFONATE 258 SODIUM DI-N-AMYL SULFOSUCCINATE	C-10 297 POTASSIUM UNDECANOATE 373 AMMONIUM EICOSAFLUOROUNDECANOATE H/CF2/10 COO NH4 425 POTASSIUM 1-1-2-DECANE TRICARBOXYLATE 30 SODIUM DECYL ALPHA SULFOPELARGONATE 33 SODIUM /OXO/ DECYL ALPHA SULFOPELARGONATE 3 SODIUM DECYL 1 SULFATE 15 SODIUM DECYL 2 SULFATE 642 OCTYL TRIMETHYLLAMMONIUM DECANE SULFATE 346 DECYL TRIMETHYLLAMMONIUM DECYL SULFATE 182 SODIUM DECYL 1-SULFONATE 341 MAGNESIUM DECANE SULFONATE 353 OCTYL TRIMETHYLLAMMONIUM DECANE SULFONATE 288 DECYL TRIMETHYLLAMMONIUM DECANE SULFONATE 505 SODIUM DECYL BENZENE SULFONATE 50 SODIUM P DECYL BENZENE SULFONATE 173 SODIUM 2-N-DECYL BENZENE SULFONATE 140 SODIUM DECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN 511 SODIUM 2-PROPYL-HEPTYL BENZENE SULFONATE
C-6	296 POTASSIUM HEPTANOATE 374 DODECAFLUOROHEPTANOIC ACID H/CF2/6COOH 486 SODIUM HEPTANOATE 355 AMMONIUM DODECAFLUOROHEPTANOATE H/CF2/6COONH4 352 SODIUM DI-1-METHYLISOAMYL SULFOSUCCINATE 259 SODIUM DI-N-HEXYL SULFOSUCCINATE 344 SODIUM HEXANE SULFONATE 339 MAGNESIUM HEXANE SULFONATE 501 SODIUM HEXYL BENZENE SULFONATE 446 SODIUM PARA-BIS-/N-HEXYL/ BENZENE SULFONATE 704 POTASSIUM 4-HEXYL RESORCINOLATE	C-11 531 DODECANOIC ACID 273 SODIUM DODECANOATE 91 POTASSIUM DODECANOATE 627 CESIUM DODECANOATE 277 BENZYL TRIMETHYL AMMONIUM DODECANOATE 527 SODIUM UNDECANE-3-CARBOXYLATE 669 DIPOTASSIUM DECYL MALONATE 602 ALPHA SULFO LAURIC ACID 235 SODIUM ALPHA SULFO LAURIC ACID 603 SODIUM PROPYL ALPHA SULFO LAURATE 613 ALPHAPHOSPHONO DODECANOIC ACID 617 MONOSODIUM ALPHAPHOSPHONO DODECANOATE 620 DISODIUM ALPHAPHOSPHONO DODECANOATE 607 SODIUM ISOPROPYL ALPHAPHOSPHONO LAURATE 311 SODIUM UNDECYL 1-SULFATE 72 SODIUM UNDECYL 3 SULFATE
C-7	529 OCTANOIC ACID 417 PERFLUORO OCTANOIC ACID 454 3-5-7-8-TETRACHLOROPERFLUORO OCTANOIC/KEL-F/ACID 476 SODIUM OCTANOATE 44 POTASSIUM OCTANOATE 456 POTASSIUM PERFLUORO OCTANOATE 284 HEXANOLAMINE-CH3CH/OH/CH2C/CH3/2NH2-OCTANOATE 303 DI-ISOPROPYLAMMONIUM CAPRYLATE 502 SODIUM HEPTYL BENZENE SULFONATE	
C-8	375 HEXADECAFLUORONONANOIC ACID H/CF2/8COOH 487 SODIUM NONANOATE 350 POTASSIUM NONANOATE 372 AMMONIUM HEXADECAFLUORONONANOATE H/CF2/8 COO NH4 420 POTASSIUM 1-1-2-OCTANE TRICARBOXYLATE 6 SODIUM ALPHA SULFOPELARGONIC ACID 25 SODIUM ETHYL ALPHA SULFOPELARGONATE 26 SODIUM AMYL ALPHA SULFOPELARGONATE 27 SODIUM HEXYL ALPHA SULFOPELARGONATE 28 SODIUM HEPTYL ALPHA SULFOPELARGONATE 7 SODIUM OCTYL ALPHA SULFOPELARGONATE 31 SODIUM 2 OCTYL ALPHA SULFOPELARGONATE 32 SODIUM 2 ETHYLHEXYL ALPHA SULFOPELARGONATE 35 SODIUM H/CF2CF2/3CH2 ALPHA SULFOPELARGONATE 611 ALPHAPHOSPHONO PELARGONIC ACID	

PART 1. Grouped by charge and arranged by number of carbon atoms in longest hydrophobic tail and by heads—Continued

80 SODIUM UNDECYL 6 SULFATE
 462 SODIUM MONOLAURIN SULFATE
 639 SODIUM UNDECYL THIOSULFATE
 537 UNDECYL SULFONIC ACID
 418 SODIUM UNDECYL SULFONATE
 45 SODIUM P 1 METHYL DECYL BENZENE SULFONATE
 678 SODIUM 1,1-DIMETHYLNONYL BENZENE SULFONATE
 679 SODIUM 3-BUTYL HEPTYL BENZENE SULFONATE

C-12

351 POTASSIUM TRIDEcanoate
 421 POTASSIUM 1-1-2-DODECANE TRICARBOXYLATE
 34 SODIUM DODECYL ALPHA SULFOPELARGONATE
 632 POTASSIUM N-DODECYL BETA-ALANINATE
 705 DODECYL SULFURIC ACID
 1 SODIUM DODECYL 1 SULFATE
 67 SODIUM DODECYL 2 SULFATE
 634 POTASSIUM DODECYL SULFATE
 111 LITHIUM DODECYL 1 SULFATE
 23 SILVER DODECYL 1 SULFATE
 568 MAGNESIUM DODECYL SULFATE
 24 CALCIUM DODECYL 1 SULFATE
 569 STRONTIUM DODECYL SULFATE
 572 COBALTOUS DODECYL SULFATE
 573 CUPRIC DODECYL SULFATE
 570 LEAD DODECYL SULFATE
 571 MANGANESE DODECYL SULFATE
 575 NICKEL DODECYL SULFATE
 574 ZINC DODECYL SULFATE
 386 AMMONIUM DODECYL SULFATE
 387 METHYLAMMONIUM DODECYL SULFATE
 388 ETHYLAMMONIUM DODECYL SULFATE
 389 BUTYLAMMONIUM DODECYL SULFATE
 112 TETRAMETHYL AMMONIUM DODECYL 1 SULFATE
 383 BUTYL TRIMETHYLMONIUM DODECYL SULFATE
 382 ETHYL TRIMETHYLMONIUM DODECYL SULFATE
 718 TETRAETHYLAMMONIUM DODECYL SULFATE
 719 TETRABUTYLMONIUM DODECYL SULFATE
 720 1-6-DITRIMETHYLMONIUM-HEXANE/DODECYL SULFATE/2
 409 TRIETHANOLAMMONIUM DODECYL SULFATE
 410 MORPHOLINIUM DODECYL SULFATE
 391 OCTYLAMMONIUM DODECYL SULFATE
 385 OCTYL TRIMETHYLMONIUM DODECYL SULFATE
 280 DECYL TRIMETHYL AMMONIUM DODECYL SULFATE
 281 DODECYL TRIMETHYLMONIUM DODECYL SULFATE
 562 SODIUM DODECENYL SULFATE
 636 SODIUM DODECYL THIOSULFATE
 541 SODIUM DODECYL MONO-OXYETHYLENE SULFATE
 542 SODIUM DODECYL DIOXYETHYLENE SULFATE
 113 SODIUM DODECYL TRI-OXYETHYLENE SULFATE
 543 SODIUM DODECYL TETRA-OXYETHYLENE SULFATE
 114 SODIUM DODECYL /OXYETHYLENE/17.5 SULFATE
 597 SODIUM DODECYL MONO-OXYPROPYL SULFATE
 200 DODECYL SULFONIC ACID
 179 SODIUM DODECANE 1-SULFONATE
 175 SODIUM DODECANE 2-SULFONATE
 40 POTASSIUM DODECYL 1 SULFONATE
 635 LITHIUM DODECYL SULFONATE
 342 MAGNESIUM DODEGANE SULFONATE
 238 SODIUM DODECANE 1-HYDROXY 2-SULFONATE
 247 DODECANE 1-HYDROXY 2-SULFONIC ACID
 506 SODIUM DODECYL BENZENE SULFONATE
 51 SODIUM P DODECYL BENZENE SULFONATE
 171 SODIUM 2-N-DODECYL BENZENE SULFONATE
 301 SODIUM 3-N-DODECYL BENZENE SULFONATE
 302 SODIUM 4-N-DODECYL BENZENE SULFONATE
 514 SODIUM 6-N-DODECYL BENZENE SULFONATE
 512 SODIUM 2-BUTYL-OCTYL BENZENE SULFONATE
 680 SODIUM 2-PROPYL-4-METHYL-OCTYL BENZENE SULFONATE
 492 SODIUM TETRAPOPYLENE/1-3-5-7-TETRAMETHYL-
 OCTYL/BENZENE SULFONATE
 139 SODIUM DODECYL BENZENE SULFONATE BRANCHED HYDROCARBON
 CHAIN

C-13

532 TETRADECANOIC ACID
 298 SODIUM TETRADECANOATE
 92 POTASSIUM TETRADECANOATE
 670 DIPOTASSIUM DODECYL MALONATE
 189 ALPHA SULFOMYRISTIC ACID
 236 SODIUM ALPHA SULFO MYRISTIC ACID
 604 SODIUM METHYL ALPHA SULFO MYRISTATE
 233 DISODIUM ALPHA SULFO MYRISTATE
 614 ALPHAPHOSPHONO TETRADECANOIC ACID

618 MONOSODIUM ALPHAPHOSPHONO TETRADECANOATE
 621 DISODIUM ALPHAPHOSPHONO TETRADECANOATE
 624 TRISODIUM ALPHAPHOSPHONO TETRADECANOATE
 608 SODIUM METHYL ALPHAPHOSPHONO MYRISTATE
 68 SODIUM TRIDECYL 2 SULFATE
 83 SODIUM TRIDECYL 7 SULFATE
 242 TRIDECANE 1-SULFONIC ACID
 229 SODIUM TRIDECANE 1-SULFONATE
 46 SODIUM P 1 METHYL DODECYL BENZENE SULFONATE
 141 SODIUM TRIDECYL BENZENE SULFONATE BRANCHED HYDROCARBON
 CHAIN

C-14

426 POTASSIUM 1-1-2-TETRADECANE TRICARBOXYLATE
 4 SODIUM TETRADECYL 1 SULFATE
 16 SODIUM TETRADECYL 2 SULFATE
 73 SODIUM TETRADECYL 3 SULFATE
 17 SODIUM TETRADECYL 4 SULFATE
 77 SODIUM TETRADECYL 5 SULFATE
 525 SODIUM TETRADECYL 6-SULFATE
 84 SODIUM TETRADECYL 7 SULFATE
 526 SODIUM 2-DI-N-HEXYL ETHYL SULFATE
 637 LITHIUM TETRADECYL SULFATE
 576 CUPRIC TETRADECYL SULFATE
 544 SODIUM TETRADECYL MONO-OXYETHYLENE SULFATE
 545 SODIUM TETRADECYL DI-OXYETHYLENE SULFATE
 546 SODIUM TETRADECYL TRI-OXYETHYLENE SULFATE
 598 SODIUM TETRADECYL MONO-OXYPROPYL SULFATE
 599 SODIUM TETRADECYL DI-OXYPROPYL SULFATE
 243 TETRADECANE 1-SULFONIC ACID
 183 SODIUM TETRADECYL 1-SULFONATE
 176 SODIUM TETRADECANE 2-SULFONATE
 248 TETRADECANE 1-HYDROXY 2-SULFONIC ACID
 239 SODIUM TETRADECANE 1-HYDROXY 2-SULFONATE
 507 SODIUM TETRADECYL BENZENE SULFONATE
 174 SODIUM 2-N-TETRADECYL BENZENE SULFONATE
 513 SODIUM 2-AMYL-NONYL BENZENE SULFONATE

C-15

533 HEXADECANOIC ACID
 300 SODIUM HEXADECANOATE
 185 POTASSIUM HEXADECANOATE
 671 DIPOTASSIUM TETRADECYL MALONATE
 190 ALPHA SULFOPALMITIC ACID
 237 SODIUM ALPHA SULFO PALMITIC ACID
 234 DISODIUM ALPHA SULFO PALMITATE
 36 SODIUM METHYL ALPHA SULFOPALMITATE
 192 SODIUM ETHYL ALPHA SULFOPALMITATE
 193 SODIUM PROPYL ALPHA SULFOPALMITATE
 197 DISODIUM 2 SULFOETHYL ALPHA SULFOPALMITATE
 615 ALPHAPHOSPHONO HEXADECANOIC ACID
 619 MONOSODIUM ALPHAPHOSPHONO HEXADECANOATE
 622 DISODIUM ALPHAPHOSPHONO HEXADECANOATE
 625 TRISODIUM ALPHAPHOSPHONO HEXADECANOATE
 609 SODIUM ISOPROPYL ALPHAPHOSPHONO PALMITATE
 69 SODIUM PENTADECYL 2 SULFATE
 74 SODIUM PENTADECYL 3 SULFATE
 78 SODIUM PENTADECYL 5 SULFATE
 85 SODIUM PENTADECYL 8 SULFATE
 244 PENTADECANE 1-SULFONIC ACID
 230 SODIUM PENTADECANE 1-SULFONATE
 47 SODIUM P 1 METHYL TETRADECYL BENZENE SULFONATE
 142 SODIUM PENTADECYL BENZENE SULFONATE BRANCHED
 HYDROCARBON CHAIN
 596 TETRAHYDROANACARDOL AMMONIUM MONOSULFONATE

C-16

422 POTASSIUM 1-1-2-HEXADECANE TRICARBOXYLATE
 5 SODIUM HEXADECYL 1 SULFATE
 75 SODIUM HEXADECYL 4 SULFATE
 81 SODIUM HEXADECYL 6 SULFATE
 86 SODIUM HEXADECYL 8 SULFATE
 638 LITHIUM HEXADECYL SULFATE
 577 CUPRIC HEXADECYL SULFATE
 60 TRIETHANOL AMMONIUM HEXADECYL SULFATE
 52 SODIUM HEXADECYL MONO OXYETHYLENE SULFATE
 53 SODIUM HEXADECYL DI OXYETHYLENE SULFATE
 54 SODIUM HEXADECYL TRI OXYETHYLENE SULFATE
 55 SODIUM HEXADECYL TETRA OXYETHYLENE SULFATE
 600 SODIUM HEXADECYL MONO-OXYPROPYL SULFATE
 245 HEXADECANE 1-SULFONIC ACID
 184 SODIUM HEXADECYL 1-SULFONATE
 177 SODIUM HEXADECANE 2-SULFONATE
 408 POTASSIUM HEXADECANE 1-SULFONATE

PART 1. Grouped by charge and arranged by number of carbon atoms in longest hydrophobic tail and by heads—Continued

249	HEXADECANE 1-HYDROXY 2-SULFONIC ACID	C-6
240	SODIUM HEXADECANE 1-HYDROXY 2-SULFONATE	708 HEXYLAMINE
508	SODIUM HEXADECYL BENZENE SULFONATE	709 HEXYLAMINE HYDROCHLORIDE
C-17		390 HEXYLAMMONIUM DODECYL SULFATE
534	OCTADECANOIC ACID	640 HEXYL TRIMETHYLAMMONIUM HEXANE SULFATE
448	SODIUM OCTADECANOATE /STEARATE/	644 HEXYL TRIMETHYLAMMONIUM HEXANE SULFONATE
256	POTASSIUM STEARATE	641 HEXYL TRIMETHYLAMMONIUM OCTANE SULFATE
672	DIPOTASSIUM HEXADECYL MALONATE	384 HEXYL TRIMETHYLAMMONIUM DODECYL SULFATE
263	SODIUM OLEATE /CIS-9-OCTADECENOATE/	354 HEXYL BENZYL DIMETHYLAMMONIUM CHLORIDE
305	POTASSIUM OLEATE/ CIS-9-OCTADECENOATE/	
285	HEXANOLAMINE-CH ₃ CH/OH/CH ₂ C/CH ₃ /2NH ₂ -OLEATE	
264	SODIUM ELAIDATE /TRANS-9-OCTADECENOATE/	392 OCTYLAMMONIUM CHLORIDE
629	POTASSIUM ELAIDATE/TRANS-9-OCTADECENOATE/	391 OCTYLAMMONIUM DODECYL SULFATE
283	HEXANOLAMINE-CH ₃ CH/OH/CH ₂ C/CH ₃ /2NH ₂ -ELAIDATE	93 OCTYL TRIMETHYL AMMONIUM BROMIDE
630	POTASSIUM RICINOLEATE/12 HYDROXY OLEATE/	347 OCTYL TRIMETHYLAMMONIUM OCTYL SULFATE
631	POTASSIUM RICINELAIDATE/12 HYDROXY ELAIDATE/	287 OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE
255	POTASSIUM 9,10 DIHYDROXY STEARATE	642 OCTYL TRIMETHYLAMMONIUM DECANE SULFATE
191	ALPHA SULFOSTEARIK ACID	353 OCTYL TRIMETHYLAMMONIUM DECANE SULFONATE
553	XYLYL SULFOSTEARIK ACID	385 OCTYL TRIMETHYLAMMONIUM DODECYL SULFATE
9	SODIUM ALPHA SULFOSTEARIK ACID	483 DIOCYTL DIMETHYL AMMONIUM CHLORIDE
10	DISODIUM ALPHA SULFOSTEARATE	135 OCTYL C BETAINE HYDROCHLORIDE
194	SODIUM METHYL ALPHA SULFOSTEARATE	359 OCTYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
195	SODIUM ETHYL ALPHA SULFOSTEARATE	100 OCTYL PYRIDINIUM BROMIDE
196	SODIUM PROPYL ALPHA SULFOSTEARATE	451 PARA DI-ISOBUTYLPHENOXYETHOXYETHYL DIMETHYLBENZYL AMMONIUM CHLORIDE/HYAMINE 1622/
199	SODIUM ISOPROPYL ALPHA SULFOSTEARATE	
198	DISODIUM 2 SULFOETHYL ALPHA SULFOSTEARATE	
551	PHENYL SULFOSTEARIK ACID	
11	SODIUM ALPHA SULFO PHENYL STEARIC ACID	94 NONYL TRIMETHYL AMMONIUM BROMIDE
8	DISODIUM ALPHA SULFOPHENYLSTEARATE	137 ALPHA DIMETHYLAMINO CAPRIC ACID HYDROCHLORIDE
552	TOLYL SULFOSTEARIK ACID	
14	DISODIUM 9,10 DIHYDROXY ALPHA SULFOSTEARATE	
12	SODIUM 9,10 DICHLORO ALPHA SULFOSTEARIK ACID	37 DECYLAMMONIUM CHLORIDE
13	DISODIUM 9,10 DICHLORO ALPHA SULFOSTEARATE	411 DECYLAMMONIUM ACETATE
623	DISODIUM ALPHAPHOSPHONO OCTADECANOATE	203 DECYL TRIMETHYL AMMONIUM CHLORIDE
626	TRISODIUM ALPHAPHOSPHONO OCTADECANOATE	95 DECYL TRIMETHYL AMMONIUM BROMIDE
610	SODIUM METHYL ALPHAPHOSPHONO STEARATE	306 DECYL TRIMETHYLAMMONIUM SULFATE
70	SODIUM HEPTADECYL 2 SULFATE	346 DECYL TRIMETHYLAMMONIUM DECYL SULFATE
87	SODIUM HEPTADECYL 9 SULFATE	288 DECYL TRIMETHYLAMMONIUM DECANESULFONATE
246	HEPTADECANE 1-SULFONIC ACID	280 DECYL TRIMETHYL AMMONIUM DODECYL SULFATE
231	SODIUM HEPTADECANE 1-SULFONATE	356 DECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
48	SODIUM P 1 METHYL HEXADECYL BENZENE SULFONATE	360 DECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
C-18		458 DECYL PYRIDINIUM IODIDE
64	SODIUM OCTADECYL 1 SULFATE	
71	SODIUM OCTADECYL 2 SULFATE	
76	SODIUM OCTADECYL 4 SULFATE	
82	SODIUM OCTADECYL 6 SULFATE	
65	TRIETHANOLAMMONIUM OCTADECYL 1 SULFATE	C-11
61	SODIUM OLEYL/CIS 9 OCTADECENYL/ SULFATE	96 UNDECYL TRIMETHYL AMMONIUM BROMIDE
62	SODIUM ELAIDYL/TRANS 9 OCTADECENYL/SULFATE	101 UNDECYL PYRIDINIUM BROMIDE
250	OCTADECANE 1-HYDROXY 2-SULFONIC ACID	304 CATOL 607 (N-/LAURYL-COLAMINO-FORMYL-METHYL/ - PYRIDINIUM CHLORIDE)
63	SODIUM 9 10 DICHLORO OCTADECYL SULFATE	560 EMULSOL 607L (N-(2-KETO-2-(2,-LAUROYLOXYETHYL AMINO)) ETHYL) PYRIDINIUM CHLORIDE
56	SODIUM OCTADECYL MONO OXYETHYLENE SULFATE	
57	SODIUM OCTADECYL DI OXYETHYLENE SULFATE	
58	SODIUM OCTADECYL TRI OXYETHYLENE SULFATE	
59	SODIUM OCTADECYL TETRA OXYETHYLENE SULFATE	
547	SODIUM OLEYL MONO-OXYETHYLENE SULFATE	C-12
548	SODIUM OLEYL DI-OXYETHYLENE SULFATE	38 DODECYL AMMONIUM CHLORIDE
549	SODIUM OLEYL TRI-OXYETHYLENE SULFATE	628 DODECYLAMMONIUM BROMIDE
601	SODIUM OCTADECYL MONO-OXYPROPYL SULFATE	482 DODECYL AMMONIUM NITRATE
232	OCTADECANE 1-SULFONIC ACID	412 DODECYLAMMONIUM ACETATE
349	SODIUM OCTADECANE 1-SULFONATE	449 DODECYLMETHYL AMMONIUM CHLORIDE
178	SODIUM OCTADECANE 2-SULFONATE	450 DODECYLDIMETHYL AMMONIUM CHLORIDE
419	SODIUM ALPHA-HEPTYL UNDECYL SULFONATE	399 DODECYL DIMETHYL ETHYLAMMONIUM CHLORIDE
241	SODIUM OCTADECANE 1-HYDROXY 2-SULFONATE	345 DODEDECYL DIMETHYLAMMONIUM CHLORIDE
509	SODIUM OCTADECYL BENZENE SULFONATE	41 DODECYL TRIMETHYL AMMONIUM CHLORIDE
C-19		97 DODECYL TRIMETHYL AMMONIUM BROMIDE
673	DIPOTASSIUM OCTADECYL MALONATE	126 DODECYL TRIMETHYLAMMONIUM IODIDE
79	SODIUM NONADECYL 5 SULFATE	130 DODECYL TRIMETHYL AMMONIUM FLUORIDE
88	SODIUM 1 NYONYL DECYL SULFATE	131 DODECYL TRIMETHYLAMMONIUM NITRATE
C-20		129 DODECYL TRIMETHYL AMMONIUM BROMATE
683	SODIUM EICOSYLBENZENE SULFONATE	127 DODECYL TRIMETHYL AMMONIUM IODATE
C-29		307 DODECYL TRIMETHYLAMMONIUM SULFATE
89	SODIUM 1 TETRADECYL PENTADECYL SULFATE	128 DODECYL TRIMETHYL AMMONIUM FORMATE
1B. CATIONICS BY NUMBER OF CARBON ATOMS		643 DODECYL TRIMETHYLAMMONIUM OCTANE SULFATE
C-3		281 DODECYL TRIMETHYLAMMONIUM DODECYL SULFATE
706	PERFLUORO PROPYLAMINE	400 DODECYL METHYL DIETHYLAMMONIUM CHLORIDE
707	PERFLUORO PROPYLAMINE HYDROCHLORIDE	401 DODECYL TRIETHYLAMMONIUM CHLORIDE
		22 DIMETHYL DODECYL AMINE OXIDE HYDROCHLORIDE
		293 DODECYL DIMETHYL AMINE OXIDE HYDROBROMIDE
		633 N-DODECYL BETA-ALANINE HYDROCHLORIDE
		124 DODECYL N BETAINE HYDROCHLORIDE
		500 DODECYL TRI/2-HYDROXYETHYL/AMMONIUM CHLORIDE
		279 DODECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
		403 DODECYL DIMETHYLPHENYLAMMONIUM CHLORIDE
		404 DODECYL DIMETHYL 2-PHENYLETHYLAMMONIUM CHLORIDE
		C6H5CH ₂ CH ₂ /N/CH ₃ /2/C12H25

PART 1. Grouped by charge and arranged by number of carbon atoms in longest hydrophobic tail and by heads—Continued

407 DODECYL DIMETHYL 3-PHENYLPROPYLAMMONIUM CHLORIDE
C6H5CH2CH2/N/CH3/2/C12H25
405 DODECYL METHYLETHYLBENZYLAMMONIUM CHLORIDE
371 DODECYL 3-4-DIMETHOXYBENZYL DIMETHYLAMMONIUM CHLORIDE
370 DODECYL 3-4-METHYLENEDIOXYBENZYL DIMETHYLAMMONIUM CHLORIDE
365 DODECYL 2-CHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
367 DODECYL 4-CHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
366 DODECYL 2-4-DICHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
369 DODECYL 3-4-DICHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
406 DODECYL DIMETHYL META-TRIFLUOROMETHYLBENZYL AMMONIUM CHLORIDE CF3C6H4CH2/N/CH3/2/C12H25
361 DODECYL 4-NITROBENZYL DIMETHYL AMMONIUM CHLORIDE
368 DODECYL 2-HYDROXY-5-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
278 DODECYL PYRIDINIUM CHLORIDE
290 DODECYL PYRIDINIUM BROMIDE
376 DODECYL PYRIDINIUM IODIDE
717 DODECYLQUINOLINIUM BROMIDE
528 DODECYL TROPYLIUM PERCHLORATE
460 DODECYL TROPYLIUM MONOPHOSPHATE
491 DODECYL TROPYLIUM BISULFATE

C-13

402 TRIDECYL TRIMETHYLAMMONIUM CHLORIDE

C-14

39 TETRADECYL AMMONIUM CHLORIDE
413 TETRADECYLAMMONIUM ACETATE
42 TETRADECYL TRIMETHYL AMMONIUM CHLORIDE
98 TETRADECYL TRIMETHYL AMMONIUM BROMIDE
308 TETRADECYL TRIMETHYLAMMONIUM SULFATE
291 TETRADECYL TRIPROPYLAAMMONIUM BROMIDE
125 TETRADECYL N BETAINE HYDROCHLORIDE
357 TETRADECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
362 TETRADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
102 TETRADECYL PYRIDINIUM BROMIDE
479 TETRADECYL PYRIDINIUM IODIDE

C-16

186 HEXADECYL AMMONIUM CHLORIDE
414 HEXADECYLAMMONIUM ACETATE
478 HEXADECYL DIMETHYLETHYLAMMONIUM BROMIDE
265 HEXADECYL TRIMETHYL AMMONIUM CHLORIDE
99 HEXADECYL TRIMETHYLAMMONIUM BROMIDE
309 HEXADECYL TRIMETHYLAMMONIUM SULFATE
666 HEXADECYL TRIBUTYLAAMMONIUM BROMATE
292 HEXADECYL TRIPROPYLAAMMONIUM BROMIDE
266 HEXADECYL DIMETHYL 2-HYDROXYETHYL AMMONIUM CHLORIDE
267 HEXADECYL DI-/2-HYDROXYETHYL/METHYL AMMONIUMCHLORIDE
268 HEXADECYL TRI-/2-HYDROXYETHYL/ AMMONIUM CHLORIDE
269 HEXADECYLDIMETHYL 2.3-DIHYDROXYPROPYLAMMONIUMCHLORIDE
275 HEXADECYL DIMETHYLBENZYLAMMONIUM CHLORIDE
363 HEXADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
274 HEXADECYL PYRIDINIUM CHLORIDE
427 HEXADECYL PYRIDINIUM BROMIDE
480 HEXADECYL PYRIDINIUM IODIDE
660 HEXADECYL PYRIDINIUM IODATE
693 N-CETYL 2-METHYL PYRIDINIUM CHLORIDE
696 N-CETYL 2-METHYL PYRIDINIUM IODIDE
694 N-CETYL 3-METHYL PYRIDINIUM CHLORIDE
697 N-CETYL-3-METHYL PYRIDINIUM IODIDE
695 N-CETYL 4-METHYL PYRIDINIUM CHLORIDE
698 N-CETYL-4-METHYL PYRIDINIUM IODIDE

C-18

187 OCTADECYL AMMONIUM CHLORIDE
415 OCTADECYLAMMONIUM ACETATE
270 OCTADECYL TRIMETHYLAMMONIUM CHLORIDE
477 OCTADECYL TRIMETHYLAMMONIUM BROMIDE
654 OCTADECYL TRIMETHYLAMMONIUM NITRATE
658 OCTADECYL TRIMETHYLAMMONIUM BROMATE
659 OCTADECYL TRIMETHYLAMMONIUM FORMATE
667 OCTADECYL TRIMETHYLAMMONIUM OXALATE
662 OCTADECYL TRIETHYLAMMONIUM BROMATE
663 OCTADECYL TRIPROPYLAAMMONIUM BROMATE
664 OCTADECYL TRIBUTYLAAMMONIUM BROMATE
665 OCTADECYL TRIAMYLAMMONIUM BROMATE
358 OCTADECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
364 OCTADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
655 OCTADECYL PYRIDINIUM CHLORIDE
657 OCTADECYL PYRIDINIUM BROMIDE
481 OCTADECYL PYRIDINIUM IODIDE
656 OCTADECYL PYRIDINIUM NITRATE
661 OCTADECYL PYRIDINIUM IODATE

1C. NONIONICS BY NUMBER OF CARBON ATOMS

C-2

645 PARA/BETA-D-GLUCOSYL/ETHYLBENZENE
C-3
43 BUTYRIC ACID
706 PERFLURO PROPYLAMINE
646 PARA/BETA-D-GLUCOSYL/PROPYL BENZENE

C-4

647 PARA/BETA-D-GLUCOSYL/BUTYLBENZENE
650 PARA/BETA-D-XYLOSYL/BUTYL BENZENE
461 BUTYL/OXYETHYLENE/1 ALCOHOL--BUTYL GLYCOL ETHER HOMOGENEOUS HEAD GROUP
393 BUTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP
394 1-METHYL PROPYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP
674 DIISOBUTYL BENZENE/OXYETHYLENE/6 ALCOHOL (DB) BRANCHED CHAIN. NATURAL OE DISTRIBUTION

C-5

700 HEXANOIC ACID

C-6

708 HEXYLAMINE
714 HEXYL DIMETHYL AMINE OXIDE
686 HEXYL SULFINYLBUTANOL
684 HEXYL SULFINYLETHANOL
685 HEXYL SULFINYLPROPANOL
687 HEXYL SULFINYLPEPTANOL
457 1-4-HEXANEDIOL
703 4-HEXYL RESORCINOL
103 HEXYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
380 HEXYL/OXYETHYLENE/4 ALCOHOL HOMOGENEOUS HEAD GROUP
381 HEXYL/OXYETHYLENE/5 ALCOHOL HOMOGENEOUS HEAD GROUP
294 HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP
395 2-ETHYL BUTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP

C-7

529 OCTANOIC ACID
715 HEPTYL DIMETHYL AMINE OXIDE
586 METHYL /OXYETHYLENE/ 7.6 OCTANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS

C-8

118 OCTYL N BETAINE
132 OCTYL C BETAINE
251 OCTYL DIMETHYL AMINE OXIDE
348 NN-DIMETHYL 1-1-DIHYDROENTADECAFLUORO OCTYL AMINE N-OXIDE /C7F15CH2N/CH3/20/
710 OCTYL DIMETHYL PHOSPHINE OXIDE
688 OCTYL SULFINYLETHANOL
689 OCTYL SULFINYLPROPANOL
690 OCTYL SULFINYLBUTANOL
691 OCTYL METHYL SULFOXIDE
648 ALPHA-D-GLUCOSYL OCTANE
424 OCTYL ALPHA-GLYCERYL ETHER
18 OCTYL BETA D GLUCOSIDE
423 OCTYL/OXYETHYLENE/1 ALCOHOL--OCTYL GLYCOL ETHER HOMOGENEOUS HEAD GROUP
104 OCTYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
105 OCTYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP
106 OCTYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP
515 ISO-OCTYL/OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
396 2-PROPYL PENTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP
335 OCTYL BENZENE/OXYETHYLENE/8.5 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
675 OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
317 T-OCTYL BENZENE/OXYETHYLENE/10 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS
318 T-OCTYL BENZENE/OXYETHYLENE/20 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS
319 T-OCTYL BENZENE/OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS
207 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL HOMOGENOUS HEAD GROUP
217 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS

PART 1. Grouped by charge and arranged by number of carbon atoms in longest hydrophobic tail and by heads—Continued

208 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL HOMOGENOUS HEAD GROUP
 218 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 209 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL HOMOGENOUS HEAD GROUP
 219 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 210 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL HOMOGENOUS HEAD GROUP
 220 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 211 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL HOMOGENOUS HEAD GROUP
 221 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 212 P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP
 222 P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 213 P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL HOMOGENOUS HEAD GROUP
 223 P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 214 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL HOMOGENOUS HEAD GROUP
 224 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 215 P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL HOMOGENOUS HEAD GROUP
 223 P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 214 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL HOMOGENOUS HEAD GROUP
 224 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 215 P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL HOMOGENOUS HEAD GROUP
 225 P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 206 TRITON X-100(P-T-OCTYL BENZENE/OXYETHYLENE/9NATURAL DISTRIBUTION OF HEAD GROUPS
 216 P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL HOMOGENOUS HEAD GROUP
 226 P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 227 P-T-OCTYL BENZENE /OXYETHYLENE/16 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 228 P-T-OCTYL BENZENE /OXYETHYLENE/40 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS

C-9

530 DECANOIC ACID
 254 NONYL DIMETHYL AMINE OXIDE
 580 METHYL /OXYETHYLENE/ 7.0 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
 581 METHYL /OXYETHYLENE/ 10.3 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
 578 METHYL /OXYETHYLENE/ 11.9 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
 582 METHYL /OXYETHYLENE/ 16.0 DECANOATE NATURAL DISTRIBUTION OF HEAD GROUPS
 516 NONYL /OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 153 NONYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 336 NONYL BENZENE/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 154 NONYL BENZENE /OXYETHYLENE/ 9.8 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 320 NONYL BENZENE/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 167 NONYL BENZENE /OXYETHYLENE/ 10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 337 NONYL BENZENE/OXYETHYLENE/10.5 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 463 NONYL BENZENE/OXYETHYLENE/11 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 321 NONYL BENZENE/OXYETHYLENE/15 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 464 NONYL BENZENE/OXYETHYLENE/15 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 168 NONYL BENZENE /OXYETHYLENE/ 15 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION

155 NONYL BENZENE /OXYETHYLENE/ 15.4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 322 NONYL BENZENE/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 156 NONYL BENZENE /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 323 NONYL BENZENE/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 721 NONYL BENZENE /OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN DIALYSED
 169 NONYL BENZENE /OXYETHYLENE/ 30 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 157 NONYL BENZENE /OXYETHYLENE/ 30.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 324 NONYL BENZENE/OXYETHYLENE/50 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 338 NONYL BENZENE /OXYETHYLENE/100 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION

C-10

119 DECYL N BETAINE
 133 DECYL C BETAINE
 252 DECYL DIMETHYL AMINE OXIDE
 711 DECYL DIMETHYL PHOSPHINE OXIDE
 587 DECYL DIMETHYLAAMMONIOPROPANE SULFONATE
 692 DECYL SULFINYLETHANOL
 19 DECYL BETA D GLUCOSIDE
 201 DECYL /OXYETHYLENE/8 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS
 202 DECYL /OXYETHYLENE/11 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS
 204 DECYL /OXYETHYLENE/12 OXYMETHYL REDUCED POLYDISPERSION
 107 DECYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
 379 DECYL /OXYETHYLENE/4 ALCOHOL HOMOGENEOUS HEAD GROUP
 519 DECYL/OXO-PROCESS/ /OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 378 DECYL/OXYETHYLENE/5 ALCOHOL HOMOGENEOUS HEAD GROUP
 143 DECYL /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 108 DECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP
 109 DECYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP
 144 DECYL /OXYETHYLENE/ 10.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 517 DECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 145 DECYL /OXYETHYLENE/ 14.9 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 146 DECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 147 DECYL /OXYETHYLENE/ 29.4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 397 2-BUTYL HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP
 398 2-BUTYL HEXYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP
 523 DECYL BENZENE PARA/OXYETHYLENE/11.0 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 522 DECYL BENZENE ORTHO/OXYETHYLENE/11.2 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS

C-11

531 DODECANOIC ACID
 120 UNDECYL N BETAINE
 713 UNDECYL DIMETHYL AMINE OXIDE
 495 SUCROSE MONOLAURATE
 276 NONAETHYLENE GLYCOL MONODODECANOATE NATURAL DISTRIBUTION OF HEAD GROUPS
 583 METHYL /OXYETHYLENE/ 6.0 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
 584 METHYL /OXYETHYLENE/ 8.4 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
 585 METHYL /OXYETHYLENE/ 11.2 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
 579 METHYL /OXYETHYLENE/ 12.5 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
 440 /OXYETHYLENE/4 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
 441 /OXYETHYLENE/8 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
 442 /OXYETHYLENE/10 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
 443 /OXYETHYLENE/20 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
 444 /OXYETHYLENE/25 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS

PART 1. Grouped by charge and arranged by number of carbon atoms in longest hydrophobic tail and by heads—Continued

520 UNDECYL/OXYETHYLENE/8 ALCOHOL NATURAL DISTRIBUTION OF
HEAD GROUPS
550 LAURIC ACID DIETHANOLAMINE CONDENSATE

C-12

121 DODECYL N BETAINE
592 DODECYL N-DIETHYL N-BETAINE
134 DODECYL C BETAINE
21 DIMETHYL DODECYL AMINE OXIDE
712 DODECYL DIMETHYL PHOSPHINE OXIDE
590 DODECYL DIMETHYLMONIOPROPANE CARBOXYLATE
588 DODECYL DIMETHYLMONIOPROPANE SULFONATE
593 DIMETHYL DODECYLAMMONIOPROPANE HYDROXY SULFONATE
594 DODECYL DIPROPYL AMMONIOPROPANE SULFONATE
595 DODECYL DIMETHYL AMMONIOPROPANE SULFATE
591 DODECYL DIMETHYLPHOSPHONIOPROPANE SULFONATE
649 ALPHA-D-GLUCOSYL DODECANE
20 DODECYL BETA D GLUCOSIDE
205 DODECYL /OXYETHYLENE/12 OXYMETHYL REDUCED
POLYDISPERSION OF HEAD GROUPS
325 DODECYL/OXYETHYLENE/4 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
377 DODECYL/OXYETHYLENE/ 5 ALCOHOL HOMOGENEOUS HEAD GROUP
110 DODECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP
468 DODECYL/OXYETHYLENE/6.2 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
488 DODECYL/OXYETHYLENE/7 ALCOHOL HOMOGENEOUS HEAD GROUP
115 DODECYL /OXYETHYLENE/7 ALCOHOL REDUCED POLYDISPERSION
OF HEAD GROUPS
469 DODECYL/OXYETHYLENE/7.6 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
163 DODECYL /OXYETHYLENE/ 8 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
489 DODECYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP
310 DODECYL/OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF
HEAD GROUPS
470 DODECYL/OXYETHYLENE/10.6 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
490 DODECYL/OXYETHYLENE/12 ALCOHOL HOMOGENEOUS HEAD GROUP
164 DODECYL /OXYETHYLENE/ 12 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
326 DODECYL/OXYETHYLENE/14 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
471 DODECYL/OXYETHYLENE/15.8 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
716 DODECYL /OXYETHYLENE/16 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
165 DODECYL /OXYETHYLENE/ 18 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
472 DODECYL/OXYETHYLENE/20.4 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
327 DODECYL/OXYETHYLENE/23 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
166 DODECYL /OXYETHYLENE/ 23 ALCOHOL NATURAL DISTRIBUTION
OF HEAD G UPS
473 DODECYL/OXYETHYLENE/27.1 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
170 DODECYL /OXYETHYLENE/ 28 ALCOHOL REDUCED DISTRIBUTION
OF HEAD GROUPS
116 DODECYL /OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSION
OF HEAD GROUPS
474 DODECYL/OXYETHYLENE/38.6 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
475 DODECYL/OXYETHYLENE/67.3 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
518 TRIMETHYLNONYL/OXYETHYLENE/6 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
158 DODECYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
159 DODECYL BENZENE /OXYETHYLENE/ 10.2 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
160 DODECYL BENZENE /OXYETHYLENE/ 15.1 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
161 DODECYL BENZENE /OXYETHYLENE/ 20.1 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
162 DODECYL BENZENE /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION

148 TRIDECYL /OXYETHYLENE/ 5.2 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
431 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/8.5 ALCOHOL
BRANCHED CHAIN, NATURAL OE DISTRIBUTION
465 TRIDECYL/OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF
HEAD GROUPS
466 TRIDECYL/SECONDARY/ /OXYETHYLENE/9 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
313 TRIDECYL/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
521 TRIDECYL/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
432 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
149 TRIDECYL /OXYETHYLENE/ 10.1 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
467 TRIDECYL/SECONDARY/ /OXYETHYLENE/12 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
314 TRIDECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
433 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/15 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
150 TRIDECYL /OXYETHYLENE/ 15.2 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
315 TRIDECYL/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
151 TRIDECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
434 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/22 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
316 TRIDECYL/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
152 TRIDECYL /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION

C-14

122 TETRADECYL N BETAINE
253 TETRADECYL DIMETHYL AMINE OXIDE
289 TETRADECYL/OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP

C-15

533 HEXADECANOIC ACID
498 SUCROSE MONOPALMITATE

C-16

123 HEXADECYL N BETAINE
589 HEXADECYL DIMETHYLAMMONIOPROPANE SULFONATE
499 SUCROSE DI-PALMITATE
282 HEXADECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD
GROUP
330 HEXADECYL/OXYETHYLENE/7 ALCOHOL HOMOGENEOUS HEAD GROUP
331 HEXADECYL /OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP
332 HEXADECYL/OXYETHYLENE/12 ALCOHOL HOMOGENEOUS HEAD
GROUP
524 HEXADECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
333 HEXADECYL/OXYETHYLENE/15 ALCOHOL HOMOGENEOUS HEAD
GROUP
535 HEXADECYL/OXYETHYLENE/18 ALCOHOL
334 HEXADECYL /OXYETHYLENE/21 ALCOHOL HOMOGENEOUS HEAD
GROUP
117 HEXADECYL /OXYETHYLENE/30 ALCOHOL REDUCED
POLYDISPERSION OF HEAD GROUPS

C-17

534 OCTADECANOIC ACID
496 SUCROSE MONOSTEARATE
553 XYLYL SULFOSTEARIC ACID

C-18

435 OCTADECYL/OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
328 OCTADECYL/OXYETHYLENE/14 ALCOHOL REDUCED
POLYDISPERSITY OF HEAD GROUPS
436 OCTADECYL/OXYETHYLENE/20 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
329 OCTADECYL/OXYETHYLENE/100 ALCOHOL REDUCED
POLYDISPERSITY OF HEAD GROUPS
437 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/12 ALCOHOL
NATURAL DISTRIBUTION OF HEAD GROUPS
438 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/16 ALCOHOL
NATURAL DISTRIBUTION OF HEAD GROUPS
439 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/20 ALCOHOL
NATURAL DISTRIBUTION OF HEAD GROUP

C-13

532 TETRADECANOIC ACID
497 SUCROSE MONOMYRISTATE
312 TRIDECYL/OXYETHYLENE/5 ALCOHOL BRANCHED CHAIN, REDUCED
OE DISTRIBUTION

Structural Indexes of Compounds with Keys to Compound Numbers—Continued

PART 2. Surfactants grouped by charge and by head group and arranged by number of carbon atoms in longest hydrophobic tail.

2A. ANIONICS BY HEAD GROUP

CARBOXYLIC ACID

- 428 PERFLUORO ACETIC ACID
- 429 PERFLUORO PROPIONIC ACID
- 43 BUTYRIC ACID
- 430 PERFLUORO BUTYRIC ACID
- 452 3-4-DICHLOROPERFLUORO BUTYRIC/KEL-F/ACID
- 700 HEXANOIC ACID
- 416 PERFLUORO HEXANOIC ACID
- 453 3-5-6-TRICHLOROPERFLUORO HEXANOIC/KEL-F/ACID
- 374 DODECAFLUOROHEPTANOIC ACID H/CF₂/6COOH
- 529 OCTANOIC ACID
- 417 PERFLUORO OCTANOIC ACID
- 454 3-5-7-8-TETRACHLOROPERFLUORO OCTANOIC/KEL-F/ACID
- 375 HEXADECAFLUORONONANOIC ACID H/CF₂/8COOH
- 530 DECANOIC ACID
- 455 PERFLUORO DECANOIC ACID
- 531 DODECANOIC ACID
- 532 TETRADECANOIC ACID
- 533 HEXADECANOIC ACID
- 534 OCTADECANOIC ACID

MONOCARBOXYLATE

- 699 SODIUM BUTYRATE
- 484 SODIUM PENTANOATE/VALERATE/
- 485 SODIUM HEXANOATE/CAPROATE/
- 188 POTASSIUM HEXANOATE
- 701 POTASSIUM PERFLUOROHEXANOATE
- 486 SODIUM HEPTANOATE
- 296 POTASSIUM HEPTANOATE
- 355 AMMONIUM DODECAFLUOROHEPTANOATE H/CF₂/6COONH₄
- 476 SODIUM OCTANOATE
- 44 POTASSIUM OCTANOATE
- 456 POTASSIUM PERFLUORO OCTANOATE
- 303 DI-ISOPROPYLAMMONIUM CAPRYLATE
- 284 HEXANOLAMINE-CH₃CH/OH/CH₂C/CH₃/2NH₂-OCTANOATE
- 487 SODIUM NONANOATE
- 350 POTASSIUM NONANOATE
- 372 AMMONIUM HEXADECAFLUORONONANOATE H/CF₂/8 COO NH₄
- 299 SODIUM DECANOATE
- 90 POTASSIUM DECANOATE
- 702 POTASSIUM PERFLUORODECANOATE
- 297 POTASSIUM UNDECANOATE
- 527 SODIUM UNDECANE-3-CARBOXYLATE
- 373 AMMONIUM EICOSAFLUOROUNDECANOATE H/CF₂/10 COO NH₄
- 273 SODIUM DODECANOATE
- 91 POTASSIUM DODECANOATE
- 627 CESIUM DODECANOATE
- 277 BENZYL TRIMETHYL AMMONIUM DODECANOATE
- 351 POTASSIUM TRIDECANOATE
- 298 SODIUM TETRADECANOATE
- 92 POTASSIUM TETRADECANOATE
- 300 SODIUM HEXADECANOATE
- 185 POTASSIUM HEXADECANOATE
- 448 SODIUM OCTADECANOATE /STEARATE/
- 256 POTASSIUM STEARATE
- 263 SODIUM OLEATE /CIS-9-OCTADECENOATE/
- 305 POTASSIUM OLEATE/ CIS-9-OCTADECENOATE/
- 285 HEXANOLAMINE-CH₃CH/OH/CH₂C/CH₃/2NH₂-OLEATE
- 264 SODIUM ELAIDATE /TRANS-9-OCTADECENOATE/
- 629 POTASSIUM ELAIDATE/TRANS-9-OCTADECENOATE/
- 283 HEXANOLAMINE-CH₃CH/OH/CH₂C/CH₃/2NH₂-ELAIDATE
- 255 POTASSIUM 9,10 DIHYDROXY STEARATE
- 630 POTASSIUM RICINOLEATE/12 HYDROXY OLEATE/
- 631 POTASSIUM RICINELAIDATE/12 HYDROXY ELAIDATE/

POLYCARBOXYLATE

- 668 DIPOTASSIUM OCTYL MALONATE
- 669 DIPOTASSIUM DECYL MALONATE
- 670 DIPOTASSIUM DODECYL MALONATE
- 671 DIPOTASSIUM TETRADECYL MALONATE
- 672 DIPOTASSIUM HEXADECYL MALONATE
- 673 DIPOTASSIUM OCTADECYL MALONATE
- 420 POTASSIUM 1-1-2-OCTANE TRICARBOXYLATE
- 425 POTASSIUM 1-1-2-DECANE TRICARBOXYLATE

- 421 POTASSIUM 1-1-2-DODECANE TRICARBOXYLATE
- 426 POTASSIUM 1-1-2-TETRADECANE TRICARBOXYLATE
- 422 POTASSIUM 1-1-2-HEXADECANE TRICARBOXYLATE

SULFO CARBOXYLIC ACID

- 6 SODIUM ALPHA SULFOPELARGONIC ACID
- 602 ALPHA SULFO LAURIC ACID
- 235 SODIUM ALPHA SULFO LAURIC ACID
- 189 ALPHA SULFOMYRISTIC ACID
- 236 SODIUM ALPHA SULFO MYRISTIC ACID
- 190 ALPHA SULFOPALMITIC ACID
- 237 SODIUM ALPHA SULFO PALMITIC ACID
- 191 ALPHA SULFOSTEARIC ACID
- 551 PHENYL SULFOSTEARIC ACID
- 552 TOLYL SULFOSTEARIC ACID
- 553 XYLYL SULFOSTEARIC ACID
- 9 SODIUM ALPHA SULFOSTEARIC ACID
- 11 SODIUM ALPHA SULFO PHENYL STEARIC ACID
- 12 SODIUM 9,10 DICHLORO ALPHA SULFOSTEARIC ACID

SULFO CARBOXYLATE

- 233 DISODIUM ALPHA SULFO MYRISTATE
- 234 DISODIUM ALPHA SULFO PALMITATE
- 197 DISODIUM 2 SULFOETHYL ALPHA SULFOPALMITATE
- 8 DISODIUM ALPHA SULFOPHENYLSTEARATE
- 10 DISODIUM ALPHA SULFOSTEARATE
- 198 DISODIUM 2 SULFOETHYL ALPHA SULFOSTEARATE
- 13 DISODIUM 9,10 DICHLORO ALPHA SULFOSTEARATE
- 14 DISODIUM 9,10 DIHYDROXY ALPHA SULFOSTEARATE

PHOSPHONO CARBOXYLIC ACID

- 611 ALPHAPHOSPHONO PELARGONIC ACID
- 612 ALPHAPHOSPHONO DECANOIC ACID
- 613 ALPHAPHOSPHONO DODECANOIC ACID
- 614 ALPHAPHOSPHONO TETRADECANOIC ACID
- 615 ALPHAPHOSPHONO HEXADECANOIC ACID

PHOSPHONO CARBOXYLATE

- 616 MONOSODIUM ALPHAPHOSPHONO DECANOATE
- 617 MONOSODIUM ALPHAPHOSPHONO DODECANOATE
- 618 MONOSODIUM ALPHAPHOSPHONO TETRADECANOATE
- 619 MONOSODIUM ALPHAPHOSPHONO HEXADECANOATE
- 620 DISODIUM ALPHAPHOSPHONO DODECANOATE
- 621 DISODIUM ALPHAPHOSPHONO TETRADECANOATE
- 622 DISODIUM ALPHAPHOSPHONO HEXADECANOATE
- 623 DISODIUM ALPHAPHOSPHONO OCTADECANOATE
- 624 TRISODIUM ALPHAPHOSPHONO TETRADECANOATE
- 625 TRISODIUM ALPHAPHOSPHONO HEXADECANOATE
- 626 TRISODIUM ALPHAPHOSPHONO OCTADECANOATE

SULFO CARBOXYLIC ESTER

- 25 SODIUM ETHYL ALPHA SULFOPELARGONATE
- 26 SODIUM AMYL ALPHA SULFOPELARGONATE
- 27 SODIUM HEXYL ALPHA SULFOPELARGONATE
- 28 SODIUM HEPTYL ALPHA SULFOPELARGONATE
- 35 SODIUM H/CF₂CF₂/3CH₂ ALPHA SULFOPELARGONATE
- 7 SODIUM OCTYL ALPHA SULFOPELARGONATE
- 31 SODIUM 2 OCTYL ALPHA SULFOPELARGONATE
- 32 SODIUM 2 ETHYLHEXYL ALPHA SULFOPELARGONATE
- 29 SODIUM NONYL ALPHA SULFOPELARGONATE
- 30 SODIUM DECYL ALPHA SULFOPELARGONATE
- 33 SODIUM /OXO/ DECYL ALPHA SULFOPELARGONATE
- 34 SODIUM DODECYL ALPHA SULFOPELARGONATE
- 603 SODIUM PROPYL ALPHA SULFO LAURATE
- 604 SODIUM METHYL ALPHA SULFO MYRISTATE
- 36 SODIUM METHYL ALPHA SULFOPALMITATE
- 192 SODIUM ETHYL ALPHA SULFOPALMITATE
- 193 SODIUM PROPYL ALPHA SULFOPALMITATE
- 194 SODIUM METHYL ALPHA SULFOSTEARATE
- 195 SODIUM ETHYL ALPHA SULFOSTEARATE
- 196 SODIUM PROPYL ALPHA SULFOSTEARATE
- 199 SODIUM ISOPROPYL ALPHA SULFOSTEARATE

PHOSPHONO ESTER

- 605 SODIUM AMYL ALPHAPHOSPHONO PELARGONATE

PART 2. Surfactants grouped by charge and by head group and arranged by number of carbon atoms in longest hydrophobic tail—Continued

606 SODIUM AMYL ALPHAPHOSPHONO CAPRATE
 607 SODIUM ISOPROPYL ALPHAPHOSPHONO LAURATE
 608 SODIUM METHYL ALPHAPHOSPHONO MYRISTATE
 610 SODIUM METHYL ALPHAPHOSPHONO STEARATE
 609 SODIUM ISOPROPYL ALPHAPHOSPHONO PALMITATE

62 SODIUM ELAIDYL/TRANS 9 OCTADECENOYL/SULFATE
 63 SODIUM 9 10 DICHLORO OCTADECYL SULFATE
 71 SODIUM OCTADECYL 2 SULFATE
 76 SODIUM OCTADECYL 4 SULFATE
 82 SODIUM OCTADECYL 6 SULFATE
 88 SODIUM 1 NYNOL DECYL SULFATE
 89 SODIUM 1 TETRADECYL PENTADECYL SULFATE

SULFATE

2 SODIUM OCTYL 1 SULFATE
 347 OCTYL TRIMETHYLLAMMONIUM OCTYL SULFATE
 643 DODECYL TRIMETHYLLAMMONIUM OCTANE SULFATE
 66 SODIUM OCTYL 2 SULFATE
 295 SODIUM NYNOL 1-SULFATE
 3 SODIUM DECYL 1 SULFATE
 642 OCTYL TRIMETHYLLAMMONIUM DECANE SULFATE
 79 SODIUM NONADECYL 5 SULFATE
 346 DECYL TRIMETHYLLAMMONIUM DECYL SULFATE
 15 SODIUM DECYL 2 SULFATE
 311 SODIUM UNDECYL 1-SULFATE
 72 SODIUM UNDECYL 3 SULFATE
 80 SODIUM UNDECYL 6 SULFATE

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705 DODECYL SULFURIC ACID
 1 SODIUM DODECYL 1 SULFATE
 634 POTASSIUM DODECYL SULFATE
 111 LITHIUM DODECYL 1 SULFATE
 23 SILVER DODECYL 1 SULFATE
 568 MAGNESIUM DODECYL SULFATE
 24 CALCIUM DODECYL 1 SULFATE
 569 STRONTIUM DODECYL SULFATE
 575 NICKEL DODECYL SULFATE
 572 COBALTOUS DODECYL SULFATE
 574 ZINC DODECYL SULFATE
 570 LEAD DODECYL SULFATE
 571 MANGANESE DODECYL SULFATE
 573 CUPRIC DODECYL SULFATE
 386 AMMONIUM DODECYL SULFATE
 387 METHYLLAMMONIUM DODECYL SULFATE
 388 ETHYLLAMMONIUM DODECYL SULFATE
 389 BUTYLLAMMONIUM DODECYL SULFATE
 112 TETRAMETHYL AMMONIUM DODECYL 1 SULFATE
 382 ETHYL TRIMETHYLLAMMONIUM DODECYL SULFATE
 383 BUTYL TRIMETHYLLAMMONIUM DODECYL SULFATE
 718 TETRAETHYLLAMMONIUM DODECYL SULFATE
 719 TETRABUTYLLAMMONIUM DODECYL SULFATE
 720 1-6-DITRIMETHYLLAMMONIUM-HEXANE/DODECYL SULFATE/2
 409 TRIETHANOLAMMONIUM DODECYL SULFATE
 410 MORPHOLINIUM DODECYL SULFATE
 391 OCTYLAMMONIUM DODECYL SULFATE
 385 OCTYL TRIMETHYLLAMMONIUM DODECYL SULFATE
 280 DECYL TRIMETHYL AMMONIUM DODECYL SULFATE
 281 DODECYL TRIMETHYLLAMMONIUM DODECYL SULFATE
 67 SODIUM DODECYL 2 SULFATE
 562 SODIUM DODECENYL SULFATE
 68 SODIUM TRIDECYL 2 SULFATE
 83 SODIUM TRIDECYL 7 SULFATE
 4 SODIUM TETRADECYL 1 SULFATE
 526 SODIUM 2-DI-N-HEXYL ETHYL SULFATE
 637 LITHIUM TETRADECYL SULFATE
 576 CUPRIC TETRADECYL SULFATE
 16 SODIUM TETRADECYL 2 SULFATE
 73 SODIUM TETRADECYL 3 SULFATE
 17 SODIUM TETRADECYL 4 SULFATE
 77 SODIUM TETRADECYL 5 SULFATE
 525 SODIUM TETRADECYL 6-SULFATE
 84 SODIUM TETRADECYL 7 SULFATE
 69 SODIUM PENTADECYL 2 SULFATE
 74 SODIUM PENTADECYL 3 SULFATE
 78 SODIUM PENTADECYL 5 SULFATE
 85 SODIUM PENTADECYL 8 SULFATE
 5 SODIUM HEXADECYL 1 SULFATE
 638 LITHIUM HEXADECYL SULFATE
 577 CUPRIC HEXADECYL SULFATE
 60 TRIETHANOL AMMONIUM HEXADECYL SULFATE
 75 SODIUM HEXADECYL 4 SULFATE
 81 SODIUM HEXADECYL 6 SULFATE
 86 SODIUM HEXADECYL 8 SULFATE
 70 SODIUM HEPTADECYL 2 SULFATE
 87 SODIUM HEPTADECYL 9 SULFATE

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64 SODIUM OCTADECYL 1 SULFATE
 65 TRIETHANOLAMMONIUM OCTADECYL 1 SULFATE
 61 SODIUM OLEYL/CIS 9 OCTADECENOYL/ SULFATE

POLYOL SULFATE

462 SODIUM MONOLAURIN SULFATE

THIOSULFATE

639 SODIUM UNDECYL THIOSULFATE
 636 SODIUM DODECYL THIOSULFATE

ALKYL SULFONATE

343 SODIUM PENTANE SULFONATE
 344 SODIUM HEXANE SULFONATE
 339 MAGNESIUM HEXANE SULFONATE
 181 SODIUM OCTYL 1-SULFONATE
 340 MAGNESIUM OCTANE SULFONATE
 287 OCTYL TRIMETHYLLAMMONIUM OCTANE SULFONATE
 536 NYNOL SULFONIC ACID
 288 DECYL TRIMETHYLLAMMONIUM DECANESULFONATE
 182 SODIUM DECYL 1-SULFONATE
 341 MAGNESIUM DECANE SULFONATE
 353 OCTYL TRIMETHYLLAMMONIUM DECANE SULFONATE
 537 UNDECYL SULFONIC ACID
 418 SODIUM UNDECYL SULFONATE
 200 DODECYL SULFONIC ACID
 179 SODIUM DODECANE 1-SULFONATE
 40 POTASSIUM DODECYL 1 SULFONATE
 635 LITHIUM DODECYL SULFONATE
 342 MAGNESIUM DODECANE SULFONATE
 175 SODIUM DODECANE 2-SULFONATE
 242 TRIDECANE 1-SULFONIC ACID
 229 SODIUM TRIDECANE 1-SULFONATE
 243 TETRADECANE 1-SULFONIC ACID
 183 SODIUM TETRADECYL 1-SULFONATE
 176 SODIUM TETRADECANE 2-SULFONATE
 244 PENTADECANE 1-SULFONIC ACID
 230 SODIUM PENTADECANE 1-SULFONATE
 245 HEXADECANE 1-SULFONIC ACID
 184 SODIUM HEXADECYL 1-SULFONATE
 408 POTASSIUM HEXADECANE 1-SULFONATE
 177 SODIUM HEXADECANE 2-SULFONATE
 246 HEPTADECANE 1-SULFONIC ACID
 231 SODIUM HEPTADECANE 1-SULFONATE
 232 OCTADECANE 1-SULFONIC ACID
 349 SODIUM OCTADECANE 1-SULFONATE
 419 SODIUM ALPHA-HEPTYL UNDECYL SULFONATE
 178 SODIUM OCTADECANE 2-SULFONATE

HYDROXY ALKYL SULFONATE

247 DODECANE 1-HYDROXY 2-SULFONIC ACID
 238 SODIUM DODECANE 1-HYDROXY 2-SULFONATE
 248 TETRADECANE 1-HYDROXY 2-SULFONIC ACID
 239 SODIUM TETRADECANE 1-HYDROXY 2-SULFONATE
 249 HEXADECANE 1-HYDROXY 2-SULFONIC ACID
 240 SODIUM HEXADECANE 1-HYDROXY 2-SULFONATE
 250 OCTADECANE 1-HYDROXY 2-SULFONIC ACID
 241 SODIUM OCTADECANE 1-HYDROXY 2-SULFONATE

SULFOSUCCINATE DIESTER

257 SODIUM DI-N-BUTYL SULFOSUCCINATE
 261 SODIUM DI-ISOBUTYL SULFOSUCCINATE
 258 SODIUM DI-N-AMYL SULFOSUCCINATE
 259 SODIUM DI-N-HEXYL SULFOSUCCINATE
 352 SODIUM DI-1-METHYLISOAMYL SULFOSUCCINATE
 260 SODIUM DI-N-OCTYL SULFOSUCCINATE
 262 SODIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE
 286 AMMONIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE

PARA ALKARYL SULFONATE

445 SODIUM PARA-BIS-/N-BUTYL/ BENZENE SULFONATE
 446 SODIUM PARA-BIS-/N-HEXYL/ BENZENE SULFONATE
 49 SODIUM P OCTYL BENZENE SULFONATE
 447 SODIUM PARA-BIS-/N-OCTYL/ BENZENE SULFONATE
 493 SODIUM P-NONYL BENZENE SULFONATE
 50 SODIUM P DECYL BENZENE SULFONATE
 45 SODIUM P 1 METHYL DECYL BENZENE SULFONATE
 51 SODIUM P DODECYL BENZENE SULFONATE
 46 SODIUM P 1 METHYL DODECYL BENZENE SULFONATE

PART 2. Surfactants grouped by charge and by head group and arranged by number of carbon atoms in longest hydrophobic tail—Continued

47	SODIUM P 1 METHYL TETRADECYL BENZENE SULFONATE	113	SODIUM DODECYL TRI-OXYETHYLENE SULFATE
48	SODIUM P 1 METHYL HEXADECYL BENZENE SULFONATE	546	SODIUM TETRADECYL TRI-OXYETHYLENE SULFATE
ALKARYL SULFONATE (UNSPECIFIED)			
682	SODIUM DIBUTYL NAPHTHALENE SULFONATE /NEKAL/	54	SODIUM HEXADECYL TRI OXYETHYLENE SULFATE
681	SODIUM DIBUTYL BENZENE SULFONATE	58	SODIUM OCTADECYL TRI OXYETHYLENE SULFATE
501	SODIUM HEXYL BENZENE SULFONATE	549	SODIUM OLEYL TRI-OXYETHYLENE SULFATE
502	SODIUM HEPTYL BENZENE SULFONATE	543	SODIUM DODECYL TETRA-OXYETHYLENE SULFATE
C-8		55	SODIUM HEXADECYL TETRA OXYETHYLENE SULFATE
503	SODIUM OCTYL BENZENE SULFONATE	59	SODIUM OCTADECYL TETRA OXYETHYLENE SULFATE
510	SODIUM 2-ETHYL-HEXYL BENZENE SULFONATE	114	SODIUM DODECYL /OXYETHYLENE/17.5 SULFATE
676	SODIUM 1,1,3,3-TETRAMETHYL BUTYL BENZENE SULFONATE		
677	SODIUM DI-/1,1,3,3-TETRAMETHYL BUTYL/BENZENESULFONATE		
172	SODIUM 2-N-OCTYL BENZENE SULFONATE		
C-9			
504	SODIUM NONYL BENZENE SULFONATE	OTHER	
138	SODIUM NONYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN	136	SODIUM ALPHA DIMETHYL AMINO CAPRATE
272	TRI-ISOPROPYL BENZENE SULFONIC ACID	704	POTASSIUM 4-HEXYL RESORCINOLATE
271	SODIUM TRI-ISO-PROPYL BENZENE SULFONATE	632	POTASSIUM N-DODECYL BETA-ALANINATE
C-10			
505	SODIUM DECYL BENZENE SULFONATE	2B. CATIONICS BY HEAD GROUP	
173	SODIUM 2-N-DECYL BENZENE SULFONATE		
511	SODIUM 2-PROPYL-HEPTYL BENZENE SULFONATE	AMINE	
678	SODIUM 1,1-DIMETHYLNONYL BENZENE SULFONATE	708	HEXYLAMINE
679	SODIUM 3-BUTYL HEPTYL BENZENE SULFONATE	706	PERFLUORO PROPYLAMINE
140	SODIUM DECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN		
561	SANTOMERSE D /SODIUM DECYLBENZENE SULFONATE/	R-H3	
C-12		707	PERFLUORO PROPYLAMINE HYDROCHLORIDE
680	SODIUM 2-PROPYL-4-METHYL-OCTYL BENZENE SULFONATE	709	HEXYLAMINE HYDROCHLORIDE
492	SODIUM TETRAPROPYLENE/1-3-5-7-TETRAMETHYL-OCTYL/BENZENE SULFONATE	390	HEXYLAMMONIUM DODECYL SULFATE
506	SODIUM DODECYL BENZENE SULFONATE	392	OCTYLYLAMMONIUM CHLORIDE
139	SODIUM DODECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN	391	OCTYLYLAMMONIUM DODECYL SULFATE
171	SODIUM 2-N-DODECYL BENZENE SULFONATE	37	DECYLYLAMMONIUM CHLORIDE
301	SODIUM 3-N-DODECYL BENZENE SULFONATE	411	DECYLYLAMMONIUM ACETATE
302	SODIUM 4-N-DODECYL BENZENE SULFONATE	38	DODECYL AMMONIUM CHLORIDE
514	SODIUM 6-N-DODECYL BENZENE SULFONATE	628	DODECYLAMMONIUM BROMIDE
512	SODIUM 2-BUTYL-OCTYL BENZENE SULFONATE	482	DODECYL AMMONIUM NITRATE
141	SODIUM TRIDECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN	412	DODECYLAMMONIUM ACETATE
554	SANTOMERSE 3 /SODIUM DODECYL BENZENE SULFONATE/	39	TETRADECYL AMMONIUM CHLORIDE
C-14		413	TETRADECYLAMMONIUM ACETATE
507	SODIUM TETRADECYL BENZENE SULFONATE	186	HEXADECYL AMMONIUM CHLORIDE
174	SODIUM 2-N-TETRADECYL BENZENE SULFONATE	414	HEXADECYLAMMONIUM ACETATE
513	SODIUM 2-AMYL-NONYL BENZENE SULFONATE	187	OCTADECYL AMMONIUM CHLORIDE
596	TETRAHYDROANACARDOL AMMONIUM MONOSULFONATE	415	OCTADECYLAMMONIUM ACETATE
142	SODIUM PENTADECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN		
508	SODIUM HEXADECYL BENZENE SULFONATE	RR-H2	
509	SODIUM OCTADECYL BENZENE SULFONATE	449	DODECYLMETHYL AMMONIUM CHLORIDE
683	SODIUM EICOSYLBENZENE SULFONATE		
557	ARESKET 300 /MONOBUTYL BIPHENYL SODIUM MONOSULFONATE/	RRR-H	
556	ARESKAP 100 /MONOBUTYL PHENYLPHENOL SODIUM MONOSULFONATE/	450	DODECYLDIMETHYL AMMONIUM CHLORIDE
558	ARESKLENE 400 /DIBUTYL PHENYLPHENOL DISODIUMDISULFONATE		
OXYPROPYL SULFATE		R-(CH₃)₃	
597	SODIUM DODECYL MONO-OXYPROPYL SULFATE	644	HEXYL TRIMETHYLAMMONIUM HEXANE SULFONATE
598	SODIUM TETRADECYL MONO-OXYPROPYL SULFATE	640	HEXYL TRIMETHYLAMMONIUM HEXANE SULFATE
600	SODIUM HEXADECYL MONO-OXYPROPYL SULFATE	641	HEXYL TRIMETHYLAMMONIUM OCTANE SULFATE
601	SODIUM OCTADECYL MONO-OXYPROPYL SULFATE	93	OCTYL TRIMETHYL AMMONIUM BROMIDE
599	SODIUM TETRADECYL DI-OXYPROPYL SULFATE		
OXYETHYLENE SULFATE		C-8	
541	SODIUM DODECYL MONO-OXYETHYLENE SULFATE	347	OCTYL TRIMETHYLAMMONIUM OCTYL SULFATE
544	SODIUM TETRADECYL MONO-OXYETHYLENE SULFATE	287	OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE
52	SODIUM HEXADECYL MONO OXYETHYLENE SULFATE	642	OCTYL TRIMETHYLAMMONIUM DECANE SULFATE
56	SODIUM OCTADECYL MONO OXYETHYLENE SULFATE	353	OCTYL TRIMETHYLAMMONIUM DECANE SULFONATE
547	SODIUM OLEYL MONO-OXYETHYLENE SULFATE	385	OCTYL TRIMETHYLAMMONIUM DODECYL SULFATE
542	SODIUM DODECYL DIOXYETHYLENE SULFATE		
545	SODIUM TETRADECYL DI-OXYETHYLENE SULFATE		
53	SODIUM HEXADECYL DI OXYETHYLENE SULFATE		
57	SODIUM OCTADECYL DI OXYETHYLENE SULFATE		
548	SODIUM OLEYL DI-OXYETHYLENE SULFATE		
C-9		C-9	
		94	NONYL TRIMETHYL AMMONIUM BROMIDE
C-10		C-10	
203	DECYL TRIMETHYL AMMONIUM CHLORIDE	203	DECYL TRIMETHYL AMMONIUM CHLORIDE
95	DECYL TRIMETHYL AMMONIUM BROMIDE	95	DECYL TRIMETHYL AMMONIUM BROMIDE
306	DECYL TRIMETHYLAMMONIUM SULFATE	306	DECYL TRIMETHYLAMMONIUM SULFATE
346	DECYL TRIMETHYLAMMONIUM DECYL SULFATE	346	DECYL TRIMETHYLAMMONIUM DECYL SULFATE
280	DECYL TRIMETHYL AMMONIUM DODECYL SULFATE	280	DECYL TRIMETHYL AMMONIUM DODECYL SULFATE
288	DECYL TRIMETHYLAMMONIUM DECANESULFONATE	288	DECYL TRIMETHYLAMMONIUM DECANESULFONATE
C-11		C-11	
		96	UNDECYL TRIMETHYL AMMONIUM BROMIDE
C-12		C-12	
41	DODECYL TRIMETHYL AMMONIUM CHLORIDE	41	DODECYL TRIMETHYL AMMONIUM CHLORIDE
126	DODECYL TRIMETHYLAMMONIUM IODIDE	126	DODECYL TRIMETHYLAMMONIUM IODIDE
97	DODECYL TRIMETHYL AMMONIUM BROMIDE	97	DODECYL TRIMETHYL AMMONIUM BROMIDE
130	DODECYL TRIMETHYL AMMONIUM FLUORIDE	130	DODECYL TRIMETHYL AMMONIUM FLUORIDE
131	DODECYL TRIMETHYLAMMONIUM NITRATE	131	DODECYL TRIMETHYLAMMONIUM NITRATE
127	DODECYL TRIMETHYL AMMONIUM IODATE	127	DODECYL TRIMETHYL AMMONIUM IODATE
129	DODECYL TRIMETHYL AMMONIUM BROMATE	129	DODECYL TRIMETHYL AMMONIUM BROMATE

PART 2. Surfactants grouped by charge and by head group and arranged by number of carbon atoms in longest hydrophobic tail—Continued

307 DODECYL TRIMETHYLAMMONIUM SULFATE
 128 DODECYL TRIMETHYL AMMONIUM FORMATE
 643 DODECYL TRIMETHYLAMMONIUM OCTANE SULFATE
 281 DODECYL TRIMETHYLAMMONIUM DODECYL SULFATE
 384 HEXYL TRIMETHYLAMMONIUM DODECYL SULFATE

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402 TRIDECYL TRIMETHYLAMMONIUM CHLORIDE

C-14

42 TETRADECYL TRIMETHYL AMMONIUM CHLORIDE
 98 TETRADECYL TRIMETHYL AMMONIUM BROMIDE
 308 TETRADECYL TRIMETHYLAMMONIUM SULFATE

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265 HEXADECYL TRIMETHYL AMMONIUM CHLORIDE
 99 HEXADECYL TRIMETHYLAMMONIUM BROMIDE
 309 HEXADECYL TRIMETHYLAMMONIUM SULFATE
 559 CATOL 605 / (N-(2-KETO-2-(2,-LAUROYL
 OXYETHYLAMINO))ETHYL TRIMETHYLAMMONIUM CHLORIDE/

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270 OCTADECYL TRIMETHYLAMMONIUM CHLORIDE
 477 OCTADECYL TRIMETHYLAMMONIUM BROMIDE
 654 OCTADECYL TRIMETHYLAMMONIUM NITRATE
 658 OCTADECYL TRIMETHYLAMMONIUM BROMATE
 659 OCTADECYL TRIMETHYLAMMONIUM FORMATE
 667 OCTADECYL TRIMETHYLAMMONIUM OXALATE

RR-(CH₃)₂

399 DODECYL DIMETHYL ETHYLAMMONIUM CHLORIDE
 403 DODECYL DIMETHYLPHENYLAMMONIUM CHLORIDE
 478 HEXADECYL DIMETHYLETHYLAMMONIUM BROMIDE
 266 HEXADECYL DIMETHYL 2-HYDROXYETHYL AMMONIUM CHLORIDE
 269 HEXADECYLDIMETHYL 2,3-DIHYDROXYPROPYLAMMONIUM CHLORIDE
 483 DIOCTYL DIMETHYL AMMONIUM CHLORIDE
 345 DIDODECYL DIMETHYLAMMONIUM CHLORIDE

BENZYL

354 HEXYL BENZYL DIMETHYLAMMONIUM CHLORIDE
 356 DECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
 279 DODECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
 357 TETRADECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
 275 HEXADECYL DIMETHYLBENZYLAMMONIUM CHLORIDE
 358 OCTADECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
 406 DODECYL DIMETHYL META-TRIFLUOROMETHYLBENZYL AMMONIUM
 CHLORIDE CF₃CH₂CH₂/N/CH₃/2/C12H25
 359 OCTYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
 360 DECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
 361 DODECYL 4-NITROBENZYL DIMETHYL AMMONIUM CHLORIDE
 362 TETRADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
 363 HEXADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
 364 OCTADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
 365 DODECYL 2-CHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
 367 DODECYL 4-CHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
 366 DODECYL 2-4-DICHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
 369 DODECYL 3-4-DICHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
 371 DODECYL 3-4-DIMETHOXYBENZYL DIMETHYLAMMONIUM CHLORIDE
 370 DODECYL 3-4-METHYLENEDIOXYBENZYL DIMETHYLAMMONIUM
 CHLORIDE
 368 DODECYL 2-HYDROXY-5-NITROBENZYL DIMETHYLAMMONIUM
 CHLORIDE
 451 PARA DI-ISOBUTYLPHENOXYETHOXETHYL DIMETHYLBENZYL
 AMMONIUM CHLORIDE/HYAMINE 1622/
 404 DODECYL DIMETHYL 2-PHENYLETHYLAMMONIUM CHLORIDE
 C6H₅CH₂CH₂/N/CH₃/2/C12H25
 407 DODECYL DIMETHYL 3-PHENYLPROPYLAMMONIUM CHLORIDE
 C6H₅CH₂CH₂/N/CH₃/2/C12H25

RRR-CH₃

400 DODECYL METHYL DIETHYLAMMONIUM CHLORIDE
 405 DODECYL METHYLETHYLBENZYLAMMONIUM CHLORIDE
 267 HEXADECYL DI-/2-HYDROXYETHYL/METHYL AMMONIUMCHLORIDE

R-R3

662 OCTADECYL TRIETHYLAMMONIUM BROMATE
 401 DODECYL TRIETHYLAMMONIUM CHLORIDE
 663 OCTADECYL TRIPROPYLAMMONIUM BROMATE
 292 HEXADECYL TRIPROPYLAMMONIUM BROMIDE
 291 TETRADECYL TRIPROPYLAMMONIUM BROMIDE
 664 OCTADECYL TRIBUTYLAMMONIUM BROMATE
 666 HEXADECYL TRIBUTYLAMMONIUM BROMATE
 665 OCTADECYL TRIAMYLAMMONIUM BROMATE

268 HEXADECYL TRI-/2-HYDROXYETHYL/ AMMONIUM CHLORIDE
 500 DODECYL TRI/2-HYDROXYETHYL/AMMONIUM CHLORIDE

RRR-O

293 DODECYL DIMETHYL AMINE OXIDE HYDROBROMIDE
 22 DIMETHYL DODECYL AMINE OXIDE HYDROCHLORIDE

BETAINE

135 OCTYL C BETAINE HYDROCHLORIDE
 124 DODECYL N BETAINE HYDROCHLORIDE
 125 TETRADECYL N BETAINE HYDROCHLORIDE

AMINO ACID

633 N-DODECYL BETA-ALANINE HYDROCHLORIDE
 137 ALPHA DIMETHYLAMINO CAPRIC ACID HYDROCHLORIDE

PYRIDINIUM

100 OCTYL PYRIDINIUM BROMIDE
 458 DECYL PYRIDINIUM IODIDE
 101 UNDECYL PYRIDINIUM BROMIDE
 278 DODECYL PYRIDINIUM CHLORIDE
 290 DODECYL PYRIDINIUM BROMIDE
 376 DODECYL PYRIDINIUM IODIDE
 102 TETRADECYL PYRIDINIUM BROMIDE
 479 TETRADECYL PYRIDINIUM IODIDE
 304 CATOL 607 (N-/LAURYL-COLAMINO-FORMYL-METHYL/-
 PYRIDINIUM CHLORIDE)

C-16

274 HEXADECYL PYRIDINIUM CHLORIDE
 427 HEXADECYL PYRIDINIUM BROMIDE
 480 HEXADECYL PYRIDINIUM IODIDE
 660 HEXADECYL PYRIDINIUM IODATE
 693 N-CETYL 2-METHYL PYRIDINIUM CHLORIDE
 696 N-CETYL 2-METHYL PYRIDINIUM IODIDE
 694 N-CETYL 3-METHYL PYRIDINIUM CHLORIDE
 697 N-CETYL-3-METHYL PYRIDINIUM IODIDE
 695 N-CETYL 4-METHYL PYRIDINIUM CHLORIDE
 698 N-CETYL-4-METHYL PYRIDINIUM IODIDE
 560 EMULSOL 607L (N-(2-KETO-2-(2,-LAUROYLOXETHYL AMINO))
 ETHYL) PYRIDINIUM CHLORIDE
 655 OCTADECYL PYRIDINIUM CHLORIDE
 657 OCTADECYL PYRIDINIUM BROMIDE
 481 OCTADECYL PYRIDINIUM IODIDE
 656 OCTADECYL PYRIDINIUM NITRATE
 661 OCTADECYL PYRIDINIUM IODATE

QUINOLINIUM

717 DODECYLQUINOLINIUM BROMIDE

TROPYLUM

528 DODECYL TROPYLUM PERCHLORATE
 460 DODECYL TROPYLUM MONOPHOSPHATE
 491 DODECYL TROPYLUM BISULFATE

2C. NONIONICS BY HEAD GROUP

CARBOXYLIC ACID

43 BUTYRIC ACID
 700 HEXANOIC ACID
 529 OCTANOIC ACID
 530 DECANOIC ACID
 531 DODECANOIC ACID
 532 TETRADECANOIC ACID
 533 HEXADECANOIC ACID
 534 OCTADECANOIC ACID

AMINE

706 PERFLURO PROPYLAMINE
 708 HEXYLAMINE

N-BETAINE

118 OCTYL N BETAINE
 119 DECYL N BETAINE
 120 UNDECYL N BETAINE
 121 DODECYL N BETAINE
 592 DODECYL N-DIETHYL N-BETAINE
 122 TETRADECYL N BETAINE
 123 HEXADECYL N BETAINE

C-BETAINE

132 OCTYL C BETAINE

PART 2. *Surfactants grouped by charge and by head group and arranged by number of carbon atoms in longest hydrophobic tail—Continued*

133 DECYL C BETAINE
134 DODECYL C BETAINE

AMINE OXIDE

714 HEXYL DIMETHYL AMINE OXIDE
715 HEPTYL DIMETHYL AMINE OXIDE
251 OCTYL DIMETHYL AMINE OXIDE
254 NONYL DIMETHYL AMINE OXIDE
252 DECYL DIMETHYL AMINE OXIDE
713 UNDECYL DIMETHYL AMINE OXIDE
21 DIMETHYL DODECYL AMINE OXIDE
253 TETRADECYL DIMETHYL AMINE OXIDE
348 NN-DIMETHYL 1-1-DIHYDROPENTADECAFLUORO OCTYL AMINE N-OXIDE /C7F15CH2N/CH3/20/

PHOSPHINE OXIDE

710 OCTYL DIMETHYL PHOSPHINE OXIDE
711 DECYL DIMETHYL PHOSPHINE OXIDE
712 DODECYL DIMETHYL PHOSPHINE OXIDE

-ONIO -ATE

590 DODECYL DIMETHYLLAMMONIOPROPANE CARBOXYLATE
587 DECYL DIMETHYLLAMMONIOPROPANE SULFONATE
588 DODECYL DIMETHYLLAMMONIOPROPANE SULFONATE
594 DODECYL DIPROPYL AMMONIOPROPANE SULFONATE
589 HEXADECYL DIMETHYLLAMMONIOPROPANE SULFONATE
595 DODECYL DIMETHYL AMMONIOPROPANE SULFATE
593 DIMETHYL DODECYLLAMMONIOPROPANE HYDROXY SULFONATE
591 DODECYL DIMETHYLPHOSPHONIOPROPANE SULFONATE

SULFINYL OL

684 HEXYL SULFINYLETHANOL
688 OCTYL SULFINYLETHANOL
692 DECYL SULFINYLETHANOL
685 HEXYL SULFINYLPROPANOL
689 OCTYL SULFINYLPROPANOL
686 HEXYL SULFINYLBUTANOL
690 OCTYL SULFINYLBUTANOL
687 HEXYL SULFINYLPENTANOL

SUGAR ALKYLATE

648 ALPHA-D-GLUCOSYL OCTANE
649 ALPHA-D-GLUCOSYL DODECANE
645 PARA/BETA-D-GLUCOSYL/ETHYLBENZENE
646 PARA/BETA-D-GLUCOSYL/PROPYL BENZENE
647 PARA/BETA-D-GLUCOSYL/BUTYLBENZENE
650 PARA/BETA-D-XYLOSYL/BUTYL BENZENE

SUGAR ESTER

495 SUCROSE MONOLAURATE
497 SUCROSE MONOMYRISTATE
498 SUCROSE MONOPALMITATE
499 SUCROSE DI-PALMITATE
496 SUCROSE MONOSTEARATE

POLYOL ETHER

424 OCTYL ALPHA-GLYCERYL ETHER
18 OCTYL BETA D GLUCOSIDE
19 DECYL BETA D GLUCOSIDE
20 DODECYL BETA D GLUCOSIDE

OTHER

457 1-4-HEXANEDIOL
703 4-HEXYL RESORCINOL
691 OCTYL METHYL SULFOXIDE

METHYL-OXYETHYLENE ESTER

583 METHYL /OXYETHYLENE/ 6.0 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
580 METHYL /OXYETHYLENE/ 7.0 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
586 METHYL /OXYETHYLENE/ 7.6 OCTANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
584 METHYL /OXYETHYLENE/ 8.4 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
581 METHYL /OXYETHYLENE/ 10.3 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
585 METHYL /OXYETHYLENE/ 11.2 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
578 METHYL /OXYETHYLENE/ 11.9 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
579 METHYL /OXYETHYLENE/ 12.5 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS

582 METHYL /OXYETHYLENE/ 16.0 DECANOATE NATURAL DISTRIBUTION OF HEAD GROUPS

METHYL-OXYETHYLENE ETHER

201 DECYL /OXYETHYLENE/8 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS
202 DECYL /OXYETHYLENE/11 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS
204 DECYL /OXYETHYLENE/12 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS
205 DODECYL /OXYETHYLENE/12 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS

OXYETHYLENE SORBITAN

440 /OXYETHYLENE/4 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
441 /OXYETHYLENE/8 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
442 /OXYETHYLENE/10 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
443 /OXYETHYLENE/20 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS
444 /OXYETHYLENE/25 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS

OXYETHYLENE DIOL ESTER

276 NONAETHYLENE GLYCOL MONODODECANOATE NATURAL DISTRIBUTION OF HEAD GROUPS

OXYETHYLENE ALCOHOL

OE 1
461 BUTYL/OXYETHYLENE/1 ALCOHOL--BUTYL GLYCOL ETHER HOMOGENEOUS HEAD GROUP
423 OCTYL/OXYETHYLENE/1 ALCOHOL--OCTYL GLYCOL ETHER HOMOGENEOUS HEAD GROUP
207 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL HOMOGENOUS HEAD GROUP
217 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS

OE 2

208 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL HOMOGENOUS HEAD GROUP
218 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS

OE 3

103 HEXYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
104 OCTYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
209 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL HOMOGENOUS HEAD GROUP
219 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
107 DECYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP

OE 3

380 HEXYL/OXYETHYLENE/4 ALCOHOL HOMOGENEOUS HEAD GROUP
515 ISO-OCTYL/OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
210 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL HOMOGENOUS HEAD GROUP
220 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
516 NONYL/OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
379 DECYL/OXYETHYLENE/4 ALCOHOL HOMOGENEOUS HEAD GROUP
519 DECYL/OXO-PROCESS/ /OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
325 DODECYL/OXYETHYLENE/4 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS

OE 5

381 HEXYL/OXYETHYLENE/5 ALCOHOL HOMOGENEOUS HEAD GROUP
211 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL HOMOGENOUS HEAD GROUP
221 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
153 NONYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
378 DECYL/OXYETHYLENE/5 ALCOHOL HOMOGENEOUS HEAD GROUP

PART 2. Surfactants grouped by charge and by head group and arranged by number of carbon atoms in longest hydrophobic tail—Continued

143	DECYL /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	313	TRIDECYL/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
377	DODECYL/OXYETHYLENE/ 5 ALCOHOL HOMOGENEOUS HEAD GROUP	331	HEXADECYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP
158	DODECYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	OE 10	
312	TRIDECYL/OXYETHYLENE/5 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION	675	OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
148	TRIDECYL /OXYETHYLENE/ 5.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	216	P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL HOMOGENOUS HEAD GROUP
OE 6		317	T-OCTYL BENZENE/OXYETHYLENE/10 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS
393	BUTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP	226	P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
394	1-METHYL PROPYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP	320	NONYL BENZENE/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
674	DIISOBUTYL BENZENE/OXYETHYLENE/6 ALCOHOL (DBBRANCHED CHAIN, NATURAL OE DISTRIBUTION	167	NONYL BENZENE /OXYETHYLENE/ 10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
294	HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP	337	NONYL BENZENE/OXYETHYLENE/10.5 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
395	2-ETHYL BUTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP	144	DECYL /OXYETHYLENE/ 10.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
105	OCTYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP	517	DECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
396	2-PROPYL PENTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP	310	DODECYL/OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
212	P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP	470	DODECYL/OXYETHYLENE/10.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
222	P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	159	DODECYL BENZENE /OXYETHYLENE/ 10.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
108	DECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP	521	TRIDECYL/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
397	2-BUTYL HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP	149	TRIDECYL /OXYETHYLENE/ 10.1 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
110	DODECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP	432	TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
468	DODECYL/OXYETHYLENE/6.2 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	435	OCTADECYL /OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
518	TRIMETHYLNONYL/OXYETHYLENE/6 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	OE 7	
289	TETRADECYL /OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP	213	P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL HOMOGENOUS HEAD GROUP
282	HEXADECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENEOUS HEAD GROUP	223	P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
		488	DODECYL/OXYETHYLENE/7 ALCOHOL HOMOGENEOUS HEAD GROUP
		115	DODECYL /OXYETHYLENE/7 ALCOHOL REDUCED POLYDISPERSION OF HEAD GROUPS
		469	DODECYL/OXYETHYLENE/7.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
		330	HEXADECYL /OXYETHYLENE/7 ALCOHOL HOMOGENEOUS HEAD GROUP
OE 8		OE 11	
335	OCTYL BENZENE/OXYETHYLENE/8.5 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	463	NONYL BENZENE/OXYETHYLENE/11 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
214	P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL HOMOGENOUS HEAD GROUP	523	DECYL BENZENE PARA/OXYETHYLENE/11.0 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
224	P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	522	DECYL BENZENE ORTHO/OXYETHYLENE/11.2 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
520	UNDECYL/OXYETHYLENE/8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	OE 12	
163	DODECYL /OXYETHYLENE/ 8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	490	DODECYL/OXYETHYLENE/12 ALCOHOL HOMOGENEOUS HEAD GROUP
431	TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/8.5 ALCOHOL BRANCHED CHAIN. NATURAL OE DISTRIBUTION	164	DODECYL /OXYETHYLENE/ 12 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
OE 9		467	TRIDECYL/SECONDARY/ /OXYETHYLENE/12 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
106	OCTYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP	332	HEXADECYL/OXYETHYLENE/12 ALCOHOL HOMOGENEOUS HEAD GROUP
215	P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL HOMOGENOUS HEAD GROUP	437	OLEYL/CIS-9-OCTADECENOYL /OXYETHYLENE/12 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
225	P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	OE 14	
206	TRITON X-100(P-T-OCTYL BENZENE/OXYETHYLENE/9)NATURAL DISTRIBUTION OF HEAD GROUPS	145	DECYL /OXYETHYLENE/ 14.9 ALCOHOL BRANCHED CHAIN. NATURAL OE DISTRIBUTION
336	NONYL BENZENE/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	326	DODECYL/OXYETHYLENE/14 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS
154	NONYL BENZENE /OXYETHYLENE/ 9.8 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	314	TRIDECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
109	DECYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP	524	HEXADECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
398	2-BUTYL HEXYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP	328	OCTADECYL /OXYETHYLENE/14 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS
489	DODECYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP	OE 15	
465	TRIDECYL/OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	321	NONYL BENZENE/OXYETHYLENE/15 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
466	TRIDECYL/SECONDARY/ /OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	464	NONYL BENZENE/OXYETHYLENE/15 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS

PART 2. *Surfactants grouped by charge and by head group and arranged by number of carbon atoms in longest hydrophobic tail*—Continued

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| <p>433 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/15 ALCOHOL BRANCHED CHAIN. NATURAL OE DISTRIBUTION</p> <p>333 HEXADECYL/OXYETHYLENE/15 ALCOHOL HOMOGENEOUS HEAD GROUP</p> <p>OE 16</p> <p>227 P-T-OCTYL BENZENE /OXYETHYLENE/16 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>716 DODECYL /OXYETHYLENE/16 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS</p> <p>438 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/16 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>OE 18</p> <p>165 DODECYL /OXYETHYLENE/ 18 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>535 HEXADECYL/OXYETHYLENE/18 ALCOHOL</p> <p>OE 20</p> <p>318 T-OCTYL BENZENE/OXYETHYLENE/20 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS</p> <p>322 NONYL BENZENE/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN. REDUCED OE DISTRIBUTION</p> <p>156 NONYL BENZENE /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN. NATURAL OE DISTRIBUTION</p> <p>146 DECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>472 DODECYL/OXYETHYLENE/20.4 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>161 DODECYL BENZENE /OXYETHYLENE/ 20.1 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>315 TRIDECYL/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION</p> <p>151 TRIDECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>436 OCTADECYL/OXYETHYLENE/20 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>439 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/20 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUP</p> <p>OE 21-25</p> <p>334 HEXADECYL/OXYETHYLENE/21 ALCOHOL HOMOGENEOUS HEAD GROUP</p> <p>434 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/22 ALCOHOL BRANCHED CHAIN. NATURAL OE DISTRIBUTION</p> <p>327 DODECYL/OXYETHYLENE/23 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS</p> | <p>166 DODECYL /OXYETHYLENE/ 23 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>OE 26-30</p> <p>473 DODECYL/OXYETHYLENE/27.1 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>170 DODECYL /OXYETHYLENE/ 28 ALCOHOL REDUCED DISTRIBUTION OF HEAD GROUPS</p> <p>147 DECYL /OXYETHYLENE/ 29.4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>319 T-OCTYL BENZENE/OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS</p> <p>323 NONYL BENZENE/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN. REDUCED OE DISTRIBUTION</p> <p>721 NONYL BENZENE /OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN DIALYSED</p> <p>169 NONYL BENZENE /OXYETHYLENE/ 30 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>157 NONYL BENZENE /OXYETHYLENE/ 30.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>116 DODECYL /OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSION OF HEAD GROUPS</p> <p>162 DODECYL BENZENE /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>316 TRIDECYL/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION</p> <p>152 TRIDECYL /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION</p> <p>117 HEXADECYL /OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSION OF HEAD GROUPS</p> <p>OE 31-40</p> <p>474 DODECYL/OXYETHYLENE/38.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>228 P-T-OCTYL BENZENE /OXYETHYLENE/40 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>OE 41+</p> <p>324 NONYL BENZENE/OXYETHYLENE/50 ALCOHOL BRANCHED CHAIN. REDUCED OE DISTRIBUTION</p> <p>475 DODECYL/OXYETHYLENE/67.3 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS</p> <p>338 NONYL BENZENE/OXYETHYLENE/100 ALCOHOL BRANCHED CHAIN. NATURAL OE DISTRIBUTION</p> <p>329 OCTADECYL/OXYETHYLENE/100 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS</p> |
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Structural Indexes of Compounds with Keys to Compound Numbers—Continued

PART 3. *Grouped by structure of hydrophobic part and arranged by number of carbon atoms in longest hydrophobic tail and by charge.*

3A. ALKANES

N-ALKANES

C-3

699	SODIUM BUTYRATE
43	BUTYRIC ACID
707	PERFLUORO PROPYLAMINE HYDROCHLORIDE
706	PERFLUORO PROPYLAMINE

C-4

484	SODIUM PENTANOATE/VALERATE/
257	SODIUM DI-N-BUTYL SULFOSUCCINATE
461	BUTYL/OXYETHYLENE/1 ALCOHOL--BUTYL GLYCOL ETHER HOMOGENEOUS HEAD GROUP
393	BUTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP

C-5

700	HEXANOIC ACID
485	SODIUM HEXANOATE/CAPROATE/
188	POTASSIUM HEXANOATE
258	SODIUM DI-N-AMYL SULFOSUCCINATE
343	SODIUM PENTANE SULFONATE
700	HEXANOIC ACID

C-6

486	SODIUM HEPTANOATE
296	POTASSIUM HEPTANOATE
344	SODIUM HEXANE SULFONATE
339	MAGNESIUM HEXANE SULFONATE
259	SODIUM DI-N-HEXYL SULFOSUCCINATE
704	POTASSIUM 4-HEXYL RESORCINOLATE
708	HEXYLAMINE
709	HEXYLAMINE HYDROCHLORIDE
640	HEXYL TRIMETHYLAMMONIUM HEXANE SULFATE
644	HEXYL TRIMETHYLAMMONIUM HEXANE SULFONATE
641	HEXYL TRIMETHYLAMMONIUM OCTANE SULFATE
708	HEXYLAMINE
390	HEXYLAMMONIUM DODECYL SULFATE
714	HEXYL DIMETHYL AMINE OXIDE
684	HEXYL SULFINYLETHANOL
685	HEXYL SULFINYLPROPANOL
686	HEXYL SULFINYLBUTANOL
687	HEXYL SULFINYLPHENOL
457	1-4-HEXANEDIOL
103	HEXYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
380	HEXYL/OXYETHYLENE/4 ALCOHOL HOMOGENOUS HEAD GROUP
381	HEXYL/OXYETHYLENE/5 ALCOHOL HOMOGENOUS HEAD GROUP
294	HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP
703	4-HEXYL RESORCINOL

C-7

529	OCTANOIC ACID
476	SODIUM OCTANOATE
44	POTASSIUM OCTANOATE
284	HEXANOLAMINE-CH ₃ CH ₂ /OH/CH ₂ C/CH ₃ /2NH ₂ -OCTANOATE
303	DI-ISOPROPYLAMMONIUM CAPRYLATE
529	OCTANOIC ACID
715	HEPTYL DIMETHYL AMINE OXIDE
586	METHYL /OXYETHYLENE/ 7.6 OCTANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS

C-8

487	SODIUM NONANOATE
350	POTASSIUM NONANOATE
420	POTASSIUM 1-1-2-OCTANE TRICARBOXYLATE
6	SODIUM ALPHA SULFOPELARGONIC ACID
25	SODIUM ETHYL ALPHA SULFOPELARGONATE
26	SODIUM AMYL ALPHA SULFOPELARGONATE
27	SODIUM HEXYL ALPHA SULFOPELARGONATE
28	SODIUM HEPTYL ALPHA SULFOPELARGONATE
7	SODIUM OCTYL ALPHA SULFOPELARGONATE
31	SODIUM 2 OCTYL ALPHA SULFOPELARGONATE
32	SODIUM 2 ETHYLHEXYL ALPHA SULFOPELARGONATE

35 SODIUM H/CF2CF2/3CH2 ALPHA SULFOPELARGONATE

611 ALPHAPHOSPHONO PELARGONIC ACID

605 SODIUM AMYL ALPHAPHOSPHONO PELARGONATE

2 SODIUM OCTYL 1 SULFATE

66 SODIUM OCTYL 2 SULFATE

347 OCTYL TRIMETHYLAMMONIUM OCTYL SULFATE

643 DODECYL TRIMETHYLAMMONIUM OCTANE SULFATE

181 SODIUM OCTYL 1-SULFONATE

340 MAGNESIUM OCTANE SULFONATE

287 OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE

260 SODIUM DI-N-OCTYL SULFOSUCCINATE

392 OCTYLAMMONIUM CHLORIDE

391 OCTYLAMMONIUM DODECYL SULFATE

93 OCTYL TRIMETHYL AMMONIUM BROMIDE

347 OCTYL TRIMETHYLAMMONIUM OCTYL SULFATE

642 OCTYL TRIMETHYLAMMONIUM DECANE SULFATE

385 OCTYL TRIMETHYLAMMONIUM DODECYL SULFATE

287 OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE

353 OCTYL TRIMETHYLAMMONIUM DECANE SULFONATE

483 DIOCTYL DIMETHYL AMMONIUM CHLORIDE

135 OCTYL C BETAINE HYDROCHLORIDE

132 OCTYL C BETAINE

118 OCTYL N BETAINE

251 OCTYL DIMETHYL AMINE OXIDE

710 OCTYL DIMETHYL PHOSPHINE OXIDE

688 OCTYL SULFINYLETHANOL

689 OCTYL SULFINYLPROPANOL

690 OCTYL SULFINYLBUTANOL

648 ALPHA-D-GLUCOSYL OCTANE

424 OCTYL ALPHA-GLYCERYL ETHER

18 OCTYL BETA D GLUCOSIDE

691 OCTYL METHYL SULFOXIDE

423 OCTYL/OXYETHYLENE/1 ALCOHOL--OCTYL GLYCOL ETHER
HOMOGENOUS HEAD GROUP

104 OCTYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP

105 OCTYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP

106 OCTYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP

C-9

530	DECANOIC ACID
299	SODIUM DECANOATE
90	POTASSIUM DECANOATE
668	DIPOTASSIUM OCTYL MALONATE
29	SODIUM NONYL ALPHA SULFOPELARGONATE
606	SODIUM AMYL ALPHAPHOSPHONO CAPRATE
612	ALPHAPHOSPHONO DECANOIC ACID
616	MONOSODIUM ALPHAPHOSPHONO DECANOATE
136	SOLIUM ALPHA DIMETHYL AMINO CAPRATE
295	SODIUM NONYL 1-SULFATE
536	NONYL SULFONIC ACID
94	NONYL TRIMETHYL AMMONIUM BROMIDE
137	ALPHA DIMETHYLAMINO CAPRIC ACID HYDROCHLORIDE
530	DECANOIC ACID
254	NONYL DIMETHYL AMINE OXIDE
580	METHYL /OXYETHYLENE/ 7.0 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
581	METHYL /OXYETHYLENE/ 10.3 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
578	METHYL /OXYETHYLENE/ 11.9 DECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS
582	METHYL /OXYETHYLENE/ 16.0 DECANOATE NATURAL DISTRIBUTION OF HEAD GROUPS

C-10

297	POTASSIUM UNDECANOATE
425	POTASSIUM 1-1-2-DECANE TRICARBOXYLATE
30	SODIUM DECYL ALPHA SULFOPELARGONATE
33	SODIUM /OXO/ DECYL ALPHA SULFOPELARGONATE
3	SODIUM DECYL 1 SULFATE
15	SODIUM DECYL 2 SULFATE
346	DECYL TRIMETHYLAMMONIUM DECYL SULFATE
642	OCTYL TRIMETHYLAMMONIUM DECANE SULFATE
182	SODIUM DECYL 1-SULFONATE
341	MAGNESIUM DECANE SULFONATE
353	OCTYL TRIMETHYLAMMONIUM DECANE SULFONATE

PART 3. Grouped by structure of hydrophobic part and arranged by number of carbon atoms in longest hydrophobic tail and by charge—Continued

288	DECYL TRIMETHYLMONIUM DECANESULFONATE	34	SODIUM DODECYL ALPHA SULFOELARGONATE		
37	DECYLMONIUM CHLORIDE	632	POTASSIUM N-DODECYL BETA-ALANINATE		
411	DECYLMONIUM ACETATE	705	DODECYL SULFURIC ACID		
203	DECYL TRIMETHYL AMMONIUM CHLORIDE	1	SODIUM DODECYL 1 SULFATE		
95	DECYL TRIMETHYL AMMONIUM BROMIDE	67	SODIUM DODECYL 2 SULFATE		
306	DECYL TRIMETHYLMONIUM SULFATE	634	POTASSIUM DODECYL SULFATE		
346	DECYL TRIMETHYLMONIUM DECYL SULFATE	111	LITHIUM DODECYL 1 SULFATE		
288	DECYL TRIMETHYLMONIUM DECANESULFONATE	23	SILVER DODECYL 1 SULFATE		
280	DECYL TRIMETHYL AMMONIUM DODECYL SULFATE	568	MAGNESIUM DODECYL SULFATE		
133	DECYL C BETAINE	24	CALCIUM DODECYL 1 SULFATE		
119	DECYL N BETAINE	569	STRONTIUM DODECYL SULFATE		
252	DECYL DIMETHYL AMINE OXIDE	571	MANGANESE DODECYL SULFATE		
711	DECYL DIMETHYL PHOSPHINE OXIDE	572	COBALTOUS DODECYL SULFATE		
587	DECYL DIMETHYLMONIOPROPANE SULFONATE	575	NICKEL DODECYL SULFATE		
692	DECYL SULFINYLETHANOL	573	CUPRIC DODECYL SULFATE		
19	DECYL BETA D GLUCOSIDE	570	LEAD DODECYL SULFATE		
201	DECYL /OXYETHYLENE/8 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS	574	ZINC DODECYL SULFATE		
202	DECYL /OXYETHYLENE/11 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS	386	AMMONIUM DODECYL SULFATE		
204	DECYL /OXYETHYLENE/12 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS	387	METHYLAMMONIUM DODECYL SULFATE		
107	DECYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP	388	ETHYLAMMONIUM DODECYL SULFATE		
379	DECYL/OXYETHYLENE/4 ALCOHOL HOMOGENEOUS HEAD GROUP	389	BUTYLAMMONIUM DODECYL SULFATE		
378	DECYL/OXYETHYLENE/5 ALCOHOL HOMOGENEOUS HEAD GROUP	409	TRIETHANOLAMMONIUM DODECYL SULFATE		
108	DECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP	112	TETRAMETHYL AMMONIUM DODECYL 1 SULFATE		
109	DECYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP	382	ETHYL TRIMETHYLMONIUM DODECYL SULFATE		
C-11					
531	DODECANOIC ACID	383	383 BUTYL TRIMETHYLMONIUM DODECYL SULFATE		
273	SODIUM DODECANOATE	718	TETRAETHYLMONIUM DODECYL SULFATE		
91	POTASSIUM DODECANOATE	719	TETRABUTYLMONIUM DODECYL SULFATE		
627	CESIUM DODECANOATE	720	1-6-DITRIMETHYLMONIUM-HEXANE/DODECYL SULFATE/2		
277	BENZYL TRIMETHYL AMMONIUM DODECANOATE	410	MORPHOLINIUM DODECYL SULFATE		
527	SODIUM UNDECANE-3-CARBOXYLATE	391	OCTYLAMMONIUM DODECYL SULFATE		
669	DIPOTASSIUM DECYL MALONATE	385	OCTYL TRIMETHYLMONIUM DODECYL SULFATE		
602	ALPHA SULFO LAURIC ACID	280	DECYL TRIMETHYL AMMONIUM DODECYL SULFATE		
235	SODIUM ALPHA SULFO LAURIC ACID	281	DODECYL TRIMETHYLMONIUM DODECYL SULFATE		
603	SODIUM PROPYL ALPHA SULFO LAURATE	636	SODIUM DODECYL THIOSULFATE		
613	ALPHAPHOSPHONO DODECANOIC ACID	462	SODIUM MONOLAURIN SULFATE		
617	MONOSODIUM ALPHAPHOSPHONO DODECANOATE	541	SODIUM DODECYL MONO-OXYETHYLENE SULFATE		
620	DISODIUM ALPHAPHOSPHONO DODECANOATE	542	SODIUM DODECYL DIOXYETHYLENE SULFATE		
607	SODIUM ISOPROPYL ALPHAPHOSPHONO LAURATE	113	SODIUM DODECYL TRI-OXYETHYLENE SULFATE		
311	SODIUM UNDECYL 1-SULFATE	543	SODIUM DODECYL TETRA-OXYETHYLENE SULFATE		
72	SODIUM UNDECYL 3 SULFATE	114	SODIUM DODECYL /OXYETHYLENE/17.5 SULFATE		
80	SODIUM UNDECYL 6 SULFATE	597	SODIUM DODECYL MONO-OXYPROPYL SULFATE		
639	SODIUM UNDECYL THIOSULFATE	200	DODECYL SULFONIC ACID		
537	UNDECYL SULFONIC ACID	179	SODIUM DODECANE 1-SULFONATE		
418	SODIUM UNDECYL SULFONATE	175	SODIUM DODECANE 2-SULFONATE		
96	UNDECYL TRIMETHYL AMMONIUM BROMIDE	40	POTASSIUM DODECYL 1 SULFONATE		
531	DODECANOIC ACID	635	LITHIUM DODECYL SULFONATE		
120	UNDECYL N BETAINE	342	MAGNESIUM DODECANE SULFONATE		
713	UNDECYL DIMETHYL AMINE OXIDE	CATIONIC			
495	SUCROSE MONOLAURATE	38	DODECYL AMMONIUM CHLORIDE		
276	NONAETHYLENE GLYCOL MONODODECANOATE NATURAL DISTRIBUTION OF HEAD GROUPS	628	DODECYLAMMONIUM BROMIDE		
583	METHYL /OXYETHYLENE/ 6.0 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS	482	DODECYL AMMONIUM NITRATE		
584	METHYL /OXYETHYLENE/ 8.4 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS	412	DODECYLAMMONIUM ACETATE		
585	METHYL /OXYETHYLENE/ 11.2 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS	449	DODECYLMETHYL AMMONIUM CHLORIDE		
579	METHYL /OXYETHYLENE/ 12.5 DODECANOATE REDUCED POLYDISPERSITY OF HEAD GROUPS	450	DODECYLDIMETHYL AMMONIUM CHLORIDE		
440	/OXYETHYLENE/4 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	41	DODECYL TRIMETHYL AMMONIUM CHLORIDE		
441	/OXYETHYLENE/8 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	307	DODECYL TRIMETHYLMONIUM SULFATE		
442	/OXYETHYLENE/10 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	97	DODECYL TRIMETHYL AMMONIUM BROMIDE		
443	/OXYETHYLENE/20 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	126	DODECYL TRIMETHYLMONIUM IODIDE		
444	/OXYETHYLENE/25 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	130	DODECYL TRIMETHYL AMMONIUM FLUORIDE		
520	UNDECYL/OXYETHYLENE/8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	131	DODECYL TRIMETHYLMONIUM NITRATE		
550	LAURIC ACID DIETHANOLAMINE CONDENSATE	129	DODECYL TRIMETHYL AMMONIUM BROMATE		
C-12					
ANIONIC					
351	POTASSIUM TRIDECANOATE	127	DODECYL TRIMETHYL AMMONIUM IODATE		
421	POTASSIUM 1-1-2-DODECANE TRICARBOXYLATE	128	DODECYL TRIMETHYL AMMONIUM FORMATE		
		643	DODECYL TRIMETHYLMONIUM OCTANE SULFATE		
		281	DODECYL TRIMETHYLMONIUM DODECYL SULFATE		
		384	HEXYL TRIMETHYLMONIUM DODECYL SULFATE		
		399	DODECYL DIMETHYL ETHYLAMMONIUM CHLORIDE		
		345	DIDODECYL DIMETHYLMONIUM CHLORIDE		
		400	DODECYL METHYL DIETHYLMONIUM CHLORIDE		
		401	DODECYL TRIETHYLMONIUM CHLORIDE		
		500	DODECYL TRI-2-HYDROXYETHYL/AMMONIUM CHLORIDE		
		22	DIMETHYL DODECYL AMINE OXIDE HYDROCHLORIDE		
		293	DODECYL DIMETHYL AMINE OXIDE HYDROBROMIDE		
		124	DODECYL N BETAINE HYDROCHLORIDE		
		633	N-DODECYL BETA-ALANINE HYDROCHLORIDE		
NONIONIC					
		121	DODECYL N BETAINE		
		592	DODECYL N-DIETHYL N-BETAINE		
		134	DODECYL C BETAINE		
		21	DIMETHYL DODECYL AMINE OXIDE		
		712	DODECYL DIMETHYL PHOSPHINE OXIDE		
		590	DODECYL DIMETHYLMONIOPROPANE CARBOXYLATE		
		588	DODECYL DIMETHYLMONIOPROPANE SULFONATE		

PART 3. Grouped by structure of hydrophobic part and arranged by number of carbon atoms in longest hydrophobic tail and by charge—Continued

591	DODECYL DIMETHYLPHOSPHONIOPROPANE SULFONATE	73	SODIUM TETRADECYL 3 SULFATE
595	DODECYL DIMETHYL AMMONIOPROPANE SULFATE	17	SODIUM TETRADECYL 4 SULFATE
594	DODECYL DIPROPYL AMMONIOPROPANE SULFONATE	77	SODIUM TETRADECYL 5 SULFATE
593	DIMETHYL DODECYLAMMONIOPROPANE HYDROXY SULFONATE	525	SODIUM TETRADECYL 6-SULFATE
20	DODECYL BETA D GLUCOSIDE	84	SODIUM TETRADECYL 7 SULFATE
649	ALPHA-D-GLUCOSYL DODECANE	637	LITHIUM TETRADECYL SULFATE
205	DODECYL /OXYETHYLENE/12 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS	576	CUPRIC TETRADECYL SULFATE
325	DODECYL/OXYETHYLENE/4 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS	544	SODIUM TETRADECYL MONO-OXYETHYLENE SULFATE
377	DODECYL/OXYETHYLENE/ 5 ALCOHOL HOMOGENEOUS HEAD GROUP	545	SODIUM TETRADECYL DI-OXYETHYLENE SULFATE
110	DODECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP	546	SODIUM TETRADECYL TRI-OXYETHYLENE SULFATE
468	DODECYL/OXYETHYLENE/6.2 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	598	SODIUM TETRADECYL MONO-OXYPROPYL SULFATE
115	DODECYL /OXYETHYLENE/7 ALCOHOL REDUCED POLYDISPERSION OF HEAD GROUPS	599	SODIUM TETRADECYL DI-OXYPROPYL SULFATE
488	DODECYL/OXYETHYLENE/7 ALCOHOL HOMOGENEOUS HEAD GROUP	243	TETRADECANE 1-SULFONIC ACID
469	DODECYL/OXYETHYLENE/7.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	183	SODIUM TETRADECYL 1-SULFONATE
163	DODECYL /OXYETHYLENE/ 8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	176	SODIUM TETRADECANE 2-SULFONATE
489	DODECYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP	39	TETRADECYL AMMONIUM CHLORIDE
310	DODECYL/OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	413	TETRADECYLAMMONIUM ACETATE
470	DODECYL/OXYETHYLENE/10.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	42	TETRADECYL TRIMETHYL AMMONIUM CHLORIDE
490	DODECYL/OXYETHYLENE/12 ALCOHOL HOMOGENEOUS HEAD GROUP	98	TETRADECYL TRIMETHYL AMMONIUM BROMIDE
164	DODECYL /OXYETHYLENE/ 12 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	308	TETRADECYL TRIMETHYLAMMONIUM SULFATE
326	DODECYL/OXYETHYLENE/14 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS	291	TETRADECYL TRIPROPYLMAMMONIUM BROMIDE
471	DODECYL/OXYETHYLENE/15.8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	125	TETRADECYL N BETAINE HYDROCHLORIDE
716	DODECYL /OXYETHYLENE/16 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS	122	TETRADECYL N BETAINE
165	DODECYL /OXYETHYLENE/ 18 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	253	TETRADECYL DIMETHYL AMINE OXIDE
472	DODECYL/OXYETHYLENE/20.4 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	289	TETRADECYL/OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP
327	DODECYL/OXYETHYLENE/23 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS		
166	DODECYL /OXYETHYLENE/ 23 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS		
473	DODECYL/OXYETHYLENE/27.1 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS		
170	DODECYL /OXYETHYLENE/ 28 ALCOHOL REDUCED DISTRIBUTION OF HEAD GROUPS		
116	DODECYL /OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSION OF HEAD GROUPS		
474	DODECYL/OXYETHYLENE/38.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS		
475	DODECYL/OXYETHYLENE/67.3 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS		
C-13			
532	TETRADECANOIC ACID	533	HEXADECANOIC ACID
298	SODIUM TETRADECANOATE	300	SODIUM HEXADECANOATE
92	POTASSIUM TETRADECANOATE	185	POTASSIUM HEXADECANOATE
670	DIPOTASSIUM DODECYL MALONATE	671	DIPOTASSIUM TETRADECYL MALONATE
189	ALPHA SULFOMYRISTIC ACID	190	ALPHA SULFOPALMITIC ACID
236	SODIUM ALPHA SULFO MYRISTIC ACID	237	SODIUM ALPHA SULFO PALMITIC ACID
604	SODIUM METHYL ALPHA SULFO MYRISTATE	36	SODIUM METHYL ALPHA SULFOPALMITATE
618	MONOSODIUM ALPHAPHOSPHONO TETRADECANOATE	192	SODIUM ETHYL ALPHA SULFOPALMITATE
608	SODIUM METHYL ALPHAPHOSPHONO MYRISTATE	193	SODIUM PROPYL ALPHA SULFOPALMITATE
233	DISODIUM ALPHA SULFO MYRISTATE	234	DISODIUM ALPHA SULFO PALMITATE
614	ALPHAPHOSPHONO TETRADECANOIC ACID	197	DISODIUM 2 SULFOETHYL ALPHA SULFOPALMITATE
621	DISODIUM ALPHAPHOSPHONO TETRADECANOATE	615	ALPHAPHOSPHONO HEXADECANOIC ACID
624	TRISODIUM ALPHAPHOSPHONO TETRADECANOATE	619	MONOSODIUM ALPHAPHOSPHONO HEXADECANOATE
68	SODIUM TRIDECYL 2 SULFATE	609	SODIUM ISOPROPYL ALPHAPHOSPHONO PALMITATE
83	SODIUM TRIDECYL 7 SULFATE	622	DISODIUM ALPHAPHOSPHONO HEXADECANOATE
242	TRIDECANE 1-SULFONIC ACID	625	TRISODIUM ALPHAPHOSPHONO HEXADECANOATE
229	SODIUM TRIDECANE 1-SULFONATE	69	SODIUM PENTADECYL 2 SULFATE
402	TRIDECYL TRIMETHYLAMMONIUM CHLORIDE	74	SODIUM PENTADECYL 3 SULFATE
532	TETRADECANOIC ACID	78	SODIUM PENTADECYL 5 SULFATE
497	SUCROSE MONOMYRISTATE	85	SODIUM PENTADECYL 8 SULFATE
465	TRIDECYL/OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	244	PENTADECANE 1-SULFONIC ACID
466	TRIDECYL/SECONDARY/ /OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	230	SODIUM PENTADECANE 1-SULFONATE
		533	HEXADECANOIC ACID
		498	SUCROSE MONOPALMITATE
C-14			
426	POTASSIUM 1-1-2-TETRADECANE TRICARBOXYLATE	422	POTASSIUM 1-1-2-HEXADECANE TRICARBOXYLATE
4	SODIUM TETRADECYL 1 SULFATE	5	SODIUM HEXADECYL 1 SULFATE
16	SODIUM TETRADECYL 2 SULFATE	75	SODIUM HEXADECYL 4 SULFATE
		81	SODIUM HEXADECYL 6 SULFATE
		86	SODIUM HEXADECYL 8 SULFATE
		638	LITHIUM HEXADECYL SULFATE
		577	CUPRIC HEXADECYL SULFATE
		60	TRIETHANOL AMMONIUM HEXADECYL SULFATE
		52	SODIUM HEXADECYL MONO OXYETHYLENE SULFATE
		53	SODIUM HEXADECYL DI OXYETHYLENE SULFATE
		54	SODIUM HEXADECYL TRI OXYETHYLENE SULFATE
		55	SODIUM HEXADECYL TETRA OXYETHYLENE SULFATE
		600	SODIUM HEXADECYL MONO-OXYPROPYL SULFATE
		245	HEXADECANE 1-SULFONIC ACID
		184	SODIUM HEXADECYL 1-SULFONATE
		177	SODIUM HEXADECANE 2-SULFONATE
		408	POTASSIUM HEXADECANE 1-SULFONATE
		186	HEXADECYL AMMONIUM CHLORIDE
		414	HEXADECYLAMMONIUM ACETATE
		265	HEXADECYL TRIMETHYL AMMONIUM CHLORIDE
		99	HEXADECYL TRIMETHYLAMMONIUM BROMIDE
		309	HEXADECYL TRIMETHYLAMMONIUM SULFATE
		478	HEXADECYL DIMETHYLETHYLAMMONIUM BROMIDE
		266	HEXADECYL DIMETHYL 2-HYDROXYETHYL AMMONIUM CHLORIDE
		269	HEXADECYL DIMETHYL2,3-DIHYDROXYPROPYLAMMONIUM CHLORIDE
		267	HEXADECYL DI-/2-HYDROXYETHYL/METHYL AMMONIUM CHLORIDE
		268	HEXADECYL TRI-/2-HYDROXYETHYL/ AMMONIUM CHLORIDE
		292	HEXADECYL TRIPROPYLMAMMONIUM BROMIDE
		666	HEXADECYL TRIBUTYLAMMONIUM BROMATE
		123	HEXADECYL N BETAINE
		589	HEXADECYL DIMETHYLAMMONIOPROPANE SULFONATE
		499	SUCROSE DI-PALMITATE

PART 3. Grouped by structure of hydrophobic part and arranged by number of carbon atoms in longest hydrophobic tail and by charge—Continued

282 HEXADECYL /OXYETHYLENE/ 6 ALCOHOL HOMOCENOUS HEAD GROUP
 330 HEXADECYL/OXYETHYLENE/7 ALCOHOL HOMOCENOUS HEAD GROUP
 331 HEXADECYL/OXYETHYLENE/9 ALCOHOL HOMOCENOUS HEAD GROUP
 332 HEXADECYL/OXYETHYLENE/12 ALCOHOL HOMOCENOUS HEAD GROUP
 333 HEXADECYL/OXYETHYLENE/15 ALCOHOL HOMOCENOUS HEAD GROUP
 535 HEXADECYL/OXYETHYLENE/18 ALCOHOL
 334 HEXADECYL/OXYETHYLENE/21 ALCOHOL HOMOCENOUS HEAD GROUP
 117 HEXADECYL /OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSION OF HEAD GROUPS

C-17

534 OCTADECANOIC ACID
 448 SODIUM OCTADECANOATE /STEARATE/
 256 POTASSIUM STEARATE
 672 DIPOTASSIUM HEXADECYL MALONATE
 191 ALPHA SULFOSTEARIC ACID
 9 SODIUM ALPHA SULFOSTEARIC ACID
 194 SODIUM METHYL ALPHA SULFOSTEARATE
 195 SODIUM ETHYL ALPHA SULFOSTEARATE
 196 SODIUM PROPYL ALPHA SULFOSTEARATE
 199 SODIUM ISOPROPYL ALPHA SULFOSTEARATE
 10 DISODIUM ALPHA SULFOSTEARATE
 198 DISODIUM 2 SULFOETHYL ALPHA SULFOSTEARATE
 553 XYLYL SULFOSTEARIC ACID
 610 SODIUM METHYL ALPHAPHOSPHONO STEARATE
 623 DISODIUM ALPHAPHOSPHONO OCTADECANOATE
 626 TRISODIUM ALPHAPHOSPHONO OCTADECANOATE
 70 SODIUM HEPTADECYL 2 SULFATE
 87 SODIUM HEPTADECYL 9 SULFATE
 246 HEPTADECANE 1-SULFONIC ACID
 231 SODIUM HEPTADECANE 1-SULFONATE
 534 OCTADECANOIC ACID
 496 SUCROSE MONOSTEARATE

C-18

673 DIPOTASSIUM OCTADECYL MALONATE
 64 SODIUM OCTADECYL 1 SULFATE
 71 SODIUM OCTADECYL 2 SULFATE
 76 SODIUM OCTADECYL 4 SULFATE
 82 SODIUM OCTADECYL 6 SULFATE
 65 TRIETHANOLAMMONIUM OCTADECYL 1 SULFATE
 56 SODIUM OCTADECYL MONO OXYETHYLENE SULFATE
 57 SODIUM OCTADECYL DI OXYETHYLENE SULFATE
 58 SODIUM OCTADECYL TRI OXYETHYLENE SULFATE
 59 SODIUM OCTADECYL TETRA OXYETHYLENE SULFATE
 601 SODIUM OCTADECYL MONO-OXYPROPYL SULFATE
 232 OCTADECANE 1-SULFONIC ACID
 349 SODIUM OCTADECANE 1-SULFONATE
 178 SODIUM OCTADECANE 2-SULFONATE
 419 SODIUM ALPHA-HEPTYL UNDECYL SULFONATE
 187 OCTADECYL AMMONIUM CHLORIDE
 415 OCTADECYLAMMONIUM ACETATE
 270 OCTADECYL TRIMETHYLLAMMONIUM CHLORIDE
 477 OCTADECYL TRIMETHYLLAMMONIUM BROMIDE
 654 OCTADECYL TRIMETHYLLAMMONIUM NITRATE
 658 OCTADECYL TRIMETHYLLAMMONIUM BROMATE
 659 OCTADECYL TRIMETHYLLAMMONIUM FORMATE
 667 OCTADECYL TRIMETHYLLAMMONIUM OXALATE
 662 OCTADECYL TRIETHYLLAMMONIUM BROMATE
 663 OCTADECYL TRIPROPYLLAMMONIUM BROMATE
 664 OCTADECYL TRIBUTYLLAMMONIUM BROMATE
 665 OCTADECYL TRIAMYLAMMONIUM BROMATE
 435 OCTADECYL/OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 328 OCTADECYL/OXYETHYLENE/14 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS
 436 OCTADECYL/OXYETHYLENE/20 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
 329 OCTADECYL/OXYETHYLENE/100 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS

C-19

79 SODIUM NONADECYL 5 SULFATE
 88 SODIUM 1 NONYL DECYL SULFATE

C-20

683 SODIUM EICOSYLBENZENE SULFONATE

C-29

89 SODIUM 1 TETRADECYL PENTADECYL SULFATE

ISO-ALKANES

261 SODIUM DI-ISOBUTYL SULFOSUCCINATE
 394 1-METHYL PROPYL/OXYETHYLENE/6 ALCOHOL HOMOCENOUS HEAD GROUP
 352 SODIUM DI-1-METHYLISOAMYL SULFOSUCCINATE
 395 2-ETHYL BUTYL/OXYETHYLENE/6 ALCOHOL HOMOCENOUS HEAD GROUP
 262 SODIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE
 286 AMMONIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE
 396 2-PROPYL PENTYL/OXYETHYLENE/6 ALCOHOL HOMOCENOUS HEAD GROUP
 515 ISO-OCTYL/OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 516 NYONYL/OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 397 2-BUTYL HEXYL/OXYETHYLENE/6 ALCOHOL HOMOCENOUS HEAD GROUP
 398 2-BUTYL HEXYL/OXYETHYLENE/9 ALCOHOL HOMOCENOUS HEAD GROUP
 517 DECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 519 DECYL/OXO-PROCESS/ /OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 143 DECYL /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 144 DECYL /OXYETHYLENE/ 10.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 145 DECYL /OXYETHYLENE/ 14.9 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 146 DECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 147 DECYL /OXYETHYLENE/ 29.4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 518 TRIMETHYLNONYL/OXYETHYLENE/6 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION

C-13

312 TRIDECYL/OXYETHYLENE/5 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 148 TRIDECYL /OXYETHYLENE/ 5.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 313 TRIDECYL/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 521 TRIDECYL/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 149 TRIDECYL /OXYETHYLENE/ 10.1 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 314 TRIDECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 150 TRIDECYL /OXYETHYLENE/ 15.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 315 TRIDECYL/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 151 TRIDECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 316 TRIDECYL/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN, REDUCED OE DISTRIBUTION
 152 TRIDECYL /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 431 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/8.5 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 432 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 433 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/15 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 434 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/22 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
 526 SODIUM 2-DI-N-HEXYL ETHYL SULFATE
 524 HEXADECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION

3B. ALKYL ARYL

ALKYL AND ARYL

354 HEXYL BENZYL DIMETHYLLAMMONIUM CHLORIDE
 356 DECYL BENZYL DIMETHYLLAMMONIUM CHLORIDE
 403 DODECYL DIMETHYLPHENYLAMMONIUM CHLORIDE
 279 DODECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
 404 DODECYL DIMETHYL 2-PHENYLETHYLAMMONIUM CHLORIDE
 C6H5CH2CH2/N/CH3/2/C12H25
 407 DODECYL DIMETHYL 3-PHENYLPROPYLAMMONIUM CHLORIDE
 C6H5CH2CH2/N/CH3/2/C12H25
 405 DODECYL METHYLETHYL BENZYLAMMONIUM CHLORIDE
 357 TETRADECYL BENZYL DIMETHYLAMMONIUM CHLORIDE

PART 3. Grouped by structure of hydrophobic part and arranged by number of carbon atoms in longest hydrophobic tail and by charge – Continued

275 HEXADECYL DIMETHYLBENZYLAMMONIUM CHLORIDE
 358 OCTADECYL BENZYL DIMETHYLBENZYLAMMONIUM CHLORIDE

1-N-ALKYL ARYL

645 PARA/BETA-D-GLUCOSYL/ETHYLBENZENE
 646 PARA/BETA-D-GLUCOSYL/PROPYLBENZENE
 445 SODIUM PARA-BIS-/N-BUTYL/ BENZENE SULFONATE
 681 SODIUM DIBUTYL BENZENE SULFONATE
 647 PARA/BETA-D-GLUCOSYL/BUTYLBENZENE
 650 PARA/BETA-D-XYLOSYL/BUTYL BENZENE
 501 SODIUM HEXYL BENZENE SULFONATE
 446 SODIUM PARA-BIS-/N-HEXYL/ BENZENE SULFONATE
 502 SODIUM HEPTYL BENZENE SULFONATE
 49 SODIUM P OCTYL BENZENE SULFONATE
 503 SODIUM OCTYL BENZENE SULFONATE
 447 SODIUM PARA-BIS-/N-OCTYL/ BENZENE SULFONATE
 675 OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 493 SODIUM P NONYL BENZENE SULFONATE
 504 SODIUM NONYL BENZENE SULFONATE
 463 NONYL BENZENE/OXYETHYLENE/11 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 464 NONYL BENZENE/OXYETHYLENE/15 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 50 SODIUM P DECYLBENZENE SULFONATE
 505 SODIUM DECYLBENZENE SULFONATE
 522 DECYLBENZENE ORTHO/OXYETHYLENE/11.2 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 523 DECYLBENZENE PARA/OXYETHYLENE/11.0 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 51 SODIUM P DODECYLBENZENE SULFONATE
 506 SODIUM DODECYLBENZENE SULFONATE
 507 SODIUM TETRADECYL BENZENE SULFONATE
 596 TETRAHYDROANACARDOL AMMONIUM MONOSULFONATE
 508 SODIUM HEXADECYLBENZENE SULFONATE
 552 TOLYL SULFOSTEARIC ACID
 509 SODIUM OCTADECYLBENZENE SULFONATE

OTHER N-ALKYL ARYL

172 SODIUM 2-N-OCTYL BENZENE SULFONATE
 173 SODIUM 2-N-DECYL BENZENE SULFONATE
 45 SODIUM P 1 METHYL DECYLBENZENE SULFONATE
 171 SODIUM 2-N-DODECYLBENZENE SULFONATE
 301 SODIUM 3-N-DODECYLBENZENE SULFONATE
 302 SODIUM 4-N-DODECYLBENZENE SULFONATE
 514 SODIUM 6-N-DODECYLBENZENE SULFONATE
 46 SODIUM P 1 METHYL DODECYLBENZENE SULFONATE
 174 SODIUM 2-N-TETRADECYL BENZENE SULFONATE
 47 SODIUM P 1 METHYL TETRADECYL BENZENE SULFONATE
 48 SODIUM P 1 METHYL HEXADECYLBENZENE SULFONATE
 551 PHENYL SULFOSTEARIC ACID

ISO-ALKYL ARYL – KNOWN BRANCHING

674 DIISOBUTYL BENZENE/OXYETHYLENE/6 ALCOHOL (DB) BRANCHED
 CHAIN. NATURAL OE DISTRIBUTION
 510 SODIUM 2-ETHYL-HEXYLBENZENE SULFONATE
 676 SODIUM 1,1,3,3-TETRAMETHYL BUTYL BENZENE SULFONATE
 677 SODIUM DI-/1,1,3,3-TETRAMETHYL BUTYL/BENZENESULFONATE

P-TERTIARY OCTYL BENZENE

207 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL HOMOGENOUS
 HEAD GROUP
 217 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 208 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL HOMOGENOUS
 HEAD GROUP
 218 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 209 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL HOMOGENOUS
 HEAD GROUP
 219 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 210 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL HOMOGENOUS
 HEAD GROUP
 220 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 211 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL HOMOGENOUS
 HEAD GROUP
 221 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 212 P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL HOMOGENOUS
 HEAD GROUP
 222 P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS

213 P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL HOMOGENOUS
 HEAD GROUP
 223 P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 214 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL HOMOGENOUS
 HEAD GROUP
 224 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 215 P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL HOMOGENOUS
 HEAD GROUP
 225 P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 206 TRITON X-100(P-T-OCTYL BENZENE/OXYETHYLENE/9NATURAL
 DISTRIBUTION OF HEAD GROUPS
 216 P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL HOMOGENOUS
 HEAD GROUP
 317 T-OCTYL BENZENE/OXYETHYLENE/10 ALCOHOL REDUCED
 POLYDISPERSITY OF HEAD GROUPS
 226 P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 227 P-T-OCTYL BENZENE /OXYETHYLENE/16 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 318 T-OCTYL BENZENE/OXYETHYLENE/20 ALCOHOL REDUCED
 POLYDISPERSITY OF HEAD GROUPS
 319 T-OCTYL BENZENE/OXYETHYLENE/30 ALCOHOL REDUCED
 POLYDISPERSITY OF HEAD GROUPS
 228 P-T-OCTYL BENZENE /OXYETHYLENE/40 ALCOHOL NATURAL
 DISTRIBUTION OF HEAD GROUPS
 272 TRI-ISOPROPYL BENZENE SULFONIC ACID
 271 SODIUM TRI-ISO-PROPYLBENZENE SULFONATE
 678 SODIUM 1,1-DIMETHYLNONYLBENZENE SULFONATE
 511 SODIUM 2-PROPYL-HEPTYLBENZENE SULFONATE
 679 SODIUM 3-BUTYL HEPTYLBENZENE SULFONATE
 680 SODIUM 2-PROPYL-4-METHYL-OCTYL BENZENE SULFONATE
 492 SODIUM TETRAPROPYLENE/1-3-5-7-TETRAMETHYL-
 OCTYL/BENZENE SULFONATE
 512 SODIUM 2-BUTYL-OCTYL BENZENE SULFONATE
 513 SODIUM 2-AMYL-NONYLBENZENE SULFONATE

ISO-ALKYL ARYL – BRANCHING UNSPECIFIED

335 OCTYL BENZENE/OXYETHYLENE/8.5 ALCOHOL BRANCHED CHAIN,
 NATURAL OE DISTRIBUTION
 138 SODIUM NONYL BENZENE SULFONATE BRANCHED HYDROCARBON

ISO-NONYL BENZENE

153 NONYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED
 CHAIN, NATURAL OE DISTRIBUTION
 154 NONYL BENZENE /OXYETHYLENE/ 9.8 ALCOHOL BRANCHED
 CHAIN, NATURAL OE DISTRIBUTION
 336 NONYL BENZENE/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN,
 NATURAL OE DISTRIBUTION
 320 NONYL BENZENE/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN,
 REDUCED OE DISTRIBUTION
 167 NONYL BENZENE /OXYETHYLENE/ 10 ALCOHOL BRANCHED CHAIN,
 NATURAL OE DISTRIBUTION
 337 NONYL BENZENE/OXYETHYLENE/10.5 ALCOHOL BRANCHED CHAIN,
 NATURAL OE DISTRIBUTION
 168 NONYL BENZENE /OXYETHYLENE/ 15 ALCOHOL BRANCHED CHAIN,
 NATURAL OE DISTRIBUTION
 321 NONYL BENZENE/OXYETHYLENE/15 ALCOHOL BRANCHED CHAIN,
 REDUCED OE DISTRIBUTION
 155 NONYL BENZENE /OXYETHYLENE/ 15.4 ALCOHOL BRANCHED
 CHAIN, NATURAL OE DISTRIBUTION
 322 NONYL BENZENE/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN,
 REDUCED OE DISTRIBUTION
 156 NONYL BENZENE /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED
 CHAIN, NATURAL OE DISTRIBUTION
 323 NONYL BENZENE/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN,
 REDUCED OE DISTRIBUTION
 721 NONYL BENZENE /OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN
 DIALYSED
 169 NONYL BENZENE /OXYETHYLENE/ 30 ALCOHOL BRANCHED CHAIN,
 NATURAL OE DISTRIBUTION
 157 NONYL BENZENE /OXYETHYLENE/ 30.2 ALCOHOL BRANCHED
 CHAIN, NATURAL OE DISTRIBUTION
 324 NONYL BENZENE/OXYETHYLENE/50 ALCOHOL BRANCHED CHAIN,
 REDUCED OE DISTRIBUTION
 338 NONYL BENZENE/OXYETHYLENE/100 ALCOHOL BRANCHED CHAIN,
 NATURAL OE DISTRIBUTION
 140 SODIUM DECYLBENZENE SULFONATE BRANCHED HYDROCARBON
 CHAIN
 139 SODIUM DODECYLBENZENE SULFONATE BRANCHED HYDROCARBON
 CHAIN

PART 3. Grouped by structure of hydrophobic part and arranged by number of carbon atoms in longest hydrophobic tail and by charge—Continued

- 158 DODECYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
- 159 DODECYL BENZENE /OXYETHYLENE/ 10.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
- 160 DODECYL BENZENE /OXYETHYLENE/ 15.1 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
- 161 DODECYL BENZENE /OXYETHYLENE/ 20.1 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
- 162 DODECYL BENZENE /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION
- 141 SODIUM TRIDECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN
- 142 SODIUM PENTADECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN

ALKYL-ARYL — PRESENCE OF BRANCHING UNCERTAIN

- 682 SODIUM DIBUTYL NAPHTHALENE SULFONATE /NEKAL/
- 11 SODIUM ALPHA SULFO PHENYL STEARIC ACID
- 8 DISODIUM ALPHA SULFOPHENYLSTEARATE

3C. UNSATURATED AND SUBSTITUTED

UNSATURATED HYDROCARBON

- 562 SODIUM DODECENYL SULFATE
- 263 SODIUM OLEATE /CIS-9-OCTADECENOATE/
- 305 POTASSIUM OLEATE/ CIS-9-OCTADECENOATE/
- 285 HEXANOLAMINE-CH₃CH/OH/CH₂/CH₃/2NH₂-OLEATE
- 61 SODIUM OLEYL/CIS 9 OCTADECENOYL/ SULFATE
- 547 SODIUM OLEYL MONO-OXYETHYLENE SULFATE
- 548 SODIUM OLEYL DI-OXYETHYLENE SULFATE
- 549 SODIUM OLEYL TRI-OXYETHYLENE SULFATE
- 437 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/12 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
- 438 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/16 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS
- 439 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/20 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUP
- 264 SODIUM ELAIDATE /TRANS-9-OCTADECENOATE/
- 629 POTASSIUM ELAIDATE/TRANS-9-OCTADECENOATE/
- 283 HEXANOLAMINE-CH₃CH/OH/CH₂/CH₃/2NH₂-ELAIDATE
- 62 SODIUM ELAIDYL/TRANS 9 OCTADECENOYL/SULFATE

OXYGEN

- 451 PARA DI-ISOBUTYLPHENOXYETHOXYETHYL DIMETHYLBENZYL AMMONIUM CHLORIDE/HYAMINE 1622/
- 238 SODIUM DODECANE 1-HYDROXY 2-SULFONATE
- 247 DODECANE 1-HYDROXY 2-SULFONIC ACID
- 370 DODECYL 3-4-METHYLENEDIOXYBENZYL DIMETHYLMONIUM CHLORIDE
- 371 DODECYL 3-4-DIMETHOXYBENZYL DIMETHYLMONIUM CHLORIDE
- 248 TETRADECANE 1-HYDROXY 2-SULFONIC ACID
- 239 SODIUM TETRADECANE 1-HYDROXY 2-SULFONATE
- 249 HEXADECANE 1-HYDROXY 2-SULFONIC ACID
- 240 SODIUM HEXADECANE 1-HYDROXY 2-SULFONATE
- 255 POTASSIUM 9,10 DIHYDROXY STEARATE
- 630 POTASSIUM RICINOLEATE/12 HYDROXY OLEATE/
- 631 POTASSIUM RICINELAIDATE/12 HYDROXY ELAIDATE/
- 14 DISODIUM 9,10 DIHYDROXY ALPHA SULFOSTEARATE
- 250 OCTADECANE 1-HYDROXY 2-SULFONIC ACID
- 241 SODIUM OCTADECANE 1-HYDROXY 2-SULFONATE

NITROBENZYL

- 359 OCTYL 4-NITROBENZYL DIMETHYLMONIUM CHLORIDE
- 360 DECYL 4-NITROBENZYL DIMETHYLMONIUM CHLORIDE
- 361 DODECYL 4-NITROBENZYL DIMETHYL AMMONIUM CHLORIDE

- 368 DODECYL 2-HYDROXY-5-NITROBENZYL DIMETHYLMONIUM CHLORIDE
- 362 TETRADECYL 4-NITROBENZYL DIMETHYLMONIUM CHLORIDE
- 363 HEXADECYL 4-NITROBENZYL DIMETHYLMONIUM CHLORIDE
- 364 OCTADECYL 4-NITROBENZYL DIMETHYLMONIUM CHLORIDE

HALOGEN

- 428 PERFLUORO ACETIC ACID
- 429 PERFLUORO PROPIONIC ACID
- 430 PERFLUORO BUTYRIC ACID
- 706 PERFLUORO PROPYLAMINE
- 416 PERFLUORO HEXANOIC ACID
- 701 POTASSIUM PERFLUOROHEXANOATE
- 417 PERFLUORO OCTANOIC ACID
- 702 POTASSIUM PERFLUORODECANOATE
- 452 3-4-DICHLOROPERFLUORO BUTYRIC/KEL-F/ACID
- 453 3-5-6-TRICHLOROPERFLUORO HEXANOIC/KEL-F/ACID
- 454 3-5-7-8-TETRACHLOROPERFLUORO OCTANOIC/KEL-F/ACID
- 374 DODECAFLUOROHEPTANOIC ACID H/CF₂/6COOH
- 355 AMMONIUM DODECAFLUOROHEPTANOATE H/CF₂/6COONH₄
- 375 HEXADECAFLUORONONANOIC ACID H/CF₂/8COOH
- 372 AMMONIUM HEXADECAFLUORONONANOATE H/CF₂/8 COO NH₄
- 373 AMMONIUM EICOSAFLUOROUNDECANOATE H/CF₂/10 COO NH₄
- 348 NN-DIMETHYL 1-1-DIHYDROPERDAFLUORO OCTYL AMINE N-OXIDE /C7F15CH₂N/CH₃/20/
- 365 DODECYL 2-CHLOROBENZYL DIMETHYLMONIUM CHLORIDE
- 367 DODECYL 4-CHLOROBENZYL DIMETHYLMONIUM CHLORIDE
- 366 DODECYL 2-4-DICHLOROBENZYL DIMETHYLMONIUM CHLORIDE
- 369 DODECYL 3-4-DICHLOROBENZYL DIMETHYLMONIUM CHLORIDE
- 406 DODECYL DIMETHYL META-TRIFLUOROMETHYLBENZYL AMMONIUM CHLORIDE CF₃C₆H₄CH₂/N/CH₃/2/C12H25
- 12 SODIUM 9,10 DICHLORO ALPHA SULFOSTEARIC ACID
- 13 DISODIUM 9,10 DICHLORO ALPHA SULFOSTEARATE
- 63 SODIUM 9,10 DICHLORO OCTADECYL SULFATE

3D. HETEROCYCLIC

- 100 OCTYL PYRIDINIUM BROMIDE
- 458 DECYL PYRIDINIUM IODIDE
- 101 UNDECYL PYRIDINIUM BROMIDE
- 278 DODECYL PYRIDINIUM CHLORIDE
- 290 DODECYL PYRIDINIUM BROMIDE
- 376 DODECYL PYRIDINIUM IODIDE
- 102 TETRADECYL PYRIDINIUM BROMIDE
- 479 TETRADECYL PYRIDINIUM IODIDE
- 274 HEXADECYL PYRIDINIUM CHLORIDE
- 427 HEXADECYL PYRIDINIUM BROMIDE
- 480 HEXADECYL PYRIDINIUM IODIDE
- 660 HEXADECYL PYRIDINIUM IODATE
- 655 OCTADECYL PYRIDINIUM CHLORIDE
- 657 OCTADECYL PYRIDINIUM BROMIDE
- 481 OCTADECYL PYRIDINIUM IODIDE
- 656 OCTADECYL PYRIDINIUM NITRATE
- 661 OCTADECYL PYRIDINIUM IODATE
- 693 N-CETYLYL 2-METHYL PYRIDINIUM CHLORIDE
- 696 N-CETYLYL 2-METHYL PYRIDINIUM IODIDE
- 694 N-CETYLYL 3-METHYL PYRIDINIUM CHLORIDE
- 697 N-CETYLYL 3-METHYL PYRIDINIUM IODIDE
- 695 N-CETYLYL 4-METHYL PYRIDINIUM CHLORIDE
- 698 N-CETYLYL 4-METHYL PYRIDINIUM IODIDE
- 304 CATOL 607 (N-/LAURYL-COLAMINO-FORMYL-METHYL/-PYRIDINIUM CHLORIDE)
- 560 EMULSOL 607L (N-(2-KETO-2-(2,-LAUROYLOXYETHYL AMINO))-ETHYL) PYRIDINIUM CHLORIDE
- 717 DODECYLQUINOLINIUM BROMIDE
- 491 DODECYL TROPYLUM BISULFATE
- 528 DODECYL TROPYLUM PERCHLORATE
- 460 DODECYL TROPYLUM MONOPHOSPHATE

Structural Indexes of Compounds with Keys to Compound Numbers—Continued

PART 4. *Polyoxyethylenes grouped by distribution of heads arranged by number of carbon atoms in longest hydrophobic tail and by heads.*

4A. POLYOXYETHYLENES — HOMOGENEOUS HEAD GROUPS

OE 1

- 461 BUTYL/OXYETHYLENE/1 ALCOHOL--BUTYL GLYCOL ETHER
HOMOGENEOUS HEAD GROUP
- 423 OCTYL/OXYETHYLENE/1 ALCOHOL--OCTYL GLYCOL ETHER
HOMOGENEOUS HEAD GROUP
- 207 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL HOMOGENOUS
HEAD GROUP
- 208 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL HOMOGENOUS
HEAD GROUP
- 103 HEXYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
- 104 OCTYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP
- 209 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL HOMOGENOUS
HEAD GROUP
- 107 DECYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENOUS HEAD GROUP

OE 4

- 380 HEXYL/OXYETHYLENE/4 ALCOHOL HOMOGENEOUS HEAD GROUP
- 210 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL HOMOGENOUS
HEAD GROUP
- 379 DECYL/OXYETHYLENE/4 ALCOHOL HOMOGENEOUS HEAD GROUP
- 381 HEXYL /OXYETHYLENE/5 ALCOHOL HOMOGENEOUS HEAD GROUP
- 211 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL HOMOGENOUS
HEAD GROUP
- 378 DECYL/OXYETHYLENE/5 ALCOHOL HOMOGENEOUS HEAD GROUP
- 377 DODECYL/OXYETHYLENE/ 5 ALCOHOL HOMOGENEOUS HEAD GROUP

OE 6

- 393 BUTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP
- 394 1-METHYL PROPYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD
GROUP
- 294 HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENOUS HEAD GROUP
- 395 2-ETHYL BUTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD
GROUP
- 105 OCTYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP
- 396 2-PROPYL PENTYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD
GROUP
- 212 P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL HOMOGENOUS
HEAD GROUP
- 108 DECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENOUS HEAD GROUP
- 397 2-BUTYL HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD
GROUP
- 110 DODECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENEOUS HEAD GROUP
- 289 TETRADECYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP
- 282 HEXADECYL /OXYETHYLENE/ 6 ALCOHOL HOMOGENEOUS HEAD
GROUP

OE 7

- 213 P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL HOMOGENOUS
HEAD GROUP
- 488 DODECYL/OXYETHYLENE/7 ALCOHOL HOMOGENEOUS HEAD GROUP
- 330 HEXADECYL/OXYETHYLENE/7 ALCOHOL HOMOGENEOUS HEAD GROUP
- 214 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL HOMOGENOUS
HEAD GROUP
- 106 OCTYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP
- 215 P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL HOMOGENOUS
HEAD GROUP
- 109 DECYL /OXYETHYLENE/ 9 ALCOHOL HOMOGENOUS HEAD GROUP
- 398 2-BUTYL HEXYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD
GROUP
- 331 HEXADECYL/OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP
- 489 DODECYL /OXYETHYLENE/9 ALCOHOL HOMOGENEOUS HEAD GROUP

OE 10

- 216 P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL HOMOGENOUS
HEAD GROUP
- 490 DODECYL/OXYETHYLENE/12 ALCOHOL HOMOGENEOUS HEAD GROUP
- 332 HEXADECYL/OXYETHYLENE/12 ALCOHOL HOMOGENEOUS HEAD
GROUP
- 333 HEXADECYL/OXYETHYLENE/15 ALCOHOL HOMOGENEOUS HEAD
GROUP
- 334 HEXADECYL/OXYETHYLENE/21 ALCOHOL HOMOGENEOUS HEAD
GROUP

4B. POLYOXYETHYLENES — REDUCED POLYDISPERSITY OF HEAD GROUPS

OE 4

- 325 DODECYL/OXYETHYLENE/4 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
- 312 TRIDECYL/OXYETHYLENE/5 ALCOHOL BRANCHED CHAIN, REDUCED
OE DISTRIBUTION
- 583 METHYL /OXYETHYLENE/ 6.0 DODECANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 586 METHYL /OXYETHYLENE/ 7.6 OCTANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 580 METHYL /OXYETHYLENE/ 7.0 DECANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 115 DODECYL /OXYETHYLENE/7 ALCOHOL REDUCED POLYDISPERSION
OF HEAD GROUPS
- 201 DECYL /OXYETHYLENE/8 OXYMETHYL REDUCED POLYDISPERSION
OF HEAD GROUPS
- 584 METHYL /OXYETHYLENE/ 8.4 DODECANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 313 TRIDECYL/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION

OE 10

- 317 T-OCTYL BENZENE/OXYETHYLENE/10 ALCOHOL REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 320 NONYL BENZENE/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
- 581 METHYL /OXYETHYLENE/ 10.3 DECANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 202 DECYL /OXYETHYLENE/11 OXYMETHYL REDUCED POLYDISPERSION
OF HEAD GROUPS
- 585 METHYL /OXYETHYLENE/ 11.2 DODECANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 578 METHYL /OXYETHYLENE/ 11.9 DECANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 204 DECYL /OXYETHYLENE/12 OXYMETHYL REDUCED POLYDISPERSION
OF HEAD GROUPS
- 205 DODECYL /OXYETHYLENE/12 OXYMETHYL REDUCED
POLYDISPERSION OF HEAD GROUPS
- 579 METHYL /OXYETHYLENE/ 12.5 DODECANOATE REDUCED
POLYDISPERSITY OF HEAD GROUPS

OE 14

- 326 DODECYL/OXYETHYLENE/14 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
- 328 OCTADECYL/OXYETHYLENE/14 ALCOHOL REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 314 TRIDECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
- 321 NONYL BENZENE/OXYETHYLENE/15 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
- 716 DODECYL /OXYETHYLENE/16 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
- 535 HEXADECYL/OXYETHYLENE/18 ALCOHOL

OE 20

- 318 T-OCTYL BENZENE/OXYETHYLENE/20 ALCOHOL REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 322 NONYL BENZENE/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
- 327 DODECYL/OXYETHYLENE/23 ALCOHOL REDUCED POLYDISPERSITY
OF HEAD GROUPS
- 315 TRIDECYL/OXYETHYLENE/20 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
- 319 T-OCTYL BENZENE/OXYETHYLENE/30 ALCOHOL REDUCED
POLYDISPERSITY OF HEAD GROUPS
- 323 NONYL BENZENE/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
- 721 NONYL BENZENE /OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN
DIALYSED
- 116 DODECYL /OXYETHYLENE/30 ALCOHOL REDUCED POLYDISPERSION
OF HEAD GROUPS

PART 4. *Polyoxyethylenes grouped by distribution of heads arranged by number of carbon atoms in longest hydrophobic tail and by heads*—Continued

316 TRIDECYL/OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
117 HEXADECYL /OXYETHYLENE/30 ALCOHOL REDUCED
POLYDISPERSION OF HEAD GROUPS
324 NYONYL BENZENE/OXYETHYLENE/50 ALCOHOL BRANCHED CHAIN,
REDUCED OE DISTRIBUTION
329 OCTADECYL/OXYETHYLENE/100 ALCOHOL REDUCED
POLYDISPERSITY OF HEAD GROUPS

4C. POLYOXYETHYLENES — NATURAL DISTRIBUTION OF
HEAD GROUPS

OE 1

217 P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
218 P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
219 P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
220 P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
515 ISO-OCTYL/OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
516 NYONYL/OXYETHYLENE/4 ALCOHOL BRANCHED CHAIN, NATURAL OE
DISTRIBUTION
519 DECYL/OXO-PROCESS/ /OXYETHYLENE/4 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
440 /OXYETHYLENE/4 SORBITAN MONOLAUARATE ISOMERS AND
DISTRIBUTED MULTIPLE OE CHAINS

OE 5

221 P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
153 NYONYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
143 DECYL /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
158 DODECYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
148 TRIDECYL /OXYETHYLENE/ 5.2 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
674 DIISOBUTYL BENZENE/OXYETHYLENE/6 ALCOHOL (DB) BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
222 P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
468 DODECYL/OXYETHYLENE/6.2 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
518 TRIMETHYLNONYL/OXYETHYLENE/6 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
223 P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
469 DODECYL/OXYETHYLENE/7.6 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS

OE 8

335 OCTYL BENZENE/OXYETHYLENE/8.5 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
224 P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
520 UNDECYL/OXYETHYLENE/8 ALCOHOL NATURAL DISTRIBUTION OF
HEAD GROUPS
441 /OXYETHYLENE/8 SORBITAN MONOLAUARATE ISOMERS AND
DISTRIBUTED MULTIPLE OE CHAINS
163 DODECYL /OXYETHYLENE/ 8 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
431 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/8.5 ALCOHOL
BRANCHED CHAIN, NATURAL OE DISTRIBUTION
206 TRITON X-100(P-T-OCTYL BENZENE/OXYETHYLENE/9)NATURAL
DISTRIBUTION OF HEAD GROUPS
225 P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
336 NYONYL BENZENE/OXYETHYLENE/9.5 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
154 NYONYL BENZENE /OXYETHYLENE/ 9.8 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
276 NONAETHYLENE GLYCOL MONODODECANOATE NATURAL
DISTRIBUTION OF HEAD GROUPS
465 TRIDECYL/OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF
HEAD GROUPS
466 TRIDECYL/SECONDARY/ /OXYETHYLENE/9 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS

OE 10

675 OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
226 P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
167 NYONYL BENZENE /OXYETHYLENE/ 10 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
337 NYONYL BENZENE/OXYETHYLENE/10.5 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
144 DECYL /OXYETHYLENE/ 10.0 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
517 DECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
442 /OXYETHYLENE/10 SORBITAN MONOLAUARATE ISOMERS AND
DISTRIBUTED MULTIPLE OE CHAINS
310 DODECYL/OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION OF
HEAD GROUPS
159 DODECYL BENZENE /OXYETHYLENE/ 10.2 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
470 DODECYL/OXYETHYLENE/10.6 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
521 TRIDECYL/OXYETHYLENE/10 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
432 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
149 TRIDECYL /OXYETHYLENE/ 10.1 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
435 OCTADECYL/OXYETHYLENE/10 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS

OE 11

463 NYONYL BENZENE/OXYETHYLENE/11 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
523 DECYL BENZENE PARA/OXYETHYLENE/11.0 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
522 DECYL BENZENE ORTHO/OXYETHYLENE/11.2 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
164 DODECYL /OXYETHYLENE/ 12 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
467 TRIDECYL/SECONDARY/ /OXYETHYLENE/12 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
437 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/12 ALCOHOL
NATURAL DISTRIBUTION OF HEAD GROUPS

OE 14

145 DECYL /OXYETHYLENE/ 14.9 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
524 HEXADECYL/OXYETHYLENE/14 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
464 NYONYL BENZENE/OXYETHYLENE/15 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
168 NYONYL BENZENE /OXYETHYLENE/ 15 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
155 NYONYL BENZENE /OXYETHYLENE/ 15.4 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION
160 DODECYL BENZENE /OXYETHYLENE/ 15.1 ALCOHOL BRANCHED
CHAIN. NATURAL OE DISTRIBUTION
150 TRIDECYL /OXYETHYLENE/ 15.2 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
433 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/15 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION

OE 16

227 P-T-OCTYL BENZENE /OXYETHYLENE/16 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS
582 METHYL /OXYETHYLENE/ 16.0 DECANOATE NATURAL
DISTRIBUTION OF HEAD GROUPS
471 DODECYL/OXYETHYLENE/15.8 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
438 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/16 ALCOHOL
NATURAL DISTRIBUTION OF HEAD GROUPS
165 DODECYL /OXYETHYLENE/ 18 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS

OE 20

151 TRIDECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION
436 OCTADECYL/OXYETHYLENE/20 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS
439 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/20 ALCOHOL
NATURAL DISTRIBUTION OF HEAD GROUP
156 NYONYL BENZENE /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION

PART 4. *Polyoxyethylenes grouped by distribution of heads arranged by number of carbon atoms in longest hydrophobic tail and by heads—Continued*

- | | |
|--|---|
| 146 DECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION | 170 DODECYL /OXYETHYLENE/ 28 ALCOHOL REDUCED DISTRIBUTION
OF HEAD GROUPS |
| 443 /OXYETHYLENE/20 SORBITAN MONOLAURATE ISOMERS AND
DISTRIBUTED MULTIPLE OE CHAINS | 147 DECYL /OXYETHYLENE/ 29.4 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION |
| 472 DODECYL/OXYETHYLENE/20.4 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS | 169 NYONYL BENZENE /OXYETHYLENE/ 30 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION |
| 161 DODECYL BENZENE /OXYETHYLENE/ 20.1 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION | 157 NYONYL BENZENE /OXYETHYLENE/ 30.2 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION |
| 434 TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/22 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION | 162 DODECYL BENZENE /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED
CHAIN, NATURAL OE DISTRIBUTION |
| 166 DODECYL /OXYETHYLENE/ 23 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS | 152 TRIDECYL /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION |
| OE-25 | 474 DODECYL/OXYETHYLENE/38.6 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS |
| 444 /OXYETHYLENE/25 SORBITAN MONOLAURATE ISOMERS AND
DISTRIBUTED MULTIPLE OE CHAINS | 228 P-T-OCTYL BENZENE /OXYETHYLENE/40 ALCOHOL NATURAL
DISTRIBUTION OF HEAD GROUPS |
| 473 DODECYL/OXYETHYLENE/27.1 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS | 475 DODECYL/OXYETHYLENE/67.3 ALCOHOL NATURAL DISTRIBUTION
OF HEAD GROUPS |
| | 338 NYONYL BENZENE/OXYETHYLENE/100 ALCOHOL BRANCHED CHAIN,
NATURAL OE DISTRIBUTION |

Structural Indexes of Compounds with Keys to Compound Numbers—Continued

PART 5. Ionics grouped by counterion and arranged by number of carbon atoms in longest hydrophobic tail and by heads.

5A. ANIONICS BY COUNTERION

HYDROGEN

- 428 PERFLUORO ACETIC ACID
- 429 PERFLUORO PROPIONIC ACID
- 43 BUTYRIC ACID
- 452 3-4-DICHLOROPERFLUORO BUTYRIC/KEL-F/ACID
- 430 PERFLUORO BUTYRIC ACID
- 700 HEXANOIC ACID
- 453 3-5-6-TRICHLOROPERFLUORO HEXANOIC/KEL-F/ACID
- 416 PERFLUORO HEXANOIC ACID
- 374 DODECAFLUOROHEPTANOIC ACID H/CF₂/6COOH
- 529 OCTANOIC ACID
- 417 PERFLUORO OCTANOIC ACID
- 375 HEXADECAFLUORONONANOIC ACID H/CF₂/8COOH
- 611 ALPHAPHOSPHONO PELARGONIC ACID
- 530 DECANOIC ACID
- 455 PERFLUORO DECANOIC ACID
- 612 ALPHAPHOSPHONO DECANOIC ACID
- 536 NONYL SULFONIC ACID
- 272 TRI-ISOPROPYL BENZENE SULFONIC ACID
- 531 DODECANOIC ACID
- 602 ALPHA SULFO LAURIC ACID
- 613 ALPHAPHOSPHONO DODECANOIC ACID
- 537 UNDECYL SULFONIC ACID

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- 705 DODECYL SULFURIC ACID
- 200 DODECYL SULFONIC ACID
- 247 DODECANE 1-HYDROXY 2-SULFONIC ACID
- 532 TETRADECANOIC ACID
- 614 ALPHAPHOSPHONO TETRADECANOIC ACID
- 189 ALPHA SULFOMYRISTIC ACID
- 242 TRIDECAINE 1-SULFONIC ACID
- 243 TETRADECANE 1-SULFONIC ACID
- 248 TETRADECANE 1-HYDROXY 2-SULFONIC ACID
- 533 HEXADECANOIC ACID
- 190 ALPHA SULFOPALMITIC ACID
- 615 ALPHAPHOSPHONO HEXADECANOIC ACID
- 244 PENTADECANE 1-SULFONIC ACID
- 245 HEXADECANE 1-SULFONIC ACID
- 249 HEXADECANE 1-HYDROXY 2-SULFONIC ACID
- 534 OCTADECANOIC ACID
- 191 ALPHA SULFOSTEARIC ACID
- 551 PHENYL SULFOSTEARIC ACID
- 552 TOLYL SULFOSTEARIC ACID
- 553 XYLYL SULFOSTEARIC ACID
- 246 HEPTADECANE 1-SULFONIC ACID
- 232 OCTADECANE 1-SULFONIC ACID
- 250 OCTADECANE 1-HYDROXY 2-SULFONIC ACID

SODIUM

- 699 SODIUM BUTYRATE
- 681 SODIUM DIBUTYL BENZENE SULFONATE
- 682 SODIUM DIBUTYL NAPHTHALENE SULFONATE /NEKAL/
- 484 SODIUM PENTANOATE/VALERATE/
- 257 SODIUM DI-N-BUTYL SULFOSUCCINATE
- 261 SODIUM DI-ISOBUTYL SULFOSUCCINATE
- 445 SODIUM PARA-BIS-/N-BUTYL/ BENZENE SULFONATE
- 556 ARESKAP 100 /MONOBUTYL PHENYLPHENOL SODIUM MONOSULFONATE/
- 557 ARESKET 300 /MONOBUTYL BIPHENYL SODIUM MONOSULFONATE/
- 558 ARESKLENE 400 /DIBUTYL PHENYLPHENOL DISODIUMDISULFONATE
- 485 SODIUM HEXANOATE/CAPROATE/
- 258 SODIUM DI-N-AMYL SULFOSUCCINATE
- 343 SODIUM PENTANE SULFONATE

C-6

- 486 SODIUM HEPTANOATE
- 259 SODIUM DI-N-HEXYL SULFOSUCCINATE
- 352 SODIUM DI-1-METHYLISOAMYL SULFOSUCCINATE
- 344 SODIUM HEXANE SULFONATE
- 501 SODIUM HEXYL BENZENE SULFONATE

- 446 SODIUM PARA-BIS-/N-HEXYL/ BENZENE SULFONATE
- 476 SODIUM OCTANOATE
- 502 SODIUM HEPTYL BENZENE SULFONATE

C-8

- 487 SODIUM NONANOATE
- 260 SODIUM DI-N-OCTYL SULFOSUCCINATE
- 262 SODIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE
- 6 SODIUM ALPHA SULFOPELARGONIC ACID
- 25 SODIUM ETHYL ALPHA SULFOPELARGONATE
- 26 SODIUM AMYL ALPHA SULFOPELARGONATE
- 27 SODIUM HEXYL ALPHA SULFOPELARGONATE
- 28 SODIUM HEPTYL ALPHA SULFOPELARGONATE
- 7 SODIUM OCTYL ALPHA SULFOPELARGONATE
- 31 SODIUM 2 OCTYL ALPHA SULFOPELARGONATE
- 32 SODIUM 2 ETHYLHEXYL ALPHA SULFOPELARGONATE
- 35 SODIUM H/CF₂CF₂/3CH₂ ALPHA SULFOPELARGONATE
- 605 SODIUM AMYL ALPHAPHOSPHONO PELARGONATE
- 181 SODIUM OCTYL 1-SULFONATE
- 2 SODIUM OCTYL 1 SULFATE
- 66 SODIUM OCTYL 2 SULFATE
- 49 SODIUM P OCTYL BENZENE SULFONATE
- 503 SODIUM OCTYL BENZENE SULFONATE
- 172 SODIUM 2-N-OCTYL BENZENE SULFONATE
- 510 SODIUM 2-ETHYL-HEXYL BENZENE SULFONATE
- 676 SODIUM 1,1,3,3-TETRAMETHYL BUTYL BENZENE SULFONATE
- 677 SODIUM DI-/1,1,3,3-TETRAMETHYL BUTYL/BENZENESULFONATE
- 447 SODIUM PARA-BIS-/N-OCTYL/ BENZENE SULFONATE

C-9

- 299 SODIUM DECANOATE
- 29 SODIUM NONYL ALPHA SULFOPELARGONATE
- 616 MONOSODIUM ALPHAPHOSPHONO DECANOATE
- 606 SODIUM AMYL ALPHAPHOSPHONO CAPRATE
- 136 SODIUM ALPHA DIMETHYL AMINO CAPRATE
- 295 SODIUM NONYL 1-SULFATE
- 493 SODIUM P-NONYL BENZENE SULFONATE
- 504 SODIUM NONYL BENZENE SULFONATE
- 271 SODIUM TRI-ISO-PROPYL BENZENE SULFONATE
- 138 SODIUM NONYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN

C-10

- 30 SODIUM DECYL ALPHA SULFOPELARGONATE
- 33 SODIUM /OXO/ DECYL ALPHA SULFOPELARGONATE
- 3 SODIUM DECYL 1 SULFATE
- 15 SODIUM DECYL 2 SULFATE
- 182 SODIUM DECYL 1-SULFONATE
- 50 SODIUM P DECYL BENZENE SULFONATE
- 505 SODIUM DECYL BENZENE SULFONATE
- 173 SODIUM 2-N-DECYL BENZENE SULFONATE
- 511 SODIUM 2-PROPYL-HEPTYL BENZENE SULFONATE
- 140 SODIUM DECYL BENZENE SULFONATE BRANCHED HYDROCARBON CHAIN
- 561 SANTOMERSE D /SODIUM DECYLBENZENE SULFONATE/

C-11

- 273 SODIUM DODECANOATE
- 527 SODIUM UNDECANE-3-CARBOXYLATE
- 235 SODIUM ALPHA SULFO LAURIC ACID
- 603 SODIUM PROPYL ALPHA SULFO LAURATE
- 607 SODIUM ISOPROPYL ALPHAPHOSPHONO LAURATE
- 620 DISODIUM ALPHAPHOSPHONO DODECANOATE
- 617 MONOSODIUM ALPHAPHOSPHONO DODECANOATE
- 311 SODIUM UNDECYL 1-SULFATE
- 72 SODIUM UNDECYL 3 SULFATE
- 80 SODIUM UNDECYL 6 SULFATE
- 639 SODIUM UNDECYL THIOSULFATE
- 462 SODIUM MONOLAURIN SULFATE
- 418 SODIUM UNDECYL SULFONATE
- 45 SODIUM P 1 METHYL DECYL BENZENE SULFONATE
- 678 SODIUM 1,1-DIMETHYLNONYL BENZENE SULFONATE
- 679 SODIUM 3-BUTYL HEPTYL BENZENE SULFONATE

PART 5. Ionics grouped by counterion and arranged by number of carbon atoms in longest hydrophobic tail
and by heads – Continued

C-12

34 SODIUM DODECYL ALPHA SULFOPELARGONATE
 1 SODIUM DODECYL 1 SULFATE
 67 SODIUM DODECYL 2 SULFATE
 562 SODIUM DODECENYL SULFATE
 636 SODIUM DODECYL THIOSULFATE
 541 SODIUM DODECYL MONO-OXYETHYLENE SULFATE
 542 SODIUM DODECYL DIOXYETHYLENE SULFATE
 113 SODIUM DODECYL TRI-OXYETHYLENE SULFATE
 543 SODIUM DODECYL TETRA-OXYETHYLENE SULFATE
 114 SODIUM DODECYL /OXYETHYLENE/17.5 SULFATE
 597 SODIUM DODECYL MONO-OXYPROPYL SULFATE
 179 SODIUM DODECANE 1-SULFONATE
 175 SODIUM DODECANE 2-SULFONATE
 238 SODIUM DODECANE 1-HYDROXY 2-SULFONATE
 51 SODIUM P DODECYL BENZENE SULFONATE
 506 SODIUM DODECYL BENZENE SULFONATE
 171 SODIUM 2-N-DODECYL BENZENE SULFONATE
 301 SODIUM 3-N-DODECYL BENZENE SULFONATE
 302 SODIUM 4-N-DODECYL BENZENE SULFONATE
 514 SODIUM 6-N-DODECYL BENZENE SULFONATE
 512 SODIUM 2-BUTYL-OCTYL BENZENE SULFONATE
 492 SODIUM TETRAPROPYLENE/1-3-5-7-TETRAMETHYL-
 OCTYL/BENZENE SULFONATE
 680 SODIUM 2-PROPYL-4-METHYL-OCTYL BENZENE SULFONATE
 139 SODIUM DODECYL BENZENE SULFONATE BRANCHED HYDROCARBON
 CHAIN
 554 SANTOMERSE 3 /SODIUM DODECYL BENZENE SULFONATE/

C-13

298 SODIUM TETRADECANOATE
 236 SODIUM ALPHA SULFO MYRISTIC ACID
 604 SODIUM METHYL ALPHA SULFO MYRISTATE
 233 DISODIUM ALPHA SULFO MYRISTATE
 618 MONOSODIUM ALPHAPHOSPHONO TETRADECANOATE
 608 SODIUM METHYL ALPHAPHOSPHONO MYRISTATE
 621 DISODIUM ALPHAPHOSPHONO TETRADECANOATE
 624 TRISODIUM ALPHAPHOSPHONO TETRADECANOATE
 68 SODIUM TRIDECYL 2 SULFATE
 83 SODIUM TRIDECYL 7 SULFATE
 229 SODIUM TRIDECANE 1-SULFONATE
 46 SODIUM P 1 METHYL DODECYL BENZENE SULFONATE
 141 SODIUM TRIDECYL BENZENE SULFONATE BRANCHED HYDROCARBON
 CHAIN

C-14

4 SODIUM TETRADECYL 1 SULFATE
 16 SODIUM TETRADECYL 2 SULFATE
 73 SODIUM TETRADECYL 3 SULFATE
 17 SODIUM TETRADECYL 4 SULFATE
 77 SODIUM TETRADECYL 5 SULFATE
 525 SODIUM TETRADECYL 6-SULFATE
 84 SODIUM TETRADECYL 7 SULFATE
 526 SODIUM 2-DI-N-HEXYL ETHYL SULFATE
 544 SODIUM TETRADECYL MONO-OXYETHYLENE SULFATE
 545 SODIUM TETRADECYL DI-OXYETHYLENE SULFATE
 546 SODIUM TETRADECYL TRI-OXYETHYLENE SULFATE
 598 SODIUM TETRADECYL MONO-OXYPROPYL SULFATE
 599 SODIUM TETRADECYL DI-OXYPROPYL SULFATE
 183 SODIUM TETRADECYL 1-SULFONATE
 176 SODIUM OCTADECANE 2-SULFONATE
 239 SODIUM TETRADECANE 1-HYDROXY 2-SULFONATE
 507 SODIUM TETRADECYL BENZENE SULFONATE
 174 SODIUM 2-N-TETRADECYL BENZENE SULFONATE
 513 SODIUM 2-AMYL-NONYL BENZENE SULFONATE

C-15

300 SODIUM HEXADECANOATE
 237 SODIUM ALPHA SULFO PALMITIC ACID
 36 SODIUM METHYL ALPHA SULFOPALMITATE
 192 SODIUM ETHYL ALPHA SULFOPALMITATE
 193 SODIUM PROPYL ALPHA SULFOPALMITATE
 234 DISODIUM ALPHA SULFO PALMITATE
 197 DISODIUM 2 SULFOETHYL ALPHA SULFOPALMITATE
 619 MONOSODIUM ALPHAPHOSPHONO HEXADECANOATE
 609 SODIUM ISOPROPYL ALPHAPHOSPHONO PALMITATE
 622 DISODIUM ALPHAPHOSPHONO HEXADECANOATE
 625 TRISODIUM ALPHAPHOSPHONO HEXADECANOATE
 69 SODIUM PENTADECYL 2 SULFATE
 74 SODIUM PENTADECYL 3 SULFATE
 78 SODIUM PENTADECYL 5 SULFATE
 85 SODIUM PENTADECYL 8 SULFATE
 230 SODIUM PENTADECANE 1-SULFONATE
 47 SODIUM P 1 METHYL TETRADECYL BENZENE SULFONATE

142 SODIUM PENTADECYL BENZENE SULFONATE BRANCHED
 HYDROCARBON CHAIN

C-16

5 SODIUM HEXADECYL 1 SULFATE
 75 SODIUM HEXADECYL 4 SULFATE
 81 SODIUM HEXADECYL 6 SULFATE
 86 SODIUM HEXADECYL 8 SULFATE
 52 SODIUM HEXADECYL MONO OXYETHYLENE SULFATE
 53 SODIUM HEXADECYL DI OXYETHYLENE SULFATE
 54 SODIUM HEXADECYL TRI OXYETHYLENE SULFATE
 55 SODIUM HEXADECYL TETRA OXYETHYLENE SULFATE
 600 SODIUM HEXADECYL MONO-OXYPROPYL SULFATE
 184 SODIUM HEXADECYL 1-SULFONATE
 177 SODIUM HEXADECANE 2-SULFONATE
 240 SODIUM HEXADECANE 1-HYDROXY 2-SULFONATE
 508 SODIUM HEXADECYL BENZENE SULFONATE

C-17

448 SODIUM OCTADECANOATE /STEARATE/
 263 SODIUM OLEATE /CIS-9-OCTADECENOATE/
 264 SODIUM ELAIDATE /TRANS-9-OCTADECENOATE/
 9 SODIUM ALPHA SULFOSTEARIC ACID
 194 SODIUM METHYL ALPHA SULFOSTEARATE
 195 SODIUM ETHYL ALPHA SULFOSTEARATE
 196 SODIUM PROPYL ALPHA SULFOSTEARATE
 199 SODIUM ISOPROPYL ALPHA SULFOSTEARATE
 10 DISODIUM ALPHA SULFOSTEARATE
 198 DISODIUM 2 SULFOETHYL ALPHA SULFOSTEARATE
 11 SODIUM ALPHA SULFO PHENYL STEARIC ACID
 8 DISODIUM ALPHA SULFOPHENYLSTEARATE
 12 SODIUM 9,10 DICHLORO ALPHA SULFOSTEARIC ACID
 13 DISODIUM 9,10 DICHLORO ALPHA SULFOSTEARATE
 14 DISODIUM 9,10 DIHYDROXY ALPHA SULFOSTEARATE
 610 SODIUM METHYL ALPHAPHOSPHONO STEARATE
 623 DISODIUM ALPHAPHOSPHONO OCTADECANOATE
 626 TRISODIUM ALPHAPHOSPHONO OCTADECANOATE
 70 SODIUM HEPTADECYL 2 SULFATE
 87 SODIUM HEPTADECYL 9 SULFATE
 231 SODIUM HEPTADECANE 1-SULFONATE
 48 SODIUM P 1 METHYL HEXADECYL BENZENE SULFONATE

C-18

64 SODIUM OCTADECYL 1 SULFATE
 71 SODIUM OCTADECYL 2 SULFATE
 76 SODIUM OCTADECYL 4 SULFATE
 82 SODIUM OCTADECYL 6 SULFATE
 61 SODIUM OLEYL/CIS 9 OCTADECENOYL/ SULFATE
 62 SODIUM ELAIDYL/TRANS 9 OCTADECENOYL/SULFATE
 63 SODIUM 9,10 DICHLORO OCTADECYL SULFATE
 56 SODIUM OCTADECYL MONO OXYETHYLENE SULFATE
 547 SODIUM OLEYL MONO-OXYETHYLENE SULFATE
 57 SODIUM OCTADECYL DI OXYETHYLENE SULFATE
 548 SODIUM OLEYL DI-OXYETHYLENE SULFATE
 58 SODIUM OCTADECYL TRI OXYETHYLENE SULFATE
 549 SODIUM OLEYL TRI-OXYETHYLENE SULFATE
 59 SODIUM OCTADECYL TETRA OXYETHYLENE SULFATE
 601 SODIUM OCTADECYL MONO-OXYPROPYL SULFATE
 349 SODIUM OCTADECANE 1-SULFONATE
 178 SODIUM OCTADECANE 2-SULFONATE
 419 SODIUM ALPHA-HEPTYL UNDECYL SULFONATE
 241 SODIUM OCTADECANE 1-HYDROXY 2-SULFONATE
 509 SODIUM OCTADECYL BENZENE SULFONATE
 79 SODIUM NONADECYL 5 SULFATE
 88 SODIUM 1 NONYL DECYL SULFATE
 683 SODIUM EICOSYLBENZENE SULFONATE
 89 SODIUM 1 TETRADECYL PENTADECYL SULFATE

POTASSIUM

188 POTASSIUM HEXANOATE
 701 POTASSIUM PERFLUOROHEXANOATE
 296 POTASSIUM HEPTANOATE
 704 POTASSIUM 4-HEXYL RESORCINOLATE
 44 POTASSIUM OCTANOATE
 456 POTASSIUM PERFLUORO OCTANOATE
 350 POTASSIUM NONANOATE
 420 POTASSIUM 1-1-2-OCTANE TRICARBOXYLATE
 90 POTASSIUM DECANOATE
 702 POTASSIUM PERFLUORODECANOATE
 668 DIPOTASSIUM OCTYL MALONATE
 297 POTASSIUM UNDECANOATE
 425 POTASSIUM 1-1-2-DECANE TRICARBOXYLATE
 91 POTASSIUM DODECANOATE
 669 DIPOTASSIUM DECYL MALONATE

PART 5. *Ionics grouped by counterion and arranged by number of carbon atoms in longest hydrophobic tail and by heads*—Continued

C-12

- 351 POTASSIUM TRIDECANOATE
- 421 POTASSIUM 1-1-2-DODECANE TRICARBOXYLATE
- 632 POTASSIUM N-DODECYL BETA-ALANINATE
- 634 POTASSIUM DODECYL SULFATE
- 40 POTASSIUM DODECYL 1 SULFONATE
- 92 POTASSIUM TETRADECANOATE
- 670 DIPOTASSIUM DODECYL MALONATE
- 426 POTASSIUM 1-1-2-TETRADECANE TRICARBOXYLATE
- 185 POTASSIUM HEXADECANOATE
- 671 DIPOTASSIUM TETRADECYL MALONATE
- 422 POTASSIUM 1-1-2-HEXADECANE TRICARBOXYLATE
- 408 POTASSIUM HEXADECANE 1-SULFONATE
- 256 POTASSIUM STEARATE
- 305 POTASSIUM OLEATE/ CIS-9-OCTADECENOATE/
- 629 POTASSIUM ELAIDATE/TRANS-9-OCTADECENOATE/
- 255 POTASSIUM 9,10 DIHYDROXY STEARATE
- 630 POTASSIUM RICINOLEATE/12 HYDROXY OLEATE/
- 631 POTASSIUM RICINELAIDATE/12 HYDROXY ELAIDATE/
- 672 DIPOTASSIUM HEXADECYL MALONATE
- 673 DIPOTASSIUM OCTADECYL MALONATE
- 494 POTASSIUM DILINOLEATE

OTHER MONOVALENT METALS

- 111 LITHIUM DODECYL 1 SULFATE
- 635 LITHIUM DODECYL SULFONATE
- 637 LITHIUM TETRADECYL SULFATE
- 638 LITHIUM HEXADECYL SULFATE
- 627 CESIUM DODECANOATE
- 23 SILVER DODECYL 1 SULFATE

POLYVALENT METALS

- 339 MAGNESIUM HEXANE SULFONATE
- 340 MAGNESIUM OCTANE SULFONATE
- 341 MAGNESIUM DECANE SULFONATE
- 568 MAGNESIUM DODECYL SULFATE
- 342 MAGNESIUM DODECANE SULFONATE
- 24 CALCIUM DODECYL 1 SULFATE
- 569 STRONTIUM DODECYL SULFATE
- 573 CUPRIC DODECYL SULFATE
- 576 CUPRIC TETRADECYL SULFATE
- 577 CUPRIC HEXADECYL SULFATE
- 572 COBALTOUS DODECYL SULFATE
- 574 ZINC DODECYL SULFATE
- 571 MANGANESE DODECYL SULFATE
- 570 LEAD DODECYL SULFATE
- 575 NICKEL DODECYL SULFATE

AMMONIUM

- 355 AMMONIUM DODECAFLUOROHEPTANOATE H/CF2/6COONH4
- 286 AMMONIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE
- 372 AMMONIUM HEXADECALUORONANOATE H/CF2/8 COO NH4
- 373 AMMONIUM EICOSAFLUOROUNDECANOATE H/CF2/10 COO NH4
- 386 AMMONIUM DODECYL SULFATE
- 387 METHYLAMMONIUM DODECYL SULFATE
- 388 ETHYLAMMONIUM DODECYL SULFATE
- 389 BUTYLAMMONIUM DODECYL SULFATE
- 303 DI-ISOPROPYLAMMONIUM CAPRYLATE
- 596 TETRAHYDROANACARDOL AMMONIUM MONOSULFONATE
- 409 TRIETHANOLAMMONIUM DODECYL SULFATE
- 60 TRIETHANOL AMMONIUM HEXADECYL SULFATE
- 65 TRIETHANOLAMMONIUM OCTADECYL 1 SULFATE
- 284 HEXANOLAMINE-CH3CH/OH/CH2C/CH3/2NH2-OCTANOATE
- 285 HEXANOLAMINE-CH3CH/OH/CH2C/CH3/2NH2-OLEATE
- 283 HEXANOLAMINE-CH3CH/OH/CH2C/CH3/2NH2-ELAIDATE

QUATERNARIES

- 112 TETRAMETHYL AMMONIUM DODECYL 1 SULFATE
- 382 ETHYL TRIMETHYLAMMONIUM DODECYL SULFATE
- 383 BUTYL TRIMETHYLAMMONIUM DODECYL SULFATE
- 277 BENZYL TRIMETHYL AMMONIUM DODECANOATE
- 720 1-6-DITRIMETHYLAMMONIUM-HEXANE/DODECYL SULFATE/2
- 718 TETRAETHYLAMMONIUM DODECYL SULFATE
- 719 TETRABUTYLAMMONIUM DODECYL SULFATE
- 410 MORPHOLINIUM DODECYL SULFATE

SURFACTANT

- 390 HEXYLAMMONIUM DODECYL SULFATE
- 391 OCTYLAMMONIUM DODECYL SULFATE
- 640 HEXYL TRIMETHYLAMMONIUM HEXANE SULFATE
- 644 HEXYL TRIMETHYLAMMONIUM HEXANE SULFONATE
- 641 HEXYL TRIMETHYLAMMONIUM OCTANE SULFATE
- 384 HEXYL TRIMETHYLAMMONIUM DODECYL SULFATE

- 347 OCTYL TRIMETHYLAMMONIUM OCTYL SULFATE
- 287 OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE
- 642 OCTYL TRIMETHYLAMMONIUM DECANE SULFATE
- 353 OCTYL TRIMETHYLAMMONIUM DECANE SULFONATE
- 385 OCTYL TRIMETHYLAMMONIUM DODECYL SULFATE
- 346 DECYL TRIMETHYLAMMONIUM DECYL SULFATE
- 288 DECYL TRIMETHYLAMMONIUM DECANESULFONATE
- 280 DECYL TRIMETHYL AMMONIUM DODECYL SULFATE
- 643 DODECYL TRIMETHYLAMMONIUM OCTANE SULFATE
- 281 DODECYL TRIMETHYLAMMONIUM DODECYL SULFATE

5B. CATIONICS BY COUNTERIONS

HYDROXYL

- 706 PERFLUORO PROPYLAMINE
- 708 HEXYLAMINE

CHLORIDE

- 707 PERFLUORO PROPYLAMINE HYDROCHLORIDE
- 709 HEXYLAMINE HYDROCHLORIDE
- 354 HEXYL BENZYL DIMETHYLAMMONIUM CHLORIDE
- 392 OCTYLAMMONIUM CHLORIDE
- 483 DIOCTYL DIMETHYL AMMONIUM CHLORIDE
- 135 OCTYL C BETAINE HYDROCHLORIDE
- 451 PARA DI-ISOBUTYLPHENOXYETHOXETHYL DIMETHYLBENZYL AMMONIUM CHLORIDE/HYAMINE 1622/
- 359 OCTYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 137 ALPHA DIMETHYLAMINO CAPRIC ACID HYDROCHLORIDE

C-10

- 37 DECYLAMMONIUM CHLORIDE
- 203 DECYL TRIMETHYL AMMONIUM CHLORIDE
- 356 DECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
- 360 DECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 304 CATOL 607 (N-/LAURYL-COLAMINO-FORMYL-METHYL/- PYRIDINIUM CHLORIDE)

C-12

- 38 DODECYL AMMONIUM CHLORIDE
- 449 DODECYLMETHYL AMMONIUM CHLORIDE
- 450 DODECYLDIMETHYL AMMONIUM CHLORIDE
- 41 DODECYL TRIMETHYL AMMONIUM CHLORIDE
- 399 DODECYL DIMETHYL ETHYLAMMONIUM CHLORIDE
- 345 DIDODECYL DIMETHYLAMMONIUM CHLORIDE
- 403 DODECYL DIMETHYLPHENYLAMMONIUM CHLORIDE
- 279 DODECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
- 404 DODECYL DIMETHYL 2-PHENYLETHYLAMMONIUM CHLORIDE C6H5CH2CH2/N/CH3/2/C12H25
- 407 DODECYL DIMETHYL 3-PHENYLPROPYLAMMONIUM CHLORIDE C6H5CH2CH2/N/CH3/2/C12H25
- 370 DODECYL 3-4-METHYLENEDIOXYBENZYL DIMETHYLAMMONIUM CHLORIDE
- 371 DODECYL 3-4-DIMETHOXYBENZYL DIMETHYLAMMONIUM CHLORIDE
- 361 DODECYL 4-NITROBENZYL DIMETHYL AMMONIUM CHLORIDE
- 368 DODECYL 2-HYDROXY-5-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 365 DODECYL 2-CHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 367 DODECYL 4-CHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 366 DODECYL 2-4-DICHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 369 DODECYL 3-4-DICHLOROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 406 DODECYL DIMETHYL META-TRIFLUOROMETHYL BENZYL AMMONIUM CHLORIDE CF3C6H4CH2/N/CH3/2/C12H25
- 400 DODECYL METHYL DIETHYLAMMONIUM CHLORIDE
- 405 DODECYL METHYLETHYLAMMONIUM CHLORIDE
- 401 DODECYL TRIETHYLAMMONIUM CHLORIDE
- 22 DIMETHYL DODECYL AMINE OXIDE HYDROCHLORIDE
- 633 N-DODECYL BETA-ALANINE HYDROCHLORIDE
- 124 DODECYL N BETAINE HYDROCHLORIDE
- 278 DODECYL PYRIDINIUM CHLORIDE
- 528 DODECYL TROPYLUM PERCHLORATE
- 500 DODECYL TRI/2-HYDROXYETHYL/AMMONIUM CHLORIDE

C-13

- 402 TRIDECYL TRIMETHYLAMMONIUM CHLORIDE
- 39 TETRADECYL AMMONIUM CHLORIDE
- 42 TETRADECYL TRIMETHYL AMMONIUM CHLORIDE
- 357 TETRADECYL BENZYL DIMETHYLAMMONIUM CHLORIDE
- 362 TETRADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE
- 125 TETRADECYL N BETAINE HYDROCHLORIDE

C-16

- 186 HEXADECYL AMMONIUM CHLORIDE

PART 5. *Ionics grouped by counterion and arranged by number of carbon atoms in longest hydrophobic tail and by heads*—Continued

275	HEXADECYL DIMETHYLBENZYLAMMONIUM CHLORIDE	NITRATE
265	HEXADECYL TRIMETHYL AMMONIUM CHLORIDE	482 DODECYL AMMONIUM NITRATE
266	HEXADECYL DIMETHYL 2-HYDROXYETHYL AMMONIUM CHLORIDE	131 DODECYL TRIMETHYLAMMONIUM NITRATE
269	HEXADECYL DIMETHYL 2,3-DIHYDROXYPROPYLAMMONIUM CHLORIDE	654 OCTADECYL TRIMETHYLAMMONIUM NITRATE
363	HEXADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE	656 OCTADECYL PYRIDINIUM NITRATE
267	HEXADECYL DI-/2-HYDROXYETHYL/METHYL AMMONIUMCHLORIDE	
268	HEXADECYL TRI-/2-HYDROXYETHYL/ AMMONIUM CHLORIDE	
274	HEXADECYL PYRIDINIUM CHLORIDE	
693	N-CETYL 2-METHYL PYRIDINIUM CHLORIDE	BROMATE
694	N-CETYL 3-METHYL PYRIDINIUM CHLORIDE	129 DODECYL TRIMETHYL AMMONIUM BROMATE
695	N-CETYL 4-METHYL PYRIDINIUM CHLORIDE	666 HEXADECYL TRIBUTYLAMMONIUM BROMATE
C-18		658 OCTADECYL TRIMETHYLAMMONIUM BROMATE
187	OCTADECYL AMMONIUM CHLORIDE	662 OCTADECYL TRIETHYLAMMONIUM BROMATE
270	OCTADECYL TRIMETHYLAMMONIUM CHLORIDE	663 OCTADECYL TRIPROPYLAMMONIUM BROMATE
358	OCTADECYL BENZYL DIMETHYLAMMONIUM CHLORIDE	664 OCTADECYL TRIBUTYLAMMONIUM BROMATE
364	OCTADECYL 4-NITROBENZYL DIMETHYLAMMONIUM CHLORIDE	665 OCTADECYL TRIAMYLAMMONIUM BROMATE
655	OCTADECYL PYRIDINIUM CHLORIDE	
BROMIDE		
93	OCTYL TRIMETHYL AMMONIUM BROMIDE	IODATE
100	OCTYL PYRIDINIUM BROMIDE	127 DODECYL TRIMETHYL AMMONIUM IODATE
94	NONYL TRIMETHYL AMMONIUM BROMIDE	660 HEXADECYL PYRIDINIUM IODATE
95	DECYL TRIMETHYL AMMONIUM BROMIDE	661 OCTADECYL PYRIDINIUM IODATE
96	UNDECYL TRIMETHYL AMMONIUM BROMIDE	
101	UNDECYL PYRIDINIUM BROMIDE	SULFATE
628	DODECYLAMMONIUM BROMIDE	306 DECYL TRIMETHYLAMMONIUM SULFATE
97	DODECYL TRIMETHYL AMMONIUM BROMIDE	307 DODECYL TRIMETHYLAMMONIUM SULFATE
293	DODECYL DIMETHYL AMINE OXIDE HYDROBROMIDE	491 DODECYL TROPYLUM BISULFATE
290	DODECYL PYRIDINIUM BROMIDE	308 TETRADECYL TRIMETHYLAMMONIUM SULFATE
717	DODECYLQUINOLINIUM BROMIDE	309 HEXADECYL TRIMETHYLAMMONIUM SULFATE
C-14		
98	TETRADECYL TRIMETHYL AMMONIUM BROMIDE	PHOSPHATE
291	TETRADECYL TRIPROPYLAMMONIUM BROMIDE	460 DODECYL TROPYLUM MONOPHOSPHATE
102	TETRADECYL PYRIDINIUM BROMIDE	
99	HEXADECYL TRIMETHYLAMMONIUM BROMIDE	CARBOXYLATE
478	HEXADECYL DIMETHYLETHYLAMMONIUM BROMIDE	128 DODECYL TRIMETHYL AMMONIUM FORMATE
292	HEXADECYL TRIPROPYLAMMONIUM BROMIDE	659 OCTADECYL TRIMETHYLAMMONIUM FORMATE
427	HEXADECYL PYRIDINIUM BROMIDE	415 OCTADECYLAMMONIUM ACETATE
477	OCTADECYL TRIMETHYLAMMONIUM BROMIDE	411 DECYLAMMONIUM ACETATE
657	OCTADECYL PYRIDINIUM BROMIDE	412 DODECYLAMMONIUM ACETATE
IODIDE		413 TETRADECYLAMMONIUM ACETATE
458	DECYL PYRIDINIUM IODIDE	414 HEXADECYLAMMONIUM ACETATE
126	DODECYL TRIMETHYLAMMONIUM IODIDE	667 OCTADECYL TRIMETHYLAMMONIUM OXALATE
376	DODECYL PYRIDINIUM IODIDE	
479	TETRADECYL PYRIDINIUM IODIDE	
480	HEXADECYL PYRIDINIUM IODIDE	
696	N-CETYL 2-METHYL PYRIDINIUM IODIDE	SURFACTANT
697	N-CETYL-3-METHYL PYRIDINIUM IODIDE	391 OCTYLAMMONIUM DODECYL SULFATE
698	N-CETYL-4-METHYL PYRIDINIUM IODIDE	347 OCTYL TRIMETHYLAMMONIUM OCTYL SULFATE
481	OCTADECYL PYRIDINIUM IODIDE	287 OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE
FLUORIDE		642 OCTYL TRIMETHYLAMMONIUM DECANE SULFATE
130	DODECYL TRIMETHYL AMMONIUM FLUORIDE	353 OCTYL TRIMETHYLAMMONIUM DECANE SULFONATE
		385 OCTYL TRIMETHYLAMMONIUM DODECYL SULFATE
		346 DECYL TRIMETHYLAMMONIUM DECYL SULFATE
		280 DECYL TRIMETHYL AMMONIUM DODECYL SULFATE
		288 DECYL TRIMETHYLAMMONIUM DECANESULFONATE
		643 DODECYL TRIMETHYLAMMONIUM OCTANE SULFATE
		281 DODECYL TRIMETHYLAMMONIUM DODECYL SULFATE

Structural Indexes of Compounds with Keys to Compound Numbers—Continued

PART 6. Commercial names and ill defined structures arranged alphabetically.

6. COMMERCIAL AND ILL-DEFINED

556 ARESKAP 100 /MONOBUTYL PHENYLPHENOL SODIUM MONOSULFONATE/
557 ARESKET 300 /MONOBUTYL BIPHENYL SODIUM MONOSULFONATE/
558 ARESKLENE 400 /DIBUTYL PHENYLPHENOL DISODIUMDISULFONATE
566 AQUAREX D
559 CATOL 605 /(N-(2-KETO-2-(2,-LAUROYL OXYETHYLAMINO))ETHYL TRIMETHYLMONIUM CHLORIDE/
304 CATOL 607 (N-/LAURYL-COLAMINO-FORMYL-METHYL/ - PYRIDINIUM CHLORIDE)
459 ALKYL/15-17/ /OXYETHYLENE/ /19-23/ALCOHOL / CETOMACROGOL 1000/
565 DAXAD 11
674 DIISOBUTYL BENZENE/OXYETHYLENE/6 ALCOHOL (DB) BRANCHED CHAIN, NATURAL OE DISTRIBUTION
494 POTASSIUM DILINOLEATE
651 EMASOL 1120 /ALKYL POLYOXYETHYLENE SORBITAN ESTER/

652 EMASOL 1130 /ALKYL POLYOXYETHYLENE SORBITAN ESTER/
653 EMULGEN 120 /ALKYL POLYOXYETHYLENE ETHER/
560 EMULSOL 607L (N-(2-KETO-2-(2,-LAUROYLOXYETHYL AMINO))ETHYL) PYRIDINIUM CHLORIDE
451 PARA DI-ISOBUTYLPHENOXYETHOXYETHYL DIMETHYLBENZYL AMMONIUM CHLORIDE/HYAMINE 1622/
452 3-4-DICHLOROPERFLUORO BUTYRIC/KEL-F/ACID
453 3-5-6-TRICHLOROPERFLUORO HEXANOIC/KEL-F/ACID
454 3-5-7-8-TETRACHLOROPERFLUORO OCTANOIC/KEL-F/ACID
550 LAURIC ACID DIETHANOLAMINE CONDENSATE
564 NA OSR
682 SODIUM DIBUTYL NAPHTHALENE SULFONATE /NEKAL/
538 PLURONIC L62
539 RENEX 698
561 SANTOMERSE D /SODIUM DECYLBENZENE SULFONATE/
554 SANTOMERSE 3 /SODIUM DODECYL BENZENE SULFONATE/
567 SA-178
540 SIPONIC BC
555 TERGITOL TMN
563 ZEPHIRAN /COCONUT DIMETHYLBENZYL AMMONIUM CHLORIDE/

Table of Recommended and Selected Critical Micelle Concentrations

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation	
COMPOUND NO = 1 MOL WGT -	288.3	SODIUM DODECYL 1 SULFATE							
10	8.67 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
10	8.81 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
10.0	8.55 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	1		
15	8.51 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
15	8.43 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
20	8.47 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
20	8.25 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
25	2.330X10-1 D	BA	EQUIV CONDCTNCE GRAPH	WILL MYSE	55005	T	1		
	8.081X10-3 M					M			
25	2.340X10-1 D	BA	FOTOMTR SOLUBLZTN OROT	WILL MYSE	55005	T	1		
	8.116X10-3 M					M			
25	2.36 X10-1 D	BB	DEBYE PLT LIGHT SCATTER	WILL MYSE	55005	T	1		
	8.185X10-3 M					M			
25	8.39 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
25	8.1 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	GODD HIGH	55018	K	1		
25	2.324X10-1 D	BA	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T	1		
	8.061X10-3 M					M			
25	8.3 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MIUR MATS	57025	TA	1		
25	8.16 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
25	8.27 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	T	1		
25.0	8.27 X10-3 M	BA	SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T	1		
25	8.15 X10-3 M	AB	SURFACE TENSION LOG PLOT	ELWO MYSE	66007	T	1		
25	8.2 X10-3 M	AA	SURFACE TNSN LINEAR PLOT	ELWO MYSE	66007	T	1		
25	8.4 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	ELWO MYSE	66007	T	1		
30	8.44 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
30	8.23 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
35	8.57 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
35	8.4 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	1		
35	8.39 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
40	8.88 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
40	8.60 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
45	9.10 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
45	8.86 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
50	9.61 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D		
50	9.18 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	D		
55	9.95 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1		
55	9.8 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	1		
55.0	9.49 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	1		
55	9.61 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1		
60	1.016X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2		
65	1.091X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
70	1.14 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	3		
1. E-2 M	AG NO3								
5. E O H	DIOXANE	35	5.0 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	3
1.0 E 1 H	DIOXANE	15	6.73 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.5 E 1 H	DIOXANE	15	7.31 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.0 E 1 H	DIOXANE	15	9.03 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.5 E 1 H	DIOXANE	15	1.38 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2. E O H	DIOXANE	15	2.10 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
3. E O H	DIOXANE	25	8.01 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
5. E O H	DIOXANE	25	8.06 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
7. E O H	DIOXANE	25	7.7 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.0 E 1 H	DIOXANE	25	8.5 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.5 E 1 H	DIOXANE	25	9.0 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.0 E 1 H	DIOXANE	25	1.31 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.5 E 1 H	DIOXANE	25	2.12 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
5. E O H	DIOXANE	35	8.74 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.0 E 1 H	DIOXANE	35	1.05 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.5 E 1 H	DIOXANE	35	1.90 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.00 E 2 A	DEUTERIUM OXIDE	25.0	8.05 X10-3 M	BA	SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T	2
9.27 E 0 P	ETHANOL	5	5.51 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
3. E 0 P	ETHANOL	10.0	7.33 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
5. E 0 P	ETHANOL	10.0	6.63 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
6. E 0 P	ETHANOL	10.0	6.33 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
9. E 0 P	ETHANOL	10.0	5.55 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
9.27 E 0 P	ETHANOL	10	5.50 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
9.27 E 0 P	ETHANOL	15	5.54 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
9.27 E 0 P	ETHANOL	20	5.67 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
9.91 E 0 H	ETHANOL	20	5.65 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	WARD	40004	P	3
2.002E 1 H	ETHANOL	20	8.5 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	WARD	40004	P	3
9.27 E 0 P	ETHANOL	25	5.96 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
2.498E 1 P	ETHANOL	25	1.067X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
9.27 E 0 P	ETHANOL	30	6.33 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
2.498E 1 P	ETHANOL	30	1.146X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3
9.27 E 0 P	ETHANOL	35	6.72 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L	3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations – Continued

Additives		Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
2.498E 1 P	ETHANOL	35	1.310X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P	ETHANOL	40	7.19 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P	ETHANOL	40	1.496X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P	ETHANOL	45	7.72 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P	ETHANOL	45	1.656X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P	ETHANOL	50	8.30 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P	ETHANOL	50	1.831X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
3. E 0 P	ETHANOL	55.0	8.75 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
5. E 0 P	ETHANOL	55.0	8.40 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
6. E 0 P	ETHANOL	55.0	8.42 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9. E 0 P	ETHANOL	55.0	8.92 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P	ETHANOL	55	8.96 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
1.0 E 1 P	ETHANOL	55.0	9.28 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
1.5 E 1 P	ETHANOL	55.0	1.160X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.0 E 1 P	ETHANOL	55.0	1.505X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P	ETHANOL	60	9.70 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P	ETHANOL	60	2.170X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
1. E-1 M	NA BR	21	4.1 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	M
			1.42 X10-3 M						
1. E-2 M	NA CL	21	1.62 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	M
			5.619X10-3 M						
1. E-2 M	NA CL	21	1.52 X10-1 D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T 3	M
			5.272X10-3 M						
3. E-2 M	NA CL	21	9.2 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 1	M
			3.19 X10-3 M						
1. E-1 M	NA CL	21	4.3 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	M
			1.49 X10-3 M						
3. E-2 M	NA CL	25	9.00 X10-2 D	BB	FOTOMTR SOLUBLZTN OROT	WILL MYSE	55005	T 1	M
			3.121X10-3 M						
3. E-2 M	NA CL	25	9.03 X10-2 D	BA	SPECFC CONDCTNCE GRAPH	WILL MYSE	55005	T 1	M
			3.132X10-3 M						
3. E-2 M	NA CL	25	9.18 X10-2 D	BB	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T 1	M
			3.184X10-3 M						
1.00 E-1 M	NA CL	25	4.30 X10-2 D	BA	SPECFC CONDCTNCE GRAPH	WILL MYSE	55005	T 3	M
			1.491X10-3 M						
2. E-1 M	NA CL	25.0	9.0 X10-4 M	BA	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T 3	
1. E-2 M	NA F	21	1.61 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	M
			5.584X10-3 M						
1. E-1 M	NA F	21	4.2 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	M
			1.45 X10-3 M						
1. E-2 M	NA I	21	1.62 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	M
			5.619X10-3 M						
3. E-2 M	NA I	21	9.0 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 1	M
			3.12 X10-3 M						
1. E-1 M	NA I	21	4.0 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	M
			1.38 X10-3 M						
1. E-2 M	NA NO3	35	5.7 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T 3	
5.03 E 0 H	PROPANOL-1	00.5	4.7 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.03 E 0 H	PROPANOL-1	10.5	4.0 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.03 E 0 H	PROPANOL-1	25.2	3.8 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.03 E 0 H	PROPANOL-1	33.5	4.1 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.03 E 0 H	PROPANOL-1	40.1	4.4 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.03 E 0 H	PROPANOL-1	50.0	5.1 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.71 E 0 H	PROPANOL-2	00.5	5.7 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.71 E 0 H	PROPANOL-2	10.5	4.9 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.71 E 0 H	PROPANOL-2	25.2	4.3 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	T 3	
5.71 E 0 H	PROPANOL-2	33.5	4.5 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.71 E 0 H	PROPANOL-2	40.1	5.0 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.71 E 0 H	PROPANOL-2	50.0	5.7 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.06 E 0 H	PROPIONIC ACID	25.2	4.3 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3	
5.00 E 2 Y	PRESSURE	25	9.09 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3	
1.000E 3 Y	PRESSURE	25	9.45 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3	
1.500E 3 Y	PRESSURE	25	9.36 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3	
2.000E 3 Y	PRESSURE	25	9.09 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3	
3. E 0 D	SUCROSE	25	7.1 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	NAKA KAWA	64034	G 3	
1.078E 1 C	0003	25.0	8.32 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3	
2.006E 1 C	0003	25.0	8.54 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3	
2.025E 1 C	0003	25.0	8.54 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3	
3.478E 1 C	0003	25.0	9.88 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3	
4.912E 1 C	0003	25.0	1.115X10-2 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3	
5.998E 1 C	0003	25.0	1.281X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2	
7.100E 1 C	0003	25.0	1.498X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2	
8.466E 1 C	0003	25.0	1.91 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2	
9.495E 1 C	0003	25.0	2.65 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2	
1.25 E 1 C	0004	RM	5.11 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T 3	

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molal; Y – atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

	Additives	Temp. °C.	CMC	Qual. Mat. Q = Meth.	Method	Authors	Reference	Source	Evaluation
2.5 E 1 C	0004	RM	3.88 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
5.0 E 1 C	0004	RM	2.62 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
COMPOUND NO =	2 MOL WGT -	232.2	SODIUM OCTYL 1 SULFATE						
		10	1.421X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		15	1.367X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		20	1.33 X10-1 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	3
		20	1.357X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		21	3.10 X10 0 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	3
			1.335X10-1 M					M	
		25	1.303X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		30.0	1.30 X10-1 M	CB	VELOCITY OF SOUND	SHIG	65022	T	3
		30	1.318X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		35	1.342X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		40	1.363X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		40.0	1.36 X10-1 M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T	3
		45	1.381X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		50	1.434X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
		55	1.463X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	3
3. E-2 M	NA CL	21	2.80 X10 0 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	3
			1.205X10-1 M					M	
1. E-1 M	NA CL	21	2.37 X10 0 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	3
			1.020X10-1 M					M	
3. E-1 M	NA CL	21	1.60 X10 0 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	3
			6.890X10-2 M					M	
1. E 0 M	NA CL	21	8.0 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	3
			3.44 X10-2 M					M	
COMPOUND NO =	3 MOL WGT -	260.3	SODIUM DECYL 1 SULFATE						
		0	3.88 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		5	3.64 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		10	3.48 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D
		10	3.50 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	D
		15	3.41 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1
		15	3.39 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1
		20	3.35 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1
		20	3.31 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1
		25	3.32 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1
		25.0	3.26 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE KAPA	61005	T	1
		25.0	3.35 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	MYSE KAPA	61005	T	1
		25	3.27 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1
		25.0	3.32 X10-2 M	BA	SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T	1
		30	3.31 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1
		30	3.26 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1
		35	3.27 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1
		35	3.35 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1
		40	3.32 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1
		40	3.41 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	1
		45	3.38 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	D
		45	3.49 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D
		50	3.47 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	D
		50	3.64 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D
		55	3.59 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	D
		55	3.78 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D
		60	3.73 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		65	3.88 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
1.00 E 2 A	DEUTERIUM OXIDE	25.0	3.25 X10-2 M	BA	SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T	2
1.00 E-2 M	NA CL	25.0	3.02 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	MYSE KAPA	61005	L	3
1.00 E-1 M	NA CL	25.0	1.51 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	MYSE KAPA	61005	L	3
1.0 E 0 M	NA CL	25.0	2.77 X10-3 M	BB	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T	3
1.25 E 1 C	0001	RM	2.02 X10-2 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
2.5 E 1 C	0001	RM	1.56 X10-2 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
5.0 E 1 C	0001	RM	1.08 X10-2 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
5.00 E 1 C	0004	25	3.03 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	30	3.09 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	35	3.18 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	40	3.31 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	45	3.47 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	50	3.66 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	55	3.87 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	60	4.16 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	65	4.51 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	70	4.90 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	75	5.35 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations – Continued

	Additives	Temp. °C.	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO =	4 MOL WGT -		316.4 SODIUM TETRADECYL 1 SULFATE						
5. E 0 H	DIOXANE	25	2.05 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
1.0 E 1 H	DIOXANE	30	2.08 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
1.5 E 1 H	DIOXANE	35	2.13 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
2.0 E 1 H	DIOXANE	40	2.21 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	1
2.5 E 1 H	DIOXANE	40	2.21 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	1
3.0 E 1 H	DIOXANE	45	2.31 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
3.5 E 1 H	DIOXANE	50	2.43 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		55	2.58 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		60	2.77 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		65	2.99 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		70	3.22 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		75	3.50 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
COMPOUND NO =	5 MOL WGT -		344.4 SODIUM HEXADECYL 1 SULFATE						
5. E 0 H	DIOXANE	40	5.2 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.0 E 1 H	DIOXANE	40.0	5.8 X10-4 M	BC	SPECFC CONDCTNCE GRAPH	EVAN	56006	T	3
1.5 E 1 H	DIOXANE	40	1.27 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.0 E 1 H	DIOXANE	40	2.0 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.5 E 1 H	DIOXANE	40	2.8 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
3.0 E 1 H	DIOXANE	40	3.54 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
		40	4.3 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
		40	5.0 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
COMPOUND NO =	15 MOL WGT —		260.3 SODIUM DECYL 2 SULFATE						
		10	5.15 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		15	4.92 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		20	4.70 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		25	4.56 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		30	4.52 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		35	4.50 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		40	4.50 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		45	4.52 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		50	4.57 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		55	4.65 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		60	4.79 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		65	4.95 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
COMPOUND NO =	16 MOL WGT -		316.4 SODIUM TETRADECYL 2 SULFATE						
		25	3.27 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
		30	3.28 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
		35	3.31 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
		40	3.38 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		45	3.48 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		50	3.64 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		55	3.83 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		60	4.04 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		65	4.29 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		70	4.60 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
		75	5.00 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
COMPOUND NO =	17 MOL WGT -		316.4 SODIUM TETRADECYL 4 SULFATE						
		25	5.12 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		30	5.05 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		35	5.04 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		40.0	5.15 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T	3
		40	5.12 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		45	5.23 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		50	5.38 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		55	5.57 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		60	5.85 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		65	6.21 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		70	6.62 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
		75	7.11 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3
COMPOUND NO =	18 MOL WGT -		292.4 OCTYL BETA D GLUCOSIDE						
		25	2.5 X10-2 M	BB	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations – Continued

Concentration units: A - mol %; B - vol % solvent; C - mol % surfactant mixture; D - wt/vol %; E - % saturation; H - wt % solvent; I - mol % surfactant; K - normality

counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations – Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
1.619E 1 H	PROPANOL-2		25	1.04 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3
5.66 E 0 H	TERTIARY BUTANOL		25	7.9 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3
7.88 E 0 H	TERTIARY BUTANOL		25	7.4 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3
COMPOUND NO = 41	MOL WGT -	263.9	DODECYL TRIMETHYL AMMONIUM CHLORIDE						
5. E 2 Y	PRESSURE	25.0	2.03 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
1. E 3 Y	PRESSURE	25.0	2.09 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
1.5 E 3 Y	PRESSURE	25.0	2.11 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
2. E 3 Y	PRESSURE	25.0	2.04 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
3. E 3 Y	PRESSURE	25.0	1.98 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
4. E 3 Y	PRESSURE	25.0	1.87 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
5. E 3 Y	PRESSURE	25.0	1.83 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
5.00 E 3 Y	PRESSURE	25.0	1.81 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
COMPOUND NO = 42	MOL WGT -	292.0	TETRADECYL TRIMETHYL AMMONIUM CHLORIDE						
		25	4.47 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	HOYE MARM	61002	T 3	
COMPOUND NO = 44	MOL WGT -	182.3	POTASSIUM OCTANOATE						
3.3 E-2 W	K OH	15	3.72 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	20	3.55 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	25	3.45 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	30	3.30 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	35	3.13 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	40	3.05 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	45	3.10 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	50	3.18 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
3.3 E-2 W	K OH	55	3.31 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
COMPOUND NO = 45	MOL WGT -	334.4	SODIUM P 1 METHYL DECYL BENZENE SULFONATE						
		35	2.53 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 2	
COMPOUND NO = 46	MOL WGT -	362.4	SODIUM P 1 METHYL DODECYL BENZENE SULFONATE						
		35	7.2 X10-4 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
COMPOUND NO = 47	MOL WGT -	390.5	SODIUM P 1 METHYL TETRADECYL BENZENE SULFONATE						
		40	3.1 X10-4 W	BB	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
COMPOUND NO = 49	MOL WGT -	292.3	SODIUM P OCTYL BENZENE SULFONATE						
		35	1.47 X10-2 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
COMPOUND NO = 50	MOL WGT -	320.4	SODIUM P DECYL BENZENE SULFONATE						
		50	1.47 X10-2 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
COMPOUND NO = 51	MOL WGT -	348.4	SODIUM P DODECYL BENZENE SULFONATE						
		60	1.20 X10-3 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
COMPOUND NO = 66	MOL WGT -	232.2	SODIUM OCTYL 2 SULFATE						
		40.0	1.80 X10-1 M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3	
COMPOUND NO = 68	MOL WGT -	302.3	SODIUM TRIDECYL 2 SULFATE						
		40.0	6.50 X10-3 M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3	
COMPOUND NO = 72	MOL WGT -	274.3	SODIUM UNDECYL 3 SULFATE						
		40.0	2.89 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3	
COMPOUND NO = 73	MOL WGT -	316.4	SODIUM TETRADECYL 3 SULFATE						
		40.0	4.30 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3	
COMPOUND NO = 75	MOL WGT -	344.4	SODIUM HEXADECYL 4 SULFATE						
		40.0	1.72 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3	
COMPOUND NO = 78	MOL WGT -	330.4	SODIUM PENTADECYL 5 SULFATE						
		40.0	3.40 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3	

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

	Additives		Temp. °C.	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO =	80	MOL WGT -	274.3 40.0 8.3	SODIUM UNDECYL 6 SULFATE X10-2 M CB SPECFC CONDCTNCE GRAPH			EVAN	56006	T 3	
COMPOUND NO =	81	MOL WGT -	344.4 40.0 2.35	SODIUM HEXADECYL 6 SULFATE X10-3 M CB SPECFC CONDCTNCE GRAPH			EVAN	56006	T 3	
COMPOUND NO =	83	MOL WGT -	302.3 40.0 1.93	SODIUM TRIDECYL 7 SULFATE X10-2 M CB SPECFC CONDCTNCE GRAPH			EVAN	56006	T 3	
COMPOUND NO =	85	MOL WGT -	330.4 40.0 6.65	SODIUM PENTADECYL 8 SULFATE X10-3 M CB SPECFC CONDCTNCE GRAPH			EVAN	56006	T 3	
COMPOUND NO =	86	MOL WGT -	344.4 40.0 4.25	SODIUM HEXADECYL 8 SULFATE X10-3 M CB SPECFC CONDCTNCE GRAPH			EVAN	56006	T 3	
COMPOUND NO =	95	MOL WGT -	280.3 25 6.46	DECYL TRIMETHYL AMMONIUM BROMIDE X10-2 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 2	
5.00 E 2 Y	PRESSURE		25 6.70	X10-2 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1.000E 3 Y	PRESSURE		25 6.70	X10-2 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1.500E 3 Y	PRESSURE		25 6.50	X10-2 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
3.000E 3 Y	PRESSURE		25 5.56	X10-2 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
COMPOUND NO =	97	MOL WGT -	308.4 20 1.59	DODECYL TRIMETHYL AMMONIUM BROMIDE X10-2 M BB INTERFACIAL TENSION LOGM			HAYD PHIL	58012	L D	
			25 4.48	X10-1 D BB TURBIDITY PLT LITE SCATR			ANAC JOHN	64017	K D	
			25 1.452X10-2 M						M	
			25 1.44	X10-2 N BA EQUIV CONDCTNCE GRAPH			VOEK TART	55006	T D	
			25 1.564X10-2 W BA	SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T D	
			25 1.42	X10-2 M BA EQUIV CONDCTNCE GRAPH			BRUN HOLT	61016	T D	
1. E-1 W	PHENOL		25 4.62	X10-3 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1.00 E-1 M	NA BR		25 1.38	X10-1 D BB TURBIDITY PLT LITE SCATR			ANAC JOHN	64017	K 3	
5.02 E-1 M	NA BR		25 4.474X10-3 M					M		
			25 6.2	X10-2 D BB TURBIDITY PLT LITE SCATR			ANAC JOHN	64017	K 3	
			25 2.01	X10-3 M				M		
5.00 E 2 Y	PRESSURE		25 1.61	X10-2 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1.000E 3 Y	PRESSURE		25 1.616X10-2 W BA	SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1.500E 3 Y	PRESSURE		25 1.56	X10-2 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
3.000E 3 Y	PRESSURE		25 1.272X10-2 W BA	SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
5. E-1 M	UREA		25 1.56	X10-2 M BA EQUIV CONDCTNCE GRAPH			BRUN HOLT	61016	T 3	
2.0 E 0 M	UREA		25 2.04	X10-2 M BA EQUIV CONDCTNCE GRAPH			BRUN HOLT	61016	T 3	
6.0 E 0 M	UREA		25 4.54	X10-2 M BA EQUIV CONDCTNCE GRAPH			BRUN HOLT	61016	T 3	
5.00 E 2 Y	PRESSURE		25 5.00	X10-3 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1. E-1 W	PHENOL		25 5.03	X10-3 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1.000E 3 Y	PRESSURE		25 4.88	X10-3 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
1.500E 3 Y	PRESSURE		25 3.88	X10-3 W BA SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T 3	
3.000E 3 Y	PRESSURE									
1. E-1 W	PHENOL									
COMPOUND NO =	99	MOL WGT -	364.5 25 9.20	HEXADECYL TRIMETHYLAMMONIUM BROMIDE X10-4 W BA SPECFC CONDCTNCE GRAPH			CZER	66030	T 2	
			35 9.5	X10-4 M BB EQUIV CONDCTNCE GRAPH			HART COLL	36001	P D	
			35 9.8	X10-4 M BB SPECFC CONDCTNCE GRAPH			HART COLL	36001	P D	
			35 1.020X10-3 W BA	SPECFC CONDCTNCE GRAPH			CZER	66030	T D	
			45 1.155X10-3 W BA	SPECFC CONDCTNCE GRAPH			CZER	66030	T 3	
			55 1.320X10-3 W BA	SPECFC CONDCTNCE GRAPH			CZER	66030	T 3	
COMPOUND NO =	103	MOL WGT -	234.3 15 1.07	HEXYL /OXYETHYLENE/ 3 ALCOHOL HOMOGENEOUS HEAD GROUP X10-3 M BB SURFACE TENSION LOG PLOT			CORK GOOD	64003	T 3	
			25 1.00	X10-1 M BB SURFACE TENSION LOG PLOT			CORK GOOD	64003	T 3	
			35 7.8	X10-2 M BB SURFACE TENSION LOG PLOT			CORK GOOD	64003	T 3	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molar; Y – atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

	Additives	Temp. °C.	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO =	104 MOL WGT — HOMOGENEOUS HEAD GROUP	262.4	OCTYL /OXYETHYLENE/ 3 ALCOHOL					
		15	9.3 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		25	7.5 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
COMPOUND NO =	105 MOL WGT — HOMOGENEOUS HEAD GROUP	394.6	OCTYL /OXYETHYLENE/ 6 ALCOHOL					
		15	1.19 X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		25	9.9 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		35	7.7 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
COMPOUND NO =	106 MOL WGT — HOMOGENEOUS HEAD GROUP	526.7	OCTYL /OXYETHYLENE/ 9 ALCOHOL					
		15	1.6 X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		25	1.3 X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		35	1.1 X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
COMPOUND NO =	107 MOL WGT — HOMOGENEOUS HEAD GROUP	290.4	DECYL /OXYETHYLENE/ 3 ALCOHOL					
		15	7.3 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		25	6.0 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		35	5.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
COMPOUND NO =	108 MOL WGT — HOMOGENEOUS HEAD GROUP	422.6	DECYL /OXYETHYLENE/ 6 ALCOHOL					
		15	1.14 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		25	9.0 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		35	6.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
COMPOUND NO =	109 MOL WGT — HOMOGENEOUS HEAD GROUP	554.8	DECYL /OXYETHYLENE/ 9 ALCOHOL					
		15	1.4 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		25	1.3 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		35	1.1 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
COMPOUND NO =	110 MOL WGT — HOMOGENEOUS HEAD GROUP	450.7	DODECYL /OXYETHYLENE/ 6 ALCOHOL					
		15	1.08 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		25	8.7 X10-5 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
		35	7.2 X10-5 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3
COMPOUND NO =	111 MOL WGT —	272.3	LITHIUM DODECYL 1 SULFATE					
		25	8.77 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2
		25	8.93 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2
COMPOUND NO =	112 MOL WGT —	339.5	TETRAMETHYL AMMONIUM DODECYL 1 SULFATE					
		25	5.41 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2
		25	5.52 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2
COMPOUND NO =	179 MOL WGT —	272.3	SODIUM DODECANE 1-SULFONATE					
		25	9.8 X10-3 M	AB	SURFACE TENSION LOG PLOT	BUJA GOOD	65007	L 3
		40	9.7 X10-3 M	AB	SURFACE TENSION LOG PLOT	BUJA GOOD	65007	L 3
COMPOUND NO =	181 MOL WGT —	216.2	SODIUM OCTYL 1-SULFONATE					
		23	1.55 X10-1 N	CB	UNSPECIFIED CONDUCTANCE	TART LELO	55021	T 3
		25	1.55 X10-1 M	CC	REFRACTIVE INDEX	KLEV	48005	T 3
		40	1.62 X10-1 M	CC	REFRACTIVE INDEX	KLEV	48005	T 3
COMPOUND NO =	182 MOL WGT —	244.3	SODIUM DECYL 1-SULFONATE					
		30	1.066X10 0 D	BA	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T 2
			4.363X10-2 M					M
1.0 E-1 M	NA CL	30	5.364X10-1 D	BA	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T 2
			2.195X10-2 M					M

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l (or kg); W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO =	189 MOL WGT -	308.4 25 5.2 X10-3 M	ALPHA SULFOMYRISTIC ACID CC EQUIV CONDCTNCE GRAPH			WEIL STIR	56008	K 3	
COMPOUND NO =	203 MOL WGT - 3.33 E 1 C 0041	235.8 25 6.11 X10-2 M 25 1.96 X10-2 M	DECYL TRIMETHYL AMMONIUM CHLORIDE BB UNSPECIFIED CONDUCTANCE UNSPEC SOLUBLZTN SDN 4			HOYE MARM	61002	T 3	
COMPOUND NO =	208 MOL WGT - HOMOGENEOUS HEAD GROUP	294.4 25 1.32 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	209 MOL WGT - HOMOGENEOUS HEAD GROUP	338.5 25 9.7 X10-5 M	P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	210 MOL WGT - HOMOGENEOUS HEAD GROUP	382.5 25 1.25 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	211 MOL WGT - HOMOGENEOUS HEAD GROUP	426.6 25 1.54 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	212 MOL WGT - HOMOGENEOUS HEAD GROUP	470.7 25 2.05 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	213 MOL WGT - HOMOGENEOUS HEAD GROUP	514.7 25 2.46 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	214 MOL WGT - HOMOGENEOUS HEAD GROUP	558.8 25 2.80 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	215 MOL WGT - HOMOGENEOUS HEAD GROUP	602.8 25 3.35 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	216 MOL WGT - HOMOGENEOUS HEAD GROUP	646.9 25 3.35 X10-4 M	P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL BC SURFACE TENSION LOG PLOT			CROO FORD	63017	GL 3	
COMPOUND NO =	254 MOL WGT -	187.3 25 5.4 X10-2 M	NONYL DIMETHYL AMINE OXIDE BC HEAT OF DILUTION			BENJ	64016	L 3	
COMPOUND NO = 1. E-3 M K OH	255 MOL WGT -	354.6 60 7.5 X10-3 M	POTASSIUM 9,10 DIHYDROXY STEARATE BC EQUIV CONDCTNCE GRAPH			GREG TART	48012	T 3	
COMPOUND NO =	258 MOL WGT -	360.3 25 5.3 X10-2 M	SODIUM DI-N-AMYL SULFOSUCCINATE BC SURFACE TENSION LOG PLOT			WILL DIXO	57009	AL 3	
COMPOUND NO =	259 MOL WGT -	388.4 25 1.24 X10-2 M	SODIUM DI-N-HEXYL SULFOSUCCINATE BC SURFACE TENSION LOG PLOT			WILL DIXO	57009	TL 3	
COMPOUND NO =	260 MOL WGT -	444.5 25 6.8 X10-4 M	SODIUM DI-N-OCTYL SULFOSUCCINATE BC SURFACE TENSION LOG PLOT			WILL DIXO	57009	TL 3	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

	Additives		Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 262	MOL WGT -	444.5	SODIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE	25 2.5 X10-3 M	BC SURFACE TENSION LOG PLOT	WILL DIXO	57009	TL 3		
COMPOUND NO = 270 4.74 E 0 H	MOL WGT - METHANOL	348.1	OCTADECYL TRIMETHYLAMMONIUM CHLORIDE	25 4.00 X10-4 M	BB EQUIV CONDCTNCE GRAPH	GRIE KRAU	48010	P 3		
COMPOUND NO = 272	MOL WGT -	284.3	TRI-ISOPROPYL BENZENE SULFONIC ACID	50 5.5 X10-2 M	FB EQUIV CONDCTNCE GRAPH	SHUC LING	49004	T 3		
COMPOUND NO = 273 1.1 E 1	MOL WGT - PH OF SOLUTION	222.3	SODIUM DODECANOATE	25 2.44 X10-2 M	DB SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T 3		
1.1 E 1	PH OF SOLUTION	24	2.3 X10-2 M	BC SURFACE TENSION LOG PLOT	HARV	56018	T 3			
30	2.25 X10-2 M	BC SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T 3					
COMPOUND NO = 274 1.00 E 2 E	MOL WGT - NITROBENZENE	340.0	HEXADECYL PYRIDINIUM CHLORIDE	25 9.0 X10-4 M	BB FOTOMTR SOLUBLZTN AZBZ	HART	38001	P 3		
80	2.36 X10-3 M	BC SPECFC CONDCTNCE GRAPH	HART	36002	P 3					
25	5.8 X10-4 M	BA EQUIV CONDCTNCE GRAPH	GRIE KRAU	49018	T 3					
COMPOUND NO = 278	MOL WGT -	283.9	DODECYL PYRIDINIUM CHLORIDE	25 1.47 X10-2 M	CC TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T 3		
COMPOUND NO = 279 5. E 2 Y	MOL WGT - PRESSURE	340.0	DODECYL BENZYL DIMETHYLAMMONIUM CHLORIDE	25.0 7.8 X10-3 M	CB SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3		
1. E 3 Y	PRESSURE	25.0	8.25 X10-3 M	CB SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3			
2. E 3 Y	PRESSURE	25.0	8.5 X10-3 M	CB SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3			
3. E 3 Y	PRESSURE	25.0	8.2 X10-3 M	CB SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3			
4. E 3 Y	PRESSURE	25.0	7.5 X10-3 M	CB SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3			
5. E 3 Y	PRESSURE	25.0	7.1 X10-3 M	CB SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3			
		25.0	6.8 X10-3 M	CB SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3			
COMPOUND NO = 287	MOL WGT -	365.6	OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE	25 2.016X10-2 M	CA SPECFC CONDCTNCE GRAPH	TART LING	43004	T 3		
COMPOUND NO = 288	MOL WGT -	421.7	DECYL TRIMETHYLAMMONIUM DECANESULFONATE	25 1.36 X10-3 M	BC METHOD NOT CITED	CORK GOOD	66014	T 3		
		40	1.40 X10-3 M	CA SPECFC CONDCTNCE GRAPH	TART LING	43004	P 3			
COMPOUND NO = 290 2. E-2 M	MOL WGT - K BR	328.3	DODECYL PYRIDINIUM BROMIDE	5 1.15 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3		
4. E-2 M	K BR	10	1.12 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 2			
6. E-2 M	K BR	15	1.10 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 2			
8. E-2 M	K BR	20	1.12 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 2			
6. E-2 M	LI BR	25	1.14 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 1			
6. E-2 M	RB BR	25	1.13 X10-2 M	CA SPECFC CONDCTNCE GRAPH	BENT SPAR	66038	T 1			
		30	1.18 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 2			
		35	1.22 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 2			
		40	1.28 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3			
		45	1.35 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3			
		50	1.40 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3			
		55	1.48 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3			
		60	1.54 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3			
		65	1.63 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3			
		70	1.72 X10-2 W	BA SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T 3			
		25	7.32 X10-3 M	CC TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T 3			
		25	4.88 X10-3 M	CC TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T 3			
		25	3.96 X10-3 M	CC TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T 3			
		25	3.36 X10-3 M	CC TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T 3			
		25	3.96 X10-3 M	CC TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T 3			
		25	3.35 X10-3 M	CC TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T 3			
COMPOUND NO = 295	MOL WGT -	246.2	SODIUM NONYL 1-SULFATE	21 1.59 X10 0 D	BB TURBIDITY PLT LITE SCATR	HUIS	64047	T 3		
				6.458X10-2 M				M		
3. E-2 M	NA CL	21	1.30 X10 0 D	BB TURBIDITY PLT LITE SCATR	HUIS	64047	T 3		M	
1. E-1 M	NA CL	21	5.280X10-2 M	BB TURBIDITY PLT LITE SCATR	HUIS	64047	T 3		M	
				4.001X10-2 M				T		

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l (or kg); W – molal; Y – atm. Details on page 222

Table of Recommended and Selected Critical Micelle Concentrations – Continued

		Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
3.	E-1 M	NA CL		21 6.3 X10-1 D 2.55 X10-2 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	3
COMPOUND NO =	298	MOL WGT -	250.3	SODIUM TETRADECANOATE						
			25 6.9 X10-3 M	DB	SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T	3	
			35 6.95 X10-3 M	DB	SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T	3	
COMPOUND NO =	299	MOL WGT -	194.2	SODIUM DECANOATE						
			25 9.40 X10-2 M	DB	SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T	3	
			35 9.80 X10-2 M	DB	SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T	3	
COMPOUND NO =	301	MOL WGT -	348.4	SODIUM 3-N-DODECYL BENZENE SULFONATE						
			25 1.46 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T	3	
COMPOUND NO =	302	MOL WGT -	348.4	SODIUM 4-N-DODECYL BENZENE SULFONATE						
			25 1.59 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T	3	
COMPOUND NO =	311	MOL WGT -	274.3	SODIUM UNDECYL 1-SULFATE						
			21 4.4 X10-1 D 1.60 X10-2 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	3	
3.	E-2 M	NA CL		21 2.65 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	3
1.	E-1 M	NA CL		21 9.660X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	3
3.	E-1 M	NA CL		21 1.49 X10-1 D 5.432X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	3
			21 8.5 X10-2 D	BC	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	3	
			3.09 X10-3 M							
COMPOUND NO =	330	MOL WGT -	550.9	HEXADECYL/OXYETHYLENE/7 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			25 1.74 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T	3	
COMPOUND NO =	331	MOL WGT -	639.0	HEXADECYL/OXYETHYLENE/9 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			25 2.09 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T	3	
COMPOUND NO =	332	MOL WGT -	771.2	HEXADECYL/OXYETHYLENE/12 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			25 2.34 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T	3	
COMPOUND NO =	333	MOL WGT -	903.4	HEXADECYL/OXYETHYLENE/15 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			25 3.09 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T	3	
COMPOUND NO =	334	MOL WGT -	1,167.8	HEXADECYL/OXYETHYLENE/21 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			25 3.89 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T	3	
COMPOUND NO =	340	MOL WGT -	410.8	MAGNESIUM OCTANE SULFONATE						
			23 1.10 X10-1 N	CB	UNSPECIFIED CONDUCTANCE	TART LELO	55021	T	3	
COMPOUND NO =	341	MOL WGT -	466.9	MAGNESIUM DECANE SULFONATE						
			60 2.0 X10-2 N	CC	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L	3	
COMPOUND NO =	342	MOL WGT -	523.1	MAGNESIUM DODECANE SULFONATE						
			60 3.6 X10-3 N	CB	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L	3	
COMPOUND NO =	346	MOL WGT -	437.8	DECYL TRIMETHYLAMMONIUM DECYL SULFATE						
1.00 E 2 I	NA BR		25 4.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	63014	L	3	
			25 4.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	63014	L	3	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 347 MOL WGT — 1.00 E 2 I NA BR		381.6 25	OCTYL TRIMETHYLMAMMONIUM OCTYL SULFATE 7.5 X10-3 M BB SURFACE TENSION LOG PLOT			CORK GOOD	65005	T 3
COMPOUND NO = 348 MOL WGT — N-OXIDE /C7F15CH2N/CH3/20/		443.2 25	NN-DIMETHYL 1-1-DIHYDROPENTADECAFLUORO OCTYL AMINE 4.7 X10-4 M BB SURFACE TENSION LOG PLOT			CORK GOOD	65005	T 3
COMPOUND NO = 353 MOL WGT — 1.78 E-1 M K CL 2.65 E-1 M K CL		393.7 RM RM RM	OCTYL TRIMETHYLMAMMONIUM DECAN SULFONATE 5.75 X10-3 M BB DEBYE PLT LIGHT SCATTER 5.67 X10-3 M BB DEBYE PLT LIGHT SCATTER 5.67 X10-3 M BB DEBYE PLT LIGHT SCATTER			ANAC	53002	T 3
COMPOUND NO = 376 MOL WGT —		375.4 24.9 25 25	DODECYL PYRIDINIUM IODIDE 5.26 X10-3 M CB MICELLAR SPECTRAL CHANGE 5.60 X10-3 M DC TURBIDITY PLT LITE SCATR 5.70 X10-3 M CB SURFACE TENSION UNSPEC			MUKE RAY PARR FORD OTTE	66006 60015 66028	T 3 T 3 T 3
COMPOUND NO = 380 MOL WGT — HOMOGENEOUS HEAD GROUP		278.4 20	HEXYL/OXYETHYLENE/4 ALCOHOL 9.0 X10-2 M BB REFRACTIVE INDEX			DONB JAN	63021	T 3
COMPOUND NO = 381 MOL WGT — HOMOGENEOUS HEAD GROUP		322.5 20	HEXYL/OXYETHYLENE/5 ALCOHOL 9.25 X10-2 M BB REFRACTIVE INDEX			DONB JAN	63021	T 3
COMPOUND NO = 385 MOL WGT —		437.8 25	OCTYL TRIMETHYLMAMMONIUM DODECYL SULFATE 4.3 X10-4 M BC METHOD NOT CITED			CORK GOOD	66014	T 3
COMPOUND NO = 393 MOL WGT — HOMOGENEOUS HEAD GROUP		338.5 20 30	BUTYL/OXYETHYLENE/6 ALCOHOL 7.96 X10-1 M BC SURFACE TENSION LOG PLOT 7.60 X10-1 M BC SURFACE TENSION LOG PLOT			ELWO FLOR ELWO FLOR	64049 64049	T 3 T 3
COMPOUND NO = 394 MOL WGT — HOMOGENEOUS HEAD GROUP		338.5 20 30	I-METHYL PROPYL/OXYETHYLENE/6 ALCOHOL 9.1 X10-1 M BC SURFACE TENSION LOG PLOT 8.8 X10-1 M BC SURFACE TENSION LOG PLOT			ELWO FLOR ELWO FLOR	64049 64049	T 3 T 3
COMPOUND NO = 395 MOL WGT — HOMOGENEOUS HEAD GROUP		366.6 20 30	2-ETHYL BUTYL/OXYETHYLENE/6 ALCOHOL 1.00 X10-1 M BC SURFACE TENSION LOG PLOT 9.3 X10-2 M BC SURFACE TENSION LOG PLOT			ELWO FLOR ELWO FLOR	64049 64049	T 3 T 3
COMPOUND NO = 396 MOL WGT — HOMOGENEOUS HEAD GROUP		394.6 20 30	2-PROPYL PENTYL/OXYETHYLENE/6 ALCOHOL 2.30 X10-2 M BC SURFACE TENSION LOG PLOT 2.0 X10-2 M BC SURFACE TENSION LOG PLOT			ELWO FLOR ELWO FLOR	64049 64049	T 3 T 3
COMPOUND NO = 397 MOL WGT — HOMOGENEOUS HEAD GROUP		422.7 20 25	2-BUTYL HEXYL/OXYETHYLENE/6 ALCOHOL 3.10 X10-3 M BC SURFACE TENSION LOG PLOT 2.84 X10-3 M BC SURFACE TENSION LOG PLOT			ELWO FLOR ELWO FLOR	64049 64049	T 3 T 3
COMPOUND NO = 398 MOL WGT — HOMOGENEOUS HEAD GROUP		554.9 20 30	2-BUTYL HEXYL/OXYETHYLENE/9 ALCOHOL 3.20 X10-3 M BC SURFACE TENSION LOG PLOT 2.79 X10-3 M BC SURFACE TENSION LOG PLOT			ELWO FLOR ELWO FLOR	64049 64049	T 3 T 3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
 D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality
 counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg;
 T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

	Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO =	416 MOL WGT -	314.1	PERFLUORO HEXANOIC ACID						
		0	1.09 X10-1 M	BC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3	
		18	1.06 X10-1 M	BC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3	
COMPOUND NO =	417 MOL WGT -	414.1	PERFLUORO OCTANOIC ACID						
		18	9.8 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3	
		35	9.3 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3	
		45	1.02 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3	
COMPOUND NO =	423 MOL WGT - HOMOGENEOUS HEAD GROUP	174.3	OCTYL/OXYETHYLENE/1 ALCOHOL--OCTYL GLYCOL ETHER						
		25	4.9 X10-3 M	BC	SURFACE TENSION LOG PLOT	SHIN YAMA	59013	T 3	
COMPOUND NO =	424 MOL WGT -	204.3	OCTYL ALPHA-GLYCERYL ETHER						
		25	5.8 X10-3 M	BC	SURFACE TENSION LOG PLOT	SHIN YAMA	59013	T 3	
COMPOUND NO =	427 MOL WGT -	384.5	HEXADECYL PYRIDINIUM BROMIDE						
		25	5.81 X10-4 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 2	
		35	7.5 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	HART COLL	36001	T 3	
		35	7.7 X10-4 M	BB	SPECFC CONDCTNCE GRAPH	HART COLL	36001	P 3	
6.4 E 0 H	METHANOL	25	7.51 X10-4 M	BC	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 3	
1.470E 1 H	METHANOL	25	1.18 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 2	
1.991E 1 H	METHANOL	25	1.69 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 2	
2.602E 1 H	METHANOL	25	2.81 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 3	
3.520E 1 H	METHANOL	25	6.01 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 2	
COMPOUND NO =	449 MOL WGT -	235.9	DODECYLMETHYL AMMONIUM CHLORIDE						
		30	1.46 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS BROO	49013	T 3	
COMPOUND NO =	450 MOL WGT -	249.9	DODECYLDIMETHYL AMMONIUM CHLORIDE						
		30	1.61 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS BROO	49013	T 3	
COMPOUND NO =	456 MOL WGT -	452.2	POTASSIUM PERFLUORO OCTANOATE						
		25	2.88 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T 3	
		30	2.74 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T 3	
		40	2.65 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T 3	
		55	2.76 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T 3	
		70	3.07 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T 3	
		85	3.54 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T 3	
9.4 E-3 W	K N03	30	2.43 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
1.82 E-2 W	K N03	30	2.22 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
2.96 E-2 W	K N03	30	2.01 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
8.3 E-3 W	K N03	40	2.40 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
2.39 E-2 W	K N03	40	2.07 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
4.08 E-2 W	K N03	40	1.79 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
7.31 E-2 W	K N03	40	1.46 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
1.13 E-2 W	K N03	55	2.42 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
2.30 E-2 W	K N03	55	2.17 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
3.54 E-2 W	K N03	55	1.93 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
4.65 E-2 W	K N03	55	1.82 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
6.01 E-2 W	K N03	55	1.65 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
7.77 E-2 W	K N03	55	1.49 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
1.28 E-2 W	K N03	70	2.75 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
3.77 E-2 W	K N03	70	2.22 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
8.22 E-2 W	K N03	70	1.82 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
4.65 E-2 W	K N03	85	2.59 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
7.71 E-2 W	K N03	85	2.27 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L 3	
COMPOUND NO =	458 MOL WGT -	347.3	DECYL PYRIDINIUM IODIDE						
		RM	2.25 X10-2 M	CB	MICELLAR SPECTRAL CHANGE	HARK KRIZ	51010	T 3	
COMPOUND NO =	482 MOL WGT -	248.4	DODECYL AMMONIUM NITRATE						
		30	9.9 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K 3	
COMPOUND NO =	483 MOL WGT -	306.0	DIOCTYL DIMETHYL AMMONIUM CHLORIDE						
		30	2.83 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48014	K 3	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations—Continued

Additives		Temp. °C	CMC		Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 491 MOL WGT —		356.5	DODECYL	TROPYLIUM BISULFATE						
7.8 E 1 H H ₂ SO ₄		25	2.2 X10-3 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3		
9.6 E 1 H H ₂ SO ₄		25	1.25 X10-2 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3		
9.6 E 1 H H ₂ SO ₄		25	5.4 X10-3 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3		
9. E-1 M NA ₂ SO ₄										
COMPOUND NO = 493 MOL WGT —		306.4	SODIUM	P-NONYL BENZENE SULFONATE						
		20	4.6 X10-3 M	BB	SURFACE TENSION LOG PLOT	HARR	59001	L 3		
COMPOUND NO = 500 MOL WGT —		354.0	DODECYL	TRI/2-HYDROXYETHYL/AMMONIUM CHLORIDE						
		30	1.38 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49009	K 3		
COMPOUND NO = 525 MOL WGT —		316.4	SODIUM	TETRADECYL 6-SULFATE						
		60	9.80 X10-3 M		UNSPECIFIED CONDUCTANCE	WINS	48008	L 3		
COMPOUND NO = 528 MOL WGT —		358.9	DODECYL	TROPYLIUM PERCHLORATE						
7.0 E 1 H H ClO ₄		25	8.6 X10-4 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3		
		30	1.25 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	SATA IWAM	63034	T 3		
COMPOUND NO = 568 MOL WGT —		555.1	MAGNESIUM	DODECYL SULFATE						
		30	1.25 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	SATA IWAM	63034	T 3		
COMPOUND NO = 572 MOL WGT —		589.7	COBALTOUS	DODECYL SULFATE						
		30	1.23 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	SATA IWAM	63034	T 3		
COMPOUND NO = 573 MOL WGT —		594.3	CUPRIC	DODECYL SULFATE						
		30	1.20 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	SATA IWAM	63034	T 3		
COMPOUND NO = 575 MOL WGT —		589.5	NICKEL	DODECYL SULFATE						
		30	1.24 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	SATA IWAM	63034	T 3		
COMPOUND NO = 587 MOL WGT —		307.5	DECYL	DIMETHYLAMMONIOPROPANE SULFONATE						
		30	1.20 X10-0 D	BB	TURBIDITY PLT LITE SCATR	HERR	66013	T M		
			3.902X10-2 M							
COMPOUND NO = 588 MOL WGT —		335.6	DODECYL	DIMETHYLAMMONIOPROPANE SULFONATE						
2. E-1 M NA CL		30	1.2 X10-1 D	BC	TURBIDITY PLT LITE SCATR	HERR	66013	T 3		
1. E O M NA CL		30	1.0 X10-1 D	BC	METHOD NOT CITED	HERR	66013	G 3		
			2.97 X10-3 M							
		30	5.8 X10-2 D	BC	METHOD NOT CITED	HERR	66013	G 3		
			1.72 X10-3 M							
COMPOUND NO = 590 MOL WGT —		299.6	DODECYL	DIMETHYLAMMONIOPROPANE CARBOXYLATE						
		30	1.6 X10-1 D	BC	TURBIDITY PLT LITE SCATR	HERR	66013	T M		
			5.34 X10-3 M							
COMPOUND NO = 591 MOL WGT —		352.5	DODECYL	DIMETHYLPHOSPHONIOPROPANE SULFONATE						
		30	8.5 X10-2 D	BC	TURBIDITY PLT LITE SCATR	HERR	66013	T 3		
			2.41 X10-3 M							
COMPOUND NO = 593 MOL WGT —		351.6	DIMETHYL	DODECYLAMMONIOPROPANE HYDROXY SULFONATE						
		30	7.0 X10-2 D	BC	METHOD NOT CITED	HERR	66013	T 3		
			1.99 X10-3 M							
COMPOUND NO = 594 MOL WGT —		391.7	DODECYL	DIPROPYL AMMONIOPROPANE SULFONATE						
		30	7.0 X10-2 D	BC	METHOD NOT CITED	HERR	66013	T 3		
			1.78 X10-3 M							
COMPOUND NO = 634 MOL WGT —		304.5	POTASSIUM	DODECYL SULFATE						
		40	7.8 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	MEGU KOND	56020	T 3		
COMPOUND NO = 640 MOL WGT —		325.5	HEXYL	TRIMETHYLAMMONIUM HEXANE SULFATE						
		25	1.1 X10-1 M	BC	METHOD NOT CITED	CORK GOOD	66014	T 3		
COMPOUND NO = 641 MOL WGT —	1.00 E 2 I NA BR	353.6	HEXYL	TRIMETHYLAMMONIUM OCTANE SULFATE						
		25	2.7 X10-2 M	BC	METHOD NOT CITED	CORK GOOD	66014	T 3		

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/d or kg; W—molal; Y—atm. Details on page 222.

Table of Recommended and Selected Critical Micelle Concentrations – Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO =	642	MOL WGT -	409.7 25	OCTYL TRIMETHYLLAMMONIUM DECANE SULFATE X10-3 M BC METHOD NOT CITED			CORK GOOD	66014	T 3
COMPOUND NO = 1.00 E 2 I	643	MOL WGT - NA BR	437.8 25	DODECYL TRIMETHYLLAMMONIUM OCTANE SULFATE X10-4 M BC METHOD NOT CITED			CORK GOOD	66014	T 3
COMPOUND NO = 5.01 E 0 H	654	MOL WGT - METHANOL	374.7 25	OCTADECYL TRIMETHYLLAMMONIUM NITRATE X10-4 M BA EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 2
1.227E 1 H		METHANOL	25	3.28 X10-4 M BB EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 3
1.508E 1 H		METHANOL	25	4.37 X10-4 M BB EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 3
1.985E 1 H		METHANOL	25	5.93 X10-4 M BA EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 2
2.589E 1 H		METHANOL	25	1.10 X10-3 M BA EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 2
3.463E 1 H		METHANOL	25	3.03 X10-3 M BB EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 3
COMPOUND NO =	655	MOL WGT -	368.1 25	OCTADECYL PYRIDINIUM CHLORIDE X10-4 M BB EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 3
COMPOUND NO = 2.0 E 1 H	656	MOL WGT - METHANOL	394.7 25	OCTADECYL PYRIDINIUM NITRATE X10-4 M BB EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	T 3
EVER KRAU			25	5.76 X10-4 M BB EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 3
COMPOUND NO = 2.0 E 1 H	657	MOL WGT - METHANOL	412.6 25	OCTADECYL PYRIDINIUM BROMIDE X10-4 M BB EQUIV CONDCTNCE GRAPH			EVER KRAU	48028	P 3
COMPOUND NO =	658	MOL WGT -	440.6 25	OCTADECYL TRIMETHYLLAMMONIUM BROMATE X10-4 M BB EQUIV CONDCTNCE GRAPH			GRIE KRAU	48010	P 3
COMPOUND NO =	659	MOL WGT -	357.7 25	OCTADECYL TRIMETHYLLAMMONIUM FORMATE X10-4 M BC EQUIV CONDCTNCE GRAPH			GRIE KRAU	48010	T 3
COMPOUND NO =	662	MOL WGT -	482.7 25	OCTADECYL TRIETHYLLAMMONIUM BROMATE X10-4 M BC EQUIV CONDCTNCE GRAPH			MCDO KRAU	51009	T 3
COMPOUND NO = 9.91 E 0 H	667	MOL WGT - ACETONE	713.4 25	OCTADECYL TRIMETHYLLAMMONIUM OXALATE X10-4 M BB EQUIV CONDCTNCE GRAPH			YOUN GRIE	49017	P 3
1.39 E 1 H		ACETONE	25	2.56 X10-4 M BB EQUIV CONDCTNCE GRAPH			YOUN GRIE	49017	P 3
2.07 E 1 H		ACETONE	25	6.50 X10-4 M BB EQUIV CONDCTNCE GRAPH			YOUN GRIE	49017	P 3
2.88 E 1 H		ACETONE	25	1.94 X10-3 M BC EQUIV CONDCTNCE GRAPH			YOUN GRIE	49017	P 3
2.07 E 1 H		METHANOL	25	2.50 X10-4 M BB EQUIV CONDCTNCE GRAPH			YOUN GRIE	49017	P 3
3.02 E 1 H		METHANOL	25	6.25 X10-4 M BB EQUIV CONDCTNCE GRAPH			YOUN GRIE	49017	P 3
4.01 E 1 H		METHANOL	25	2.20 X10-3 M BC EQUIV CONDCTNCE GRAPH			YOUN GRIE	49017	P 3
COMPOUND NO =	668	MOL WGT -	292.5 20	DIPOTASSIUM OCTYL MALONATE X10-1 M BC SURFACE TENSION UNSPEC			SHIN	55007	T 3
COMPOUND NO =	670	MOL WGT -	348.6 25	DIPOTASSIUM DODECYL MALONATE X10-2 M BB FOTOMTR SOLUBLZTN 2NPA			SHIN	55007	T 3
COMPOUND NO =	671	MOL WGT -	376.7 25	DIPOTASSIUM TETRADECYL MALONATE X10-2 M BC FOTOMTR SOLUBLZTN 2NPA			SHIN	55007	T 3
COMPOUND NO =	710	MOL WGT -	190.3 30	OCTYL DIMETHYL PHOSPHINE OXIDE X10-1 D BB TURBIDITY PLT LITE SCATR			HERR BRUS	66039	T M 3
			4.04 X10-2 M						
COMPOUND NO =	712	MOL WGT -	246.4 1	DODECYL DIMETHYL PHOSPHINE OXIDE X10-2 D BC TURBIDITY PLT LITE SCATR			HERR BRUS	66039	T M 3
			8.11 X10-4 M						
			30	1.4 X10-2 D BD TURBIDITY PLT LITE SCATR			HERR BRUS	66039	T M 3
			5.68 X10-4 M						
COMPOUND NO =	717	MOL WGT -	378.4 25	DODECYLQUINOLINIUM BROMIDE X10-3 M BB SURFACE TENSION LOG PLOT			FEW GILB	58031	T 3
COMPOUND NO =	720	MOL WGT -	733.2 30	1-6-DITRIMETHYLLAMMONIUM-HEXANE/DODECYL SULFATE/2 X10-4 M CB EQUIV CONDCTNCE GRAPH			MEGU KOND	59026	T 3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 1 MOL WGT -	288.3	SODIUM DODECYL 1 SULFATE						
	53005	VALUES FRM REF IN CMC			KLIN LANG	57022	R	
	59016	VALUES FRM REF IN CMC			BOTR CRES	60024	R	
	47006	VALUES FRM REF IN CMC			CORR HARK	46005	R	
		GRAPH DATA NOT RETRIEVED			KLEV	46007	R	
01.0	7.6 X10-3 M	BB	SURFACE TENSION LOG PLOT		SCHI	63026	L L	
05.0	7.5 X10-3 M	BB	SURFACE TENSION LOG PLOT		SCHI MANN	66001	L L	
08.0	8.6 X10-3 M	BB	AVER SP EQUIV COND		FLOC UBBE	53008	G L	
UNK	1.83 X10-1 D	CC	TURBIDITY PLT LITE SCATR		KUSH HUBB	55003	K L	
	6.347X10-3 M						M	
10	8.67 X10-3 M	BA	AVER SP EQUIV COND		FLOC	61007	L 1	
10	8.81 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS		GODD BENS	57011	L 1	
10.5	8.5 X10-3 M	BB	AVER SP EQUIV COND		FLOC UBBE	53008	G L	
10.0	8.55 X10-3 M	BA	EQUIV CONDCTNCE GRAPH		FLOC	57013	L 1	
10	7.4 X10-3 M	BB	SURFACE TENSION LOG PLOT		SCHI	64020	T L	
15	8.51 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS		GODD BENS	57011	L 1	
15	9.65 X10-3 M	BB	EQUIV CONDCTNCE GRAPH		SHIR MATU	65020	L L	
15	8.43 X10-3 M	BA	AVER SP EQUIV COND		FLOC	61007	L 1	
20	8.47 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS		GODD BENS	57011	L 1	
20	9. X10-3 M	DE	VISCOSITY		HESS PHIL	39009	T L	
20	8.0 X10-3 M	BB	INTERFACIAL TENSION LOGM		V VO	60025	T L	
20	7.22 X10-3 M	CG	EQUIV COND 1ST DEVIATION		WARD	40004	T L	
20	8.5 X10-3 M	CB	EQUIV CONDCTNCE GRAPH		WARD	40004	P L	
20	7.7 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCR		GODD HARV	53012	T L	
20	6.6 X10-3 M	BB	SURFACE TENSION LOG PLOT		HARR	60004	L L	
20	7.1 X10-3 M	BB	SURFACE TENSION LOG PLOT		HARR	60004	G L	
20	8.25 X10-3 M	BA	AVER SP EQUIV COND		FLOC	61007	L 1	
20	7.95 X10-3 M	BB	INTERFACIAL TENSION LOGM		HAYD PHIL	58012	L L	
20	8.0 X10-3 M	BB	SURFACE TENSION LOG PLOT		V VO	61026	TL L	
21	2.5 X10-1 D	BD	REFRACTIVE INDEX		HUIS	64047	T L	
	8.67 X10-3 M						M	
21	2.30 X10-1 D	BB	TURBIDITY PLT LITE SCATR		HUIS	64047	T L	
	7.977X10-3 M						M	
21	2.15 X10-1 D	BB	SURFACE TENSION LOG PLOT		HUIS	64047	T L	
	7.457X10-3 M						M	
23	8. X10-3 N	CD	UNSPECIFIED CONDUCTANCE		TART LELO	55021	T L	
24.7	1.3 X10-1 P	HC	SPECFC CONDCTNCE GRAPH		ROSS BRAM	57031	T L	
25	1.83 X10-1 D	CG	VISUAL SPCTR CHNGE PNCR		SCHI FOWK	57014	T L	
	6.347X10-3 M						M	
25	2.21 X10-1 D	CB	UNSPECIFIED CONDUCTANCE		SCHI FOWK	57014	T L	
	7.665X10-3 M						M	
25	7.4 X10-3 M	BB	SURFACE TENSION LOG PLOT		MIYA	60029	T L	
25	2.330X10-1 D	BA	EQUIV CONDCTNCE GRAPH		WILL MYSE	55005	T 1	
	8.081X10-3 M						M	
25	2.340X10-1 D	BA	FOTOMTR SOLUBLZTN OROT		WILL MYSE	55005	T 1	
	8.116X10-3 M						M	
25	2.36 X10-1 D	BB	DEBYE PLT LIGHT SCATTER		WILL MYSE	55005	T 1	
	8.185X10-3 M						M	
25	8.39 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS		GODD BENS	57011	L 1	
25.2	8.0 X10-3 M	BA	AVER SP EQUIV COND		FLOC UBBE	53008	T L	
25	8.2 X10-3 M	BA	SPECFC CONDCTNCE GRAPH		CORK GOOD	62006	T L	
25	5.8 X10-3 M	CC	REFRACTIVE INDEX		KLEV	48005	T L	
25	6. X10-3 M	CE	ELECTROMOTIVE FORCE		STAN RADL	60021	T L	
25	8.1 X10-3 N	CC	UNSPECIFIED CONDUCTANCE		HAFF PICC	42003	T L	
25	7.8 X10-3 M	BB	EQUIV COND 1ST DEVIATION		GODD HIGH	55018	T L	
25	8.1 X10-3 M	BA	EQUIV CONDCTNCE GRAPH		GODD HIGH	55018	K 1	
25	8.3 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT		GINN HARR	60010	G L	
25.0	7.4 X10-3 M	BB	SURFACE TENSION LOG PLOT		SCHI	63026	L L	
	2.324X10-1 D	BA	FOTOMTR SOLUBLZTN OROT		SCHO	66036	T 1	
	8.061X10-3 M						M	
25	8.11 X10-3 M	CA	SPECFC CONDCTNCE GRAPH		BENT SPAR	66038	T L	
25	8.3 X10-3 M	BB	EQUIV CONDCTNCE GRAPH		SHIR MATU	65020	L L	
25	8.8 X10-3 M	BC	VISCOSITY		MIUR MATS	57025	T L	
25	8.3 X10-3 M	BB	SURFACE TNSN LINEAR PLOT		MIUR MATS	57025	TA 1	
25	8.2 X10-3 M	BC	EQUIV CONDCTNCE GRAPH		MIUR MATS	57025	T L	
25	8.16 X10-3 M	BA	AVER SP EQUIV COND		FLOC	61007	L 1	
25	8.2 X10-3 M	BC	SPECFC CONDCTNCE GRAPH		MIUR MATS	57025	T L	
25.6	5.4 X10-3 M	CG	FOTOMTR SPCTR CHNGE PNCR		KLEV	47004	T L	
25	8.27 X10-3 W	BA	SPECFC CONDCTNCE GRAPH		HAMA	62036	T 1	
25	7.3 X10-4 M	HG	FOTOMTR SPCTR CHNGE PNCR		GINN HARR	58008	T L	
25	5.63 X10-3 M	HC	SPECFC CONDCTNCE GRAPH		GINN HARR	58008	T L	
25.0	7.2 X10-3 M	BB	SURFACE TENSION LOG PLOT		SCHI MANN	66001	L L	
	1.63 X10-3 M	HG	FOTOMTR SPCTR CHNGE PNCR		GINN HARR	58008	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
	25	4.03 X10-3 M	HC	SPECFC CONDCTNCE GRAPH	GINN HARR	58008	T L	
	25	7.2 X10-2 D	HE	METHOD NOT CITED	GINN HARR	61014	T L	
	25	7.2 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
	25	2.29 X10-1 D	DC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	M
		7.943X10-3 M						
	25.0	8.22 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	MYSE OTTE	61017	TL L	
	25.0	7.83 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	L L	
	25	8.2 X10-3 M	BC	UNSPECIFIED CONDUCTANCE	MIUR ARIS	58023	G L	
	25.0	8.27 X10-3 M	BA	SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T L	
	25	8.0 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	NAKA KAWA	64034	G L	
	25	8.15 X10-3 M	AB	SURFACE TENSION LOG PLOT	ELWO MYSE	66007	T L	
	25	8.2 X10-3 M	AA	SURFACE TNSN LINEAR PLOT	ELWO MYSE	66007	T L	
	25	8.4 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	ELWO MYSE	66007	T L	
	25	6.0 X10-3 M	GC	REFRACTIVE INDEX	LIN	57005	T L	
	26	6.02 X10-3 M	DG	VISUAL SPCTR CHNGE PNPN	CORR HARK	47006	T L	
	26	6.12 X10-3 M	DG	VISUAL SPCTR CHNGE RHD6	CORR HARK	47006	T L	
	26	6.02 X10-3 M	DG	VISUAL SPCTR CHNGE PNPN	CORR HARK	47010	T L	
	29	7.7 X10-3 M	BB	SURFACE TENSION LOG PLOT	BRAD	48022	T L	
	30	1.00 X10-2 M	GB	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
	30.0	8.0 X10-3 M	CB	VELOCITY OF SOUND	SHIG	65022	T L	
	30	8.44 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1	
	30	5. X10-3 M	CE	SURFACE TENSION MINIMUM	KLEV RAIS	54010	T L	
	30	7.0 X10-3 W	CC	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L	
	30	8.23 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 1	
	35	8.57 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1	
	35	8.4 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T 1	
	35	9.18 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L L	
	35	8.39 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 1	
	35.8	5.0 X10-3 M	CG	FOTOMTR SPCTR CHNGE PNPN	KLEV	47004	T L	
	35	7.2 X10-3 M	BC	TURBIDMTR SOLUBLZTN LOH	MIUR ARIS	58023	TA L	
	40	1.2 X10-2 M	XG	VISCOSITY MINIMUM	SATA TYUZ	53006	T L	
	40	8.88 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1	
	40.1	8.1 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G L	
	40	6.1 X10-3 M	BG	VISUAL SPCTR CHNGE PNPN	MIYA	60029	T L	
	40	6.1 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6	MIYA	60029	T L	
	40	7.00 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNPN	GODD HARV	53012	L L	
	40	8.60 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 1	
	40.0	8.65 X10-3 M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T L	
	40	7.2 X10-3 M	XG	VELOCITY OF SOUND	KUPP SURY	65028	T L	
	40	8. X10-3 M	CD	VISCOSITY	NAKA NINO	64025	T L	
	40	8.9 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	MEGU KOND	56020	T L	
	45	9.10 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1	
	45	8.86 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 1	
	45.0	4.6 X10-3 M	CG	FOTOMTR SPCTR CHNGE PNPN	KLEV	47004	T L	
	45.0	7.6 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
	45	7.6 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
	45	7.2 X10-3 M	BC	TURBIDMTR SOLUBLZTN LOH	MIUR ARIS	58023	TA L	
	50	8.1 X10-3 M	CB	EQUIV COND 1ST DEVIATION	LANG	53005	T L	
	50	9.3 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	LANG	53005	K L	
	50	8.1 X10-3 M	BB	INTERFACIAL TENSION LOGM	KLIN LANG	57022	T L	
	50	8.1 X10-3 M	BB	SURFACE TENSION LQG PLOT	KLIN LANG	57022	T L	
	50	9.61 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L D	
	50	8.00 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
	50	7.3 X10-3 W	CC	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L	
	50	8.9 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
	50	9.18 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L D	
	50	5.0 X10-3 M	DG	VISUAL SPCTR CHNGE PNPN	RAIS	52016	T L	
	50	6.8 X10-3 M	BG	VISUAL SPCTR CHNGE PNPN	WEIL STIR	63013	T L	
	50	6.8 X10-3 M	CG	VISUAL SPCTR CHNGE PNPN	WEIL STIR	59004	T L	
	54	9.2 X10-3 M	BB	SURFACE TENSION LOG PLOT	MIYA	60029	T L	
	55	9.95 X10-3 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1	
	55	9.8 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T 1	
	55.0	9.3 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G L	
	55.0	9.49 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 1	
	55	2.26 X10-1 D	CB	UNSPECIFIED CONDUCTANCE	SCHI FOWK	57014	T L	M
		7.839X10-3 M						
	55.0	7.8 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
	55	9.61 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 1	
	55.0	4.5 X10-3 M	CG	FOTOMTR SPCTR CHNGE PNPN	KLEV	47004	T L	
	55	7.3 X10-3 M	BC	TURBIDMTR SOLUBLZTN LOH	MIUR ARIS	58023	TA L	
	60	1.10 X10-2 M	GB	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
	60	1.016X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 2	
	60	2.25 X10-1 D	DC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	M
		7.804X10-3 M						
	65	1.091X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	70	1.4 X10-2 M	XG	VISCOSITY MINIMUM	SATA TYUZ	53006	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation	
	65	1.09 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3	
	70	1.4 X10-2 M	XG	VISCOSITY MINIMUM	SATA TYUZ	53006	T	L	
	70	1.14 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	3	
	70	6.6 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L	
	75	1.13 X10-4 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G	L	
	75	6.6 X10-2 D	HE	METHOD NOT CITED	GINN HARR	61014	T	L	
	90	1.20 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T	L	
UNK	1.4	X10-2 D	HG	SURFACE TENSION UNSPEC	WAN	66018	T	L	
UNK	7.8	X10-3 M	BD	EMF ALONG CONC GRADIENT	BOTR CRES	59015	T	L	
UNK	8.	X10-3 M	BD	ELECTROMOTIVE FORCE	BOTR CRES	59016	T	L	
UNK	5.9	X10-3 M	CG	VISUAL SPCTR CHNGE RHD6	STAN RADL	60021	T	L	
UNK	8.0	X10-3 M	BB	UNSPECIFIED CONDUCTANCE	BOTR CRES	59015	T	L	
UNK	8.	X10-3 M	BC	UNSPECIFIED CONDUCTANCE	BOTR CRES	59016	T	L	
UNK	8.6	X10-3 M	BC	ULTRAfiltration	HUTC	59018	K	L	
UNK	5.1	X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T	L	
RM	7.6	X10-3 M	CG	UNSPECIFIED CONDUCTANCE	PRIN HERM	56002	T	L	
RM	7.15	X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T	L	
UNK	6.2	X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G	L	
RM	7.2	X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MIUR MATS	57025	T	L	
UNK	6.5	X10-3 M	XG	METHOD NOT CITED	KLEV RAIS	54004	T	L	
RM	7.55	X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	L	
UNK	8.0	X10-3 M	BB	ELECTROMOTIVE FORCE	BOTR DE M	64032	T	L	
UNK	1.7	X10-1 D	XG	VISUAL SPCTR CHNGE PNCN	DEMC	61031	T	L	
		5.89 X10-3 M					M		
1. E-2 M	AG NO3	35	5.0 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	3
5. E-4 N	AL CL3	70	5.1 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
7. E-4 N	AL CL3	70	3.8 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
1. E-3 N	AL CL3	70	2.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
2. E-3 N	AL CL3	70	1.6 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
3. E-4 N	BZL* C6H5 /CH3/2 N I	40	4.4 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
5. E-4 N	BZL* C6H5 /CH3/2 N I	40	3.6 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
8. E-4 N	BZL* C6H5 /CH3/2 N I	40	3.2 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1.2 E-3 N	BZL* C6H5 /CH3/2 N I	40	3.0 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1.6 E-3 N	BZL* C6H5 /CH3/2 N I	40	2.8 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1.7 E-3 N	BZL* C6H5 /CH3/2 N I	40	2.6 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
5. E-4 N	(C4H9)(CH3)3 N I	40	5.2 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1. E-3 N	(C4H9)(CH3)3 N I	40	4.3 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1.4 E-3 N	(C4H9)(CH3)3 N I	40	4.0 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1.9 E-3 N	(C4H9)(CH3)3 N I	40	3.5 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
3.0 E-3 N	(C4H9)(CH3)3 N I	40	3.0 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1. E-4 N	CA CL2	70	6.0 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
3. E-4 N	CA CL2	70	5.1 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
5. E-4 N	CA CL2	70	4.3 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
1. E-3 N	CA CL2	70	3.9 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
2. E-3 N	CA CL2	70	3.0 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
5. E-3 N	CA CL2	70	2.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
1.2 E-2 N	CA CL2	70	1.8 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
2.5 E-2 N	CA CL2	70	1.4 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
5. E-2 N	CA CL2	70	1.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
1.0 E-1 N	CA CL2	70	1.0 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
3.0 E-1 N	CA CL2	70	7.7 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
9.45 E-4 N	CS CL	40	6.90 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
2.18 E-3 N	CS CL	40	6.15 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
5.15 E-3 N	CS CL	40	5.20 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
8.64 E-3 N	CS CL	40	4.50 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1.18 E-2 N	CS CL	40	3.75 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
1.39 E-2 N	CS CL	40	3.60 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
2.357E-2 N	CS CL	40	2.75 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
2.59 E-2 N	CS CL	40	2.65 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
3.79 E-2 N	CS CL	40	2.10 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
2.04 E-2 N	CS2 S04	40	3.05 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
3.31 E-2 N	CS2 S04	40	2.30 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L	L
E 0	CU S04				GRAPH DATA NOT RETRIEVED	SATA IWAM	63034	R	
2.22 E 1 Q	N-C10 GLYCEROL ETHER	25	8.4 X10-2 D	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOKW	57014	T	L
			2.91 X10-3 M				M		
2.22 E 1 Q	N-C10 GLYCEROL ETHER	25	9.7 X10-2 D	CB	UNSPECIFIED CONDUCTANCE	SCHI FOKW	57014	T	L
			3.36 X10-3 M				M		
5. E 0 H	DIOXANE	15	6.73 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.0 E 1 H	DIOXANE	15	7.31 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.5 E 1 H	DIOXANE	15	9.03 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.0 E 1 H	DIOXANE	15	1.38 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2.5 E 1 H	DIOXANE	15	2.10 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
2. E 0 H	DIOXANE	25	8.01 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
3. E 0 H	DIOXANE	25	8.06 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
5. E 0 H	DIOXANE	25	7.7 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
7. E 0 H	DIOXANE	25	8.5 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3
1.0 E 1 H	DIOXANE	25	9.0 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L	3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.5 E 1 H DIOXANE	25	1.31 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
2.0 E 1 H DIOXANE	25	2.12 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
2.5 E 1 H DIOXANE	25	3.0 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
5. E 0 H DIOXANE	35	8.74 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
1.0 E 1 H DIOXANE	35	1.05 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
1.5 E 1 H DIOXANE	35	1.90 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
7. E-2 P DECANOL-1	24.7	7. X10-2 P	HD	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L	
2.22 E 1 Q 3.5(CH ₃) ₂ C ₆ H ₃ GLET*	25	1.70 X10-1 D 5.896X10-3 M	CB	UNSPECIFIED CONDUCTANCE	SCHI FOWK	57014	T L	M
2.22 E 1 Q 3.5(CH ₃) ₂ C ₆ H ₃ GLET*	25	1.4 X10-1 D 4.85 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T L	M
1.00 E 2 A DEUTERIUM OXIDE	25.0	8.05 X10-3 M	BA	SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T 2	
9.27 E 0 P ETHANOL	5	5.51 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
3. E 0 P ETHANOL	10.0	7.33 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
5. E 0 P ETHANOL	10.0	6.63 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
6. E 0 P ETHANOL	10.0	6.33 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9. E 0 P ETHANOL	10.0	5.55 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	10	5.50 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	15	5.54 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	20	5.67 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.91 E 0 H ETHANOL	20	5.18 X10-3 M	CG	EQUIV COND 1ST DEVIATION	WARD	40004	T L	
9.91 E 0 H ETHANOL	20	5.65 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	WARD	40004	P 3	
2.002E 1 H ETHANOL	20	8.5 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	WARD	40004	P 3	
2.002E 1 H ETHANOL	20	7.22 X10-3 M	CG	EQUIV COND 1ST DEVIATION	WARD	40004	T L	
2.978E 1 H ETHANOL	20	1.30 X10-2 M	CC	EQUIV CONDCTNCE GRAPH	WARD	40004	P L	
2.978E 1 H ETHANOL	20	1.04 X10-2 M	CG	EQUIV COND 1ST DEVIATION	WARD	40004	T L	
9.27 E 0 P ETHANOL	25	5.96 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	25	1.067X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	30	6.33 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	30	1.146X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	35	6.72 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	35	1.310X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	40	7.19 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	40	1.496X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	45	7.72 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	45	1.656X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	50	8.30 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	50	1.831X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
3. E 0 P ETHANOL	55.0	8.75 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
5. E 0 P ETHANOL	55.0	8.40 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
6. E 0 P ETHANOL	55.0	8.42 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9. E 0 P ETHANOL	55.0	8.92 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
9.27 E 0 P ETHANOL	55	8.96 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
1.0 E 1 P ETHANOL	55.0	9.28 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
1.5 E 1 P ETHANOL	55.0	1.160X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.0 E 1 P ETHANOL	55.0	1.505X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	55.0	2.005X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L L	
9.27 E 0 P ETHANOL	60	9.70 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
2.498E 1 P ETHANOL	60	2.170X10-2 M	BB	EQUIV CONDCTNCE GRAPH	FLOC	57013	L 3	
1.13 E 2 I HEXANOL-1	UNK	5.1 X10-3 M	BC	ULTRAFILTRATION	HUTC	59018	K L	
2. E-2 N K CL	70	3.1 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
5. E-2 N K CL	70	2.4 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
0.10 E-1 N K CL	70	1.6 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
1. E-3 N K OH	UNK	5. X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L	
4.08 E-3 N K2 S04	40	6.00 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
7.42 E-3 N K2 S04	40	5.05 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
1.50 E-2 N K2 S04	40	3.95 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
2.16 E-2 N K2 S04	40	3.45 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
2.67 E-2 N K2 S04	40	3.10 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
3.60 E-2 N K2 S04	40	2.50 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
4.53 E-2 N K2 S04	40	2.35 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
5.61 E-3 N LI2 S04	40	6.10 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
1.00 E-2 N LI2 S04	40	5.35 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
1.49 E-2 N LI2 S04	40	4.80 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
2.27 E-2 N LI2 S04	40	4.00 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
2.74 E-2 N LI2 S04	40	3.65 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
3.37 E-2 N LI2 S04	40	3.40 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
4.36 E-2 N LI2 S04	40	2.95 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
5.38 E-2 N LI2 S04	40	2.65 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L	
7. E-2 P LAURYL ALCOHOL	24.7	9. X10-2 P	HD	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L	
5. E-1 I LAURYL ALCOHOL	25	8.1 X10-3 M	BC	UNSPECIFIED CONDUCTANCE	MIUR ARIS	58023	G L	
6.67 E-1 I LAURYL ALCOHOL	25	8.1 X10-3 M	BC	UNSPECIFIED CONDUCTANCE	MIUR ARIS	58023	G L	
1. E 0 I LAURYL ALCOHOL	25	8.0 X10-3 M	BC	UNSPECIFIED CONDUCTANCE	MIUR ARIS	58023	G L	
3.33 E 0 I LAURYL ALCOHOL	25	7.0 X10-3 M	BC	UNSPECIFIED CONDUCTANCE	MIUR ARIS	58023	G L	
6.25 E 0 I LAURYL ALCOHOL	25	6.5 X10-3 M	BC	UNSPECIFIED CONDUCTANCE	MIUR ARIS	58023	G L	

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
2.22 E 1 Q	LAURYL ALCOHOL	25	6.6 X10-2 D 2.28 X10-3 M	CB	UNSPECIFIED CONDUCTANCE	SCHI FOWK	57014	T L
2.22 E 1 Q	LAURYL ALCOHOL	55	7.6 X10-2 D 2.63 X10-3 M	CB	UNSPECIFIED CONDUCTANCE	SCHI FOWK	57014	T L
1. E-3 N	MG CL2	70	3.9 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
5. E-3 N	MG CL2	70	2.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
2.5 E-2 N	MG CL2	70	1.5 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
5. E-2 N	MG CL2	70	1.25 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
1.0 E-1 N	MG CL2	70	1.0 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
2.5 E-1 N	MG CL2	70	8.2 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
1. E-1 M	NA BR	21	4.13 X10-2 D 1.432X10-3 M	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L
1. E-1 M	NA BR	21	4.1 X10-2 D 1.42 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3
3. E-1 M	NA BR	21	2.3 X10-2 D 7.97 X10-4 M	BC	TURBIDITY PLT LITE SCATR	HUIS	64047	T L
2. E-1 M	NA CL	10.0	7.7 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	63026	L L
1.00 E-1 M	NA CL	17.0	4.0 X10-2 D 1.38 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI	62009	T L
1.00 E-1 M	NA CL	18.0	4.0 X10-2 D 1.38 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI	62009	T L
1. E-2 M	NA CL	20	5.13 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T L
5. E-2 M	NA CL	20	2.24 X10-3 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
1. E-1 K	NA CL	20	1.4 X10-3 M	BC	INTERFACIAL TENSION LOGM	V VO	60025	T L
1.0 E-1 M	NA CL	20	1.41 X10-3 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
1.0 E-1 M	NA CL	20	1.51 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T L
1.00 E-1 M	NA CL	20.0	4.1 X10-2 D 1.42 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI	62009	T L
2. E-1 M	NA CL	20	7.2 X10-4 M	CC	SURFACE TENSION LOG PLOT	HARR	59001	K L
2.0 E-1 M	NA CL	20	7.59 X10-4 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T L
2.5 E-1 M	NA CL	20	5.89 X10-4 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
5.0 E-1 M	NA CL	20	3.24 X10-4 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
1. E-2 M	NA CL	21	1.62 X10-1 D 5.619X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3
1. E-2 M	NA CL	21	1.52 X10-1 D 5.272X10-3 M	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T 3
3. E-2 M	NA CL	21	9.2 X10-2 D 3.19 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 1
3. E-2 M	NA CL	21	8.7 X10-2 D 3.01 X10-3 M	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L
1. E-1 M	NA CL	21	4.2 X10-2 D 1.45 X10-3 M	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L
1. E-1 M	NA CL	21	4.3 X10-2 D 1.49 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3
3. E-1 M	NA CL	21	1.9 X10-2 D 6.59 X10-4 M	BC	TURBIDITY PLT LITE SCATR	HUIS	64047	T L
3. E-1 M	NA CL	21	2.4 X10-2 D 8.32 X10-4 M	BC	SURFACE TENSION LOG PLOT	HUIS	64047	T L
2. E-2 M	NA CL	25	1.10 X10-1 D 3.815X10-3 M	BE	DEBYE PLT LIGHT SCATTER	WILL MYSE	55005	M L
2. E-2 M	NA CL	25	1.05 X10-1 D 3.642X10-3 M	BD	TURBIDITY PLT LITE SCATR	WILL MYSE	55005	T M
3. E-2 M	NA CL	25	9.00 X10-2 D 3.121X10-3 M	BB	FOTOMTR SOLUBLZTN OROT	WILL MYSE	55005	T 1
3. E-2 M	NA CL	25	9.03 X10-2 D 3.132X10-3 M	BA	SPECFC CONDCTNCE GRAPH	WILL MYSE	55005	T 1
3. E-2 M	NA CL	25	8.9 X10-2 D 3.08 X10-3 M	BC	DEBYE PLT LIGHT SCATTER	WILL MYSE	55005	T L
3. E-2 M	NA CL	25	8.3 X10-2 D 2.87 X10-3 M	BB	TURBIDITY PLT LITE SCATR	WILL MYSE	55005	T M
3. E-2 M	NA CL	25	9.18 X10-2 D 3.184X10-3 M	BB	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T 1
7.5 E-2 M	NA CL	25	1. X10-2 D 3.4 X10-4 M	BE	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T L
1. E-1 M	NA CL	25	1.7 X10-3 M	BC	ULTRACENTRIFUGATION	ANAC JOHN	64017	T L
1. E-1 M	NA CL	25	3.0 X10-2 D 1.04 X10-3 M	BC	TURBIDITY PLT LITE SCATR	WILL MYSE	55005	M M
1. E-1 M	NA CL	25	3.7 X10-2 D 1.28 X10-3 M	BC	DEBYE PLT LIGHT SCATTER	WILL MYSE	55005	T L
1.0 E-1 M	NA CL	25	4.10 X10-2 D 1.422X10-3 M	BC	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T M
1.00 E-1 M	NA CL	25	4.15 X10-2 D 1.439X10-3 M	BC	FOTOMTR SOLUBLZTN OROT	WILL MYSE	55005	T L
1.00 E-1 M	NA CL	25	4.30 X10-2 D 1.491X10-3 M	BA	SPECFC CONDCTNCE GRAPH	WILL MYSE	55005	T 3 M

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2.	E-1 M	NA CL	25	2.8 X10-2 D 9.71 X10-4 M	BD	DEBYE PLT LIGHT SCATTER	WILL MYSE	55005	T M	L
2.	E-1 M	NA CL	25	2.5 X10-2 D 8.67 X10-4 M	BC	TURBIDITY PLT LITE SCATR	WILL MYSE	55005	T M	L
2.	E-1 M	NA CL	25.0	7.5 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
2.	E-1 M	NA CL	25.0	9.0 X10-4 M	BA	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T	3
3.	E-1 M	NA CL	25.0	7. X10-4 M	BD	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T	L
4.	E-1 M	NA CL	25	1.7 X10-2 D 5.89 X10-4 M	BD	DEBYE PLT LIGHT SCATTER	WILL MYSE	55005	T M	L
4.	E-1 M	NA CL	25	1.6 X10-2 D 5.54 X10-4 M	BD	TURBIDITY PLT LITE SCATR	WILL MYSE	55005	T M	L
4.0	E-1 M	NA CL	25	1.894X10-2 D 6.569X10-4 M	BC	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T M	L
2.05	E-3 M	NA CL	26	5.11 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
4.88	E-3 M	NA CL	26	4.47 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
9.36	E-3 M	NA CL	26	4.19 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
1.85	E-2 M	NA CL	26	3.40 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
3.11	E-2 M	NA CL	26	2.75 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
4.60	E-2 M	NA CL	26	2.11 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
7.53	E-2 M	NA CL	26	1.73 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
1.47	E-1 M	NA CL	26	1.35 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
2.43	E-1 M	NA CL	26	1.11 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
2.69	E-1 M	NA CL	26	1.10 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
3.16	E-1 M	NA CL	26	9.73 X10-4 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
3.54	E-1 M	NA CL	26	8.66 X10-4 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T	L
5.	E-2 M	NA CL	30	2.34 X10-3 M	CC	VELOCITY OF SOUND	SHIG	66010	G	L
1.	E-1 M	NA CL	30	1.63 X10-3 M	CC	VELOCITY OF SOUND	SHIG	66010	G	L
1.00	E-1 M	NA CL	30.0	4.8 X10-2 D 1.66 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
2.	E-1 M	NA CL	30	3.0 X10-2 D 1.04 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T M	L
5.	E-1 M	NA CL	30	5.9 X10-4 M	CD	VELOCITY OF SOUND	SHIG	66010	G	L
1.	E-2 M	NA CL	40	5.37 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T	L
6.75	E-2 M	NA CL	40	1.6 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	56020	T	L
1.0	E-1 M	NA CL	40	1.62 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T	L
2.0	E-1 M	NA CL	40	8.71 X10-4 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T	L
1.00	E-1 M	NA CL	50.2	7.2 X10-2 D 2.49 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
2.	E-1 M	NA CL	50	4.0 X10-2 D 1.38 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T M	L
2.	E-1 M	NA CL	55.0	1.1 X10-3 M	BD	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
1.	E-2 M	NA CL	60	6.17 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T	L
1.0	E-1 M	NA CL	60	2.04 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T	L
2.0	E-1 M	NA CL	60	1.45 X10-3 M	BB	SURFACE TNSN LINEAR PLOT	MATI PETH	58020	T	L
1.00	E-1 M	NA CL	69.8	1.10 X10-1 D 3.815X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.	E-2 N	NA CL	70	4.5 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
2.	E-2 N	NA CL	70	3.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
5.	E-2 N	NA CL	70	2.3 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
1.0	E-1 N	NA CL	70	1.6 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
2.	E-1 M	NA CL	70	6.0 X10-2 D 2.08 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T M	L
2.0	E-1 N	NA CL	70	1.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
5.0	E-1 N	NA CL	70	7.5 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L	L
6.1	E-3 M	NA CL	RM	5.7 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T	L
1.09	E-2 M	NA CL	RM	5.0 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T	L
1.83	E-2 M	NA CL	RM	4.1 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T	L
2.	E-2 M	NA CL	UNK	7.1 X10-2 D 2.46 X10-3 M	CD	TURBIDITY PLT LITE SCATR	KUSH HUBB	55003	K M	L
3.0	E-2 M	NA CL	RM	3.5 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	G	L
3.21	E-2 M	NA CL	RM	3.4 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T	L
4.	E-2 M	NA CL	UNK	6.0 X10-2 D 2.08 X10-3 M	CE	TURBIDITY PLT LITE SCATR	KUSH HUBB	55003	K M	L
5.59	E-2 M	NA CL	RM	3.0 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T	L
6.	E-2 M	NA CL	UNK	4.8 X10-2 D 1.66 X10-3 M	CE	TURBIDITY PLT LITE SCATR	KUSH HUBB	55003	K M	L
1.2	E-1 M	NA CL	UNK	2.6 X10-2 D 9.01 X10-4 M	CE	TURBIDITY PLT LITE SCATR	KUSH HUBB	55003	K M	L
2.0	E-1 M	NA CL	UNK	1.1 X10-2 D 3.81 X10-4 M	CE	TURBIDITY PLT LITE SCATR	KUSH HUBB	55003	K M	L
1.	E-2 M	NA F	21	1.61 X10-1 D 5.584X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	3
3.	E-2 M	NA F	21	8.5 X10-2 D 2.94 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
 D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality
 counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg;
 T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
1.	E-1 M	NA F	21	4.15 X10-2 D 1.439X10-3 M	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L
1.	E-1 M	NA F	21	4.2 X10-2 D 1.45 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3
3.	E-1 M	NA F	21	2.0 X10-2 D 6.93 X10-4 M	BC	TURBIDITY PLT LITE SCATR	HUIS	64047	T L
1.	E-2 M	NA I	21	1.62 X10-1 D 5.619X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3
3.	E-2 M	NA I	21	9.0 X10-2 D 3.12 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 1
1.	E-1 M	NA I	21	4.0 X10-2 D 1.38 X10-3 M	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L
1.	E-1 M	NA I	21	4.0 X10-2 D 1.38 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3
3.	E-1 M	NA I	21	2.3 X10-2 D 7.97 X10-4 M	BC	TURBIDITY PLT LITE SCATR	HUIS	64047	T L
4.	E-2 K	NA N03	20	2.43 X10-3 M	BB	SURFACE TENSION LOG PLOT	V VO	61026	TL L
1.	E-2 M	NA N03	35	5.7 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T 3
1.55	E-3 M	NA4 P207 PYRO	26	4.55 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
5.55	E-3 M	NA4 P207 PYRO	26	3.32 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
1.87	E-2 M	NA4 P207 PYRO	26	1.86 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
4.13	E-2 M	NA4 P207 PYRO	26	1.24 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
6.02	E-2 M	NA4 P207 PYRO	26	1.06 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
3.48	E-3 M	NA2 S04	26	4.54 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
6.87	E-3 M	NA2 S04	26	3.58 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
1.18	E-2 M	NA2 S04	26	3.12 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
1.81	E-2 M	NA2 S04	26	2.40 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
3.03	E-2 M	NA2 S04	26	1.97 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
4.00	E-2 M	NA2 S04	26	1.74 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
8.36	E-2 M	NA2 S04	26	1.36 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
1.44	E-1 M	NA2 S04	26	9.98 X10-4 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
5.68	E-3 N	NA2 S04	40	5.80 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
8.14	E-3 N	NA2 S04	40	5.50 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1.19	E-2 N	NA2 S04	40	4.70 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1.70	E-2 N	NA2 S04	40	4.05 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
2.32	E-2 N	NA2 S04	40	3.65 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
3.10	E-2 N	NA2 S04	40	3.10 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
3.82	E-2 N	NA2 S04	40	2.75 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
5.14	E-2 N	NA2 S04	40	2.40 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
2.	E 2 I	NA5 P3010 TRIPOLY	60	1.22 X10-1 D 4.231X10-3 M	DC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
2.22	E 1 Q	PENTADECANOL V.BR*	25	1.76 X10-1 D 6.104X10-3 M	CB	UNSPECIFIED CONDUCTANCE	SCHI FOWK	57014	T L
1.	E-5 M	PINACYANOL CL (DYE)	RM	1.93 X10-1 D 6.694X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MUKE MYSE	55015	G L
1.	E-5 M	PINACYANOL CL (DYE)	RM	2.01 X10-1 D 6.971X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MUKE MYSE	55015	G L
4.	E-5 M	PINACYANOL CL (DYE)	RM	2.03 X10-1 D 7.041X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MUKE MYSE	55015	G L
4.	E-5 M	PINACYANOL CL (DYE)	RM	2.10 X10-1 D 7.284X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MUKE MYSE	55015	G L
1.	E-4 M	PINACYANOL CL (DYE)	RM	2.15 X10-1 D 7.457X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MUKE MYSE	55015	G L
1.	E-4 M	PINACYANOL CL (DYE)	RM	2.29 X10-1 D 7.943X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MUKE MYSE	55015	G L
2.	E-6 M	PINACYANOL CL (DYE)	RM	1.84 X10-1 D 6.382X10-3 M	BG	FOTOMTR SPCTR CHNCE PNCN	MUKE MYSE	55015	G L
2.	E-6 M	PINACYANOL CL (DYE)	RM	1.91 X10-1 D 6.625X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	MUKE MYSE	55015	G L
5.03	E 0 H	PROPANOL-1	00.5	4.7 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.03	E 0 H	PROPANOL-1	10.5	4.0 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.03	E 0 H	PROPANOL-1	25.2	3.8 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
9.21	E 0 H	PROPANOL-1	25.2	1.31 X10-3 M	BG	AVER COND BEGINING MAXIM	FLOC UBBE	53008	G L
1.339E 1 H	PROPANOL-1	25.2	6.0 X10-4 M	BG	AVER COND BEGINING MAXIM	FLOC UBBE	53008	G L	
1.914E 1 H	PROPANOL-1	25.2	5.0 X10-4 M	BG	AVER COND BEGINING MAXIM	FLOC UBBE	53008	G L	
5.03	E 0 H	PROPANOL-1	33.5	4.1 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.03	E 0 H	PROPANOL-1	40.1	4.4 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.03	E 0 H	PROPANOL-1	50.0	5.1 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.71	E 0 H	PROPANOL-2	00.5	5.7 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.71	E 0 H	PROPANOL-2	10.5	4.9 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.71	E 0 H	PROPANOL-2	25.2	4.3 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	T 3
1.040E 1 H	PROPANOL-2	25.2	2.23 X10-3 M	BG	AVER COND BEGINING MAXIM	FLOC UBBE	53008	G L	
1.348E 1 H	PROPANOL-2	25.2	1.83 X10-3 M	BG	AVER COND BEGINING MAXIM	FLOC UBBE	53008	G L	
1.741E 1 H	PROPANOL-2	25.2	1.34 X10-3 M	BG	AVER COND BEGINING MAXIM	FLOC UBBE	53008	G L	
5.71	E 0 H	PROPANOL-2	33.5	4.5 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
5.71 E 0 H	PROPANOL-2	40.1	5.0 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.71 E 0 H	PROPANOL-2	50.0	5.7 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
5.06 E 0 H	PROPIONIC ACID	25.2	4.3 X10-3 M	BB	AVER SP EQUIV COND	FLOC UBBE	53008	G 3
1.042E 1 H	PROPIONIC ACID	25.2	4.0 X10-3 M	BD	AVER SP EQUIV COND	FLOC UBBE	53008	G L
1.519E 1 H	PROPIONIC ACID	25.2	5.5 X10-3 M	BE	AVER SP EQUIV COND	FLOC UBBE	53008	G L
5.00 E 2 Y	PRESSURE	25	9.09 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3
1.000E 3 Y	PRESSURE	25	9.45 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3
1.500E 3 Y	PRESSURE	25	9.36 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3
2.000E 3 Y	PRESSURE	25	9.09 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	HAMA	62036	G 3
1. E-3 N	PHENYL (CH3)3 N I	40	4.77 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1.4 E-3 N	PHENYL (CH3)3 N I	40	4.35 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1.9 E-3 N	PHENYL (CH3)3 N I	40	3.8 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
2.5 E-3 N	PHENYL (CH3)3 N I	40	3.47 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
3.3 E-3 N	PHENYL (CH3)3 N I	40	3.1 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
4.4 E-3 N	PHENYL (CH3)3 N I	40	2.8 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
7.5 E-3 N	PHENYL (CH3)3 N I	40	2.47 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1.3 E-2 N	PHENYL (CH3)3 N I	40	2.15 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1.86 E-2 N	PHENYL (CH3)3 N I	40	1.97 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1. E-5 M	RHODAMINE 6GPC	40	6.7 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	56020	T L
5. E-6 M	RHODAMINE 6GPC	40	6.6 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	56020	T L
3. E 0 D	SUCROSE	25	7.1 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	NAKA KAWA	64034	G 3
1.0 E 1 D	SUCROSE	25	6.1 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	NAKA KAWA	64034	G L
2.0 E 1 D	SUCROSE	25	5.5 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	NAKA KAWA	64034	G L
3.0 E 1 D	SUCROSE	25	5.1 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	NAKA KAWA	64034	G L
5. E-2 P	TRIBUTYL PHOSPHATE	24.7	1.3 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L
2.0 E-1 P	TRIBUTYL PHOSPHATE	24.7	1.8 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L
4.0 E-1 P	TRIBUTYL PHOSPHATE	24.7	1.6 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L
1.2 E-3 N	(CH3)4 N I	40	6.15 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
2.85 E-3 N	(CH3)4 N I	40	5.47 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
4.7 E-3 N	(CH3)4 N I	40	4.9 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
7.7 E-3 N	(CH3)4 N I	40	4.1 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
1.3 E-2 N	(CH3)4 N I	40	3.3 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	GODD HARV	53012	L L
3. E 0 M	UREA	10	1.06 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
6. E 0 M	UREA	10	1.41 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
1. E 0 M	UREA	25	7.4 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
2. E 0 M	UREA	25	8.0 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
3. E 0 M	UREA	25	9. X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
4.5 E 0 M	UREA	25	1.02 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
6. E 0 M	UREA	25	1.20 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
5. E-3 M	UREA	26	6.2 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
2.5 E-2 M	UREA	26	6.2 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
4.7 E-2 M	UREA	26	6.2 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
1.22 E-1 M	UREA	26	6.2 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
1.63 E-1 M	UREA	26	6.1 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
2.35 E-1 M	UREA	26	6.2 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
2.78 E-1 M	UREA	26	6.1 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
4.70 E-1 M	UREA	26	6.1 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	G L
3.22 E 0 M	UREA	26	5.72 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L
3. E 0 M	UREA	45	9.1 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
6. E 0 M	UREA	45	1.15 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L
0003				SEE CMPD NMBR IN ADDITV	SHED JAKO	63001	X	
1.078E 1 C	0003	25.0	8.32 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3
2.006E 1 C	0003	25.0	8.54 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3
2.025E 1 C	0003	25.0	8.54 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3
3.478E 1 C	0003	25.0	9.88 X10-3 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3
4.912E 1 C	0003	25.0	1.115X10-2 M	CA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 3
5.998E 1 C	0003	25.0	1.281X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2
7.100E 1 C	0003	25.0	1.498X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2
8.466E 1 C	0003	25.0	1.91 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2
9.495E 1 C	0003	25.0	2.65 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	MYSE OTTE	61017	T 2
E 0	0004	50		GRAPH DATA NOT RETRIEVED	LANG	53005	R	
1.25 E 1 C	0004	RM	5.11 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T 3
2.5 E 1 C	0004	RM	3.88 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T 3
5.0 E 1 C	0004	RM	2.62 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T 3
E 0	0005	50		SEE CMPD NMBR IN ADDITV	LANG	53005	X	
		50		GRAPH DATA NOT RETRIEVED	LANG	53005	R	
2. E 0 C	0024	70	5.7 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
5. E 0 C	0024	70	5.3 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
1.0 E 1 N	0024	70	4.9 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
2.5 E 1 C	0024	70	4.0 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
5.0 E 1 C	0024	70	3.4 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
7.5 E 1 C	0024	70	3.1 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L
E 0	0064	50		GRAPH DATA NOT RETRIEVED	LANG	53005	R	
1.25 E 1 C	0091	UNK	6.8 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
2.98 E 1 C	0091	UNK	7.8 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
3.74 E 1 C 0091	UNK	8.3 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
4.97 E 1 C 0091	UNK	9.4 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
6.53 E 1 C 0091	UNK	1.32 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
7.27 E 1 C 0091	UNK	1.59 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
7.84 E 1 C 0091	UNK	1.82 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
8.27 E 1 C 0091	UNK	1.91 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
3.80 E 1 C 0092	25	6.4 X10-3 M	CC	REFRACTIVE INDEX	KLEV	48005	T L
6.45 E 1 C 0092	25	6.2 X10-3 M	CC	REFRACTIVE INDEX	KLEV	48005	T L
1.48 E 1 C 0092	UNK	6.1 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
2.07 E 1 C 0092	UNK	6.1 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
3.34 E 1 C 0092	UNK	6.2 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
4.54 E 1 C 0092	UNK	6.2 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
5.69 E 1 C 0092	UNK	6.3 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
6.89 E 1 C 0092	UNK	6.4 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
8.41 E 1 C 0092	UNK	6.5 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
9.30 E 1 C 0092	UNK	6.4 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L
0115			SEE CMPD NMBR IN ADDITV	SCHI MANN	66001	X	
0116			SEE CMPD NMBR IN ADDITV	SCHI MANN	66001	X	
0205			SEE CMPD NMBR IN ADDITV	KURI NAKA	62010	X	
0325			SEE CMPD NMBR IN ADDITV	SCHI MANN	66001	X	
0327			SEE CMPD NMBR IN ADDITV	SCHI MANN	66001	X	
0417			SEE CMPD NMBR IN ADDITV	KLEV RAIS	54010	X	
7. E-3 M PENTANOL-1	UNK	4. X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L
1. E-3 N K OH							
2.5 E-2 M PENTANOL-1	UNK	2.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L
1. E-3 N K OH							
9. E-3 M H CL	25	3.7 X10-3 M	CC	SPECFC CONDCTNCE GRAPH	JAME PETH	60020	T L
1. E-3 M NA CL							
2. E-1 IONIC STRENGTH	25	3. X10-2 D	HG	REACTN RATE SULUBILIZATE	TONG REEV	65030	T L
1.01 E 1 PH OF SOLUTION							
5. E 1 E N-HEPTANE	UNK	4.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L
1. E-3 N K OH							
1. E-1 K NA ION	UNK	1.45 X10-3 M	BB	SURFACE TENSION LOG PLOT	ROE BRAS	54013	T L
6.5 E 0 PH OF SOLUTION							
9.25 E-3 M NA CL	RM	5.1 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T L
3.1 E-3 M K CL							
1.13 E-2 M NA CL	RM	4.8 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T L
4.9 E-3 M K CL							
2.06 E-2 M NA CL	RM	3.6 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T L
7.75 E-3 M K CL							
2.38 E-2 M NA CL	RM	3.8 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T L
2.4 E-3 M K CL							
4.44 E-2 M NA CL	RM	3.3 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T L
3.2 E-3 M K CL							
4.94 E-2 M NA CL	RM	2.9 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	T L
4.8 E-3 M K CL							
2.5 E-3 M NA CL	RM	7.1 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	G L
2.7 E-2 M LI CL							
1.2 E-2 M NA CL	RM	7.1 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	G L
1.8 E-2 M LI CL							
1.6 E-2 M NA CL	RM	7.0 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	G L
1.4 E-2 M LI CL							
2.16 E-2 M NA CL	RM	5.8 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	G L
8.4 E-3 M LI CL							
2.52 E-2 M NA CL	RM	5.3 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	G L
4.8 E-3 M LI CL							
2.75 E-2 M NA CL	RM	3.8 X10-3 M	CG	VISUAL FLUOR CHNGE RHD6	PRIN HERM	56002	G L
2.5 E-3 M LI CL							
3. E-3 M OCTANOL-1	UNK	2.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L
E-3 N K OH							
5. E-3 M OCTANOL-1	UNK	1.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L
1. E-3 N K OH							
7. E-3 M OCTANOL-1	UNK	6. X10-4 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L
1. E-3 N K OH							
2.5 E-2 M PENTAMINE	UNK	1.8 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L
1. E-3 N K OH							
558 ENTRIES FOR COMPOUND							
COMPOUND NO = 2 MOL WGT -	232.2	SODIUM OCTYL 1 SULFATE					
	10	1.421X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3
	15	1.367X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3
	20	1.33 X10-1 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L 3
	20	1.337X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation	
	21	2.96 X10 ⁻¹ 0 D 1.274X10 ⁻¹ M	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T M	L	
	21	3.10 X10 ⁻¹ 0 D 1.335X10 ⁻¹ M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T M	3	
	25	7. X10 ⁻² N	CD	SPECFC CONDCTNCE GRAPH	HAFF PICC	42003	T L		
	25	1.3 X10 ⁻¹ N	CC	UNSPECIFIED CONDUCTANCE	HAFF PICC	42003	T L		
	25	1.303X10 ⁻¹ W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3		
	25	3.0 X10 ⁻¹ 0 D	BD	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T L		
		1.29 X10 ⁻¹ M					M		
	30.0	1.30 X10 ⁻¹ M	CB	VELOCITY OF SOUND	SHIG	65022	T 3		
	30	1.318X10 ⁻¹ W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3		
	35	1.342X10 ⁻¹ W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3		
	40	1.363X10 ⁻¹ W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3		
	40.0	1.36 X10 ⁻¹ M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3		
	45	1.381X10 ⁻¹ W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3		
	50	9.8 X10 ⁻² M	BB	INTERFACIAL TENSION LOGM	KLIN LANG	57022	T L		
	50	1.434X10 ⁻¹ W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3		
	55	1.463X10 ⁻¹ W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 3		
	UNK	2.30 X10 ⁻¹ D	XG	VISUAL SPCTR CHNGE PNCN	DEMC	61031	T L		
		9.905X10 ⁻¹ M							
5. E-2 M	NA CL	20	1.12 X10 ⁻¹ M	BB	INTERFACIAL TENSION LOGM	HYD TAYL	62004	L L	
1.0 E-1 M	NA CL	20	9.75 X10 ⁻² M	BB	INTERFACIAL TENSION LOGM	HYD TAYL	62004	L L	
2.5 E-1 M	NA CL	20	7.08 X10 ⁻² M	BB	INTERFACIAL TENSION LOGM	HYD TAYL	62004	L L	
5.0 E-1 M	NA CL	20	5.01 X10 ⁻² M	BB	INTERFACIAL TENSION LOGM	HYD TAYL	62004	L L	
1. E-2 M	NA CL	21	2.90 X10 ⁻¹ D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L	
		1.248X10 ⁻¹ M					M		
3. E-2 M	NA CL	21	2.70 X10 ⁻¹ D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L	
		1.162X10 ⁻¹ M					M		
3. E-2 M	NA CL	21	2.80 X10 ⁻¹ D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	
		1.205X10 ⁻¹ M					M		
1. E-1 M	NA CL	21	2.37 X10 ⁻¹ D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	
		1.020X10 ⁻¹ M					M		
1. E-1 M	NA CL	21	2.21 X10 ⁻¹ D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L	
		9.517X10 ⁻² M					M		
3. E-1 M	NA CL	21	1.49 X10 ⁻¹ D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T L	
		6.416X10 ⁻² M					M		
3. E-1 M	NA CL	21	1.60 X10 ⁻¹ D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	
		6.890X10 ⁻² M					M		
1. E 0 M	NA CL	21	8.0 X10 ⁻¹ D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3	
		3.44 X10 ⁻² M					M		

32 ENTRIES FOR COMPOUND

COMPOUND NO =	3 MOL WGT -	260.3 SODIUM DECYL 1 SULFATE	VALUES FRM REF IN CMC	MYSE OTTE	61017	R
		61005	VALUES FRM REF IN CMC	FLOC	61007	L 3
	0	3.88 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 3
	5	3.64 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 3
	10	3.48 X10 ⁻² W	BA SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L D
	10	3.50 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L D
	15	3.41 X10 ⁻² W	BA SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1
	15	3.39 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 1
	20	3.35 X10 ⁻² W	BA SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1
	20	3.31 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 1
	21	7.8 X10 ⁻¹ D	BB SURFACE TENSION LOG PLOT	HUIS	64047	T L
		2.99 X10 ⁻² M				M
	21	7.9 X10 ⁻¹ D	BB TURBIDITY PLT LITE SCATR	HUIS	64047	T L
		3.03 X10 ⁻² M				M
	23	3.1 X10 ⁻² N	CC UNSPECIFIED CONDUCTANCE	TART LELO	55021	T L
	25	3.1 X10 ⁻² N	CC UNSPECIFIED CONDUCTANCE	HAFF PICC	42003	T L
	25	3.32 X10 ⁻² W	BA SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1
	25.0	3.26 X10 ⁻² M	BA EQUIV CONDCTNCE GRAPH	MYSE KAPA	61005	T 1
	25.0	3.35 X10 ⁻² M	BA SPECFC CONDCTNCE GRAPH	MYSE KAPA	61005	T 1
	25	3.27 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 1
	25	8.4 X10 ⁻¹ D	DB TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T L
		3.22 X10 ⁻² M				M
	25	8.5 X10 ⁻¹ D	DB REFRACTIVE INDEX	PRIN HERM	56011	T L
		3.26 X10 ⁻² M				M
	25	3.1 X10 ⁻² M	BB SURFACE TENSION LOG PLOT	SCHI	64020	T L
	25.0	3.32 X10 ⁻² M	BA SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T 1
	30	3.31 X10 ⁻² W	BA SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1
	30	3.26 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 1
	35	3.27 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 1
	35	3.35 X10 ⁻² W	BA SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1
	40	3.32 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L 1
	40	3.41 X10 ⁻² W	BA SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L 1
	45	3.38 X10 ⁻² M	BA AVER SP EQUIV COND	FLOC	61007	L D

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/l
kg; T—wt % surfactant mixture; U—mol/(l orgkg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation	
	45	3.49 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D	
	50	3.2 X10-2 M	BB	INTERFACIAL TENSION LOGM	KLIN LANG	57022	T	L	
	50	3.47 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	D	
	50	3.4 X10-2 M	BB	SURFACE TENSION LOG PLOT	KLIN LANG	57022	T	L	
	50	3.64 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D	
	55	3.59 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	D	
	55	3.78 X10-2 W	BA	SPECFC CONDCTNCE EQUATNS	GODD BENS	57011	L	D	
	60	3.73 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3	
	65	3.88 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3	
	90	4.39 X10-2 W	CA	EQUIV CONDCTNCE GRAPH	KURZ	62040	T	L	
UNK	3.0	X10-2 M	BC	ULTRAFILTRATION	HUTC	59018	K	L	
RM	3.37	X10-2 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	L	
UNK	6.8	X10-1 D	XG	VISUAL SPCTR CHNGE PNCF	DEMC	61031	T	L	
		2.61 X10-2 M					M		
1.00 E 2 A	DEUTERIUM OXIDE	25.0	3.25 X10-2 M	BA	SPECFC CONDCTNCE EQUATNS	MUKE KAPA	66002	T	2
6.6 E 1 I	HEXANOL-1	UNK	2.5 X10-2 M	BC	ULTRAFILTRATION	HUTC	59018	K	L
1.16 E 2 I	HEXANOL-1	UNK	2.2 X10-2 M	BC	ULTRAFILTRATION	HUTC	59018	K	L
1.96 E 2 I	HEXANOL-1	UNK	1.8 X10-2 M	BC	ULTRAFILTRATION	HUTC	59018	K	L
1.0 E-1 M	NA CL	20	1.41 X10-2 M	BB	INTERFACIAL TENSION LOGM	HYAD TAYL	62004	L	L
2.5 E-1 M	NA CL	20	7.41 X10-3 M	BB	INTERFACIAL TENSION LOGM	HYAD TAYL	62004	L	L
5.0 E-1 M	NA CL	20	4.42 X10-3 M	BB	INTERFACIAL TENSION LOGM	HYAD TAYL	62004	L	L
1. E-2 M	NA CL	21	6.9 X10-1 D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T	L
			2.65 X10-2 M				M		
1. E-2 M	NA CL	21	7.0 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	L
			2.68 X10-2 M				M		
3. E-2 M	NA CL	21	5.4 X10-1 D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T	L
			2.07 X10-2 M				M		
3. E-2 M	NA CL	21	5.6 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	L
			2.15 X10-2 M				M		
1. E-1 M	NA CL	21	3.5 X10-1 D	BB	SURFACE TENSION LOG PLOT	HUIS	64047	T	L
			1.34 X10-2 M				M		
1. E-1 M	NA CL	21	3.45 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	L
			1.325X10-2 M				M		
3. E-1 M	NA CL	21	1.8 X10-1 D	BC	SURFACE TENSION LOG PLOT	HUIS	64047	T	L
			6.91 X10-3 M				M		
3. E-1 M	NA CL	21	1.8 X10-1 D	BC	TURBIDITY PLT LITE SCATR	HUIS	64047	T	L
			6.91 X10-3 M				M		
1.00 E-2 M	NA CL	25.0	3.02 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	MYSE KAPA	61005	L	3
7.5 E-2 M	NA CL	25	4.2 X10-1 D	DC	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T	L
			1.61 X10-2 M				M		
1.00 E-1 M	NA CL	25.0	1.51 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	MYSE KAPA	61005	L	3
3. E-1 M	NA CL	25.0	7.3 X10-3 M	BD	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T	L
3.00 E-1 M	NA CL	25.0	8.0 X10-3 M	BD	SPECFC CONDCTNCE GRAPH	MYSE KAPA	61005	L	L
4. E-1 M	NA CL	25.0	6.5 X10-3 M	BD	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T	L
6. E-1 M	NA CL	25.0	4.6 X10-3 M	BC	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T	L
1.0 E 0 M	NA CL	25.0	2.77 X10-3 M	BB	SPECFC CONDCTNCE GRAPH	MYSE MYSE	65018	T	3
1.2 E 0 M	NA CL	25	4. X10-2 D	DD	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T	L
			1.5 X10-3 M				M		
4.98 E-2 W	NA CLO4	90	3.66 X10-2 W	CB	UNSPECIFIED CONDUCTANCE	KURZ	62040	T	L
9.89 E-2 W	NA CLO4	90	2.94 X10-2 W	CB	UNSPECIFIED CONDUCTANCE	KURZ	62040	T	L
2.002E-1 W	NA CLO4	90	2.34 X10-2 W	CB	UNSPECIFIED CONDUCTANCE	KURZ	62040	T	L
1. E 0 M	UREA	25	3.2 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
2. E 0 M	UREA	25	3.5 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3. E 0 M	UREA	25	3.7 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
4.5 E 0 M	UREA	25	4.3 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6. E 0 M	UREA	25	5.0 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
	0001			SEE CMPD NMBR IN ADDITV	MYSE OTTE	61017	X		
1.25 E 1 C	0001	RM	2.02 X10-2 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
2.5 E 1 C	0001	RM	1.56 X10-2 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
5.0 E 1 C	0001	RM	1.08 X10-2 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	3
7.5 E 1 C	0001	RM	8.55 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T	L
5.00 E 1 C	0004	25	3.03 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	30	3.09 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	35	3.18 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	40	3.31 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2
5.00 E 1 C	0004	45	3.47 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	50	3.66 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	55	3.87 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	60	4.16 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	65	4.51 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	70	4.90 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
5.00 E 1 C	0004	75	5.35 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3
89 ENTRIES FOR COMPOUND									
COMPOUND NO =	4	MOL WGT -	316.4	SODIUM TETRADECYL 1 SULFATE					
			53005	VALUES FRM REF IN CMC	KLIN LANG	57022	R		
			21.5 2.21 X10-3 M	CB EQUIV CONDCTNCE GRAPH	FLOC UBBE	53008	K	L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation	
	25.2	2.08 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	FLOC UBBE	53008	K L		
	25	1.4 X10-3 M	CD	SPECFC CONDCTNCE GRAPH	BENT SPAR	66038	T L		
	25	2.05 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 2		
	30.0	2.2 X10-3 M	CC	VELOCITY OF SOUND	SHIG	65022	T L		
	30	2.08 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 2		
	35	2.15 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 2		
	40	2.4 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	WINS	48008	L L		
	40.1	2.07 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	FLOC UBBE	53008	K L		
	40	2.21 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 1		
	40	2.21 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 1		
	40.0	2.40 X10-3 M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T L		
	45	2.31 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 2		
	50	2.1 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	LANG	53005	T L		
	50	2.2 X10-3 M	BB	INTERFACIAL TENSION LOGM	KLIN LANG	57022	T L		
	50	2.0 X10-3 M	BB	SURFACE TENSION LOG PLOT	KLIN LANG	57022	T L		
	50	2.54 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L		
	50	2.43 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3		
	50	6. X10-2 D	BD	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T L		
		1.8 X10-3 M				M			
	50	1.32 X10-3 M	BG	VISUAL SPCTR CHNGE PNPN	WEIL STIR	63013	T L		
	55	2.58 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3		
	60	2.80 X10-3 M	CB	UNSPECIFIED CONDUCTANCE	WINS	48008	L L		
	60	2.77 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3		
	65	2.99 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3		
	70	3.22 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3		
	75	3.50 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3		
	RM	1.66 X10-3 M	CG	VISUAL SPCTR CHNGE PNPN	WINS	48008	T L		
	RM	1.83 X10-3 M	BB	ELECTROMOTIVE FORCE	SHED JAKO	63001	T L		
	UNK	5.0 X10-2 D	XG	VISUAL SPCTR CHNGE PNPN	DEMC	61031	T L		
		1.58 X10-3 M				M			
5. E O H	DIOXANE	40	2.4 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
1.0 E 1 H	DIOXANE	40	2.9 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
1.5 E 1 H	DIOXANE	40	3.8 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
2.0 E 1 H	DIOXANE	40	5.2 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
2.5 E 1 H	DIOXANE	40	7.5 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
3.0 E 1 H	DIOXANE	40	1.28 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
3.5 E 1 H	DIOXANE	40	1.77 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
1. E-2 N	NA CL	23	1.9 X10-3 N	CC	UNSPECIFIED CONDUCTANCE	TART LELO	55021	T L	
. E 0	NA CL	50			THEORETICALLY ESTIMATED	PRIN HERM	56011	R	
	0001				SEE CMPD NMBR IN ADDITV	SHED JAKO	63001	X	
. E 0	0001	50			GRAPH DATA NOT RETRIEVED	LANG	53005	R	
	0003				SEE CMPD NMBR IN ADDITV	FLOC	61007	X	
42 ENTRIES FOR COMPOUND									
COMPOUND NO = 5 MOL WGT -		344.4	SODIUM HEXADECYL 1 SULFATE						
		53005		VALUES FRM REF IN CMC	KLIN LANG	57022	R		
	25	2.1 X10-4 M	CD	SPECFC CONDCTNCE GRAPH	BENT SPAR	66038	T L		
	30	4. X10-4 M	CD	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L		
	30	4.1 X10-4 M	BG	VISUAL SPCTR CHNGE PNPN	WEIL STIR	59004	T L		
	35	2. X10-4 M	XE	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L		
	35.8	4.4 X10-4 M	CG	FOTOMTR SPCTR CHNGE PNPN	CORR KLEV	46010	T L		
	40	5.2 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3		
	40.0	5.8 X10-4 M	BC	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3		
	45	6.0 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	64020	T L		
	50	5.4 X10-4 M	CC	EQUIV CONDCTNCE GRAPH	LANG	53005	T L		
	50	6.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	KLIN LANG	57022	T L		
	50	6.5 X10-4 M	BB	INTERFACIAL TENSION LOGM	KLIN LANG	57022	T L		
	50	6.65 X10-4 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L		
	50	2. X10-2 D	BE	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T L		
		5.8 X10-4 M				M			
	50	4.2 X10-4 M	BG	VISUAL SPCTR CHNGE PNPN	WEIL STIR	63013	T L		
	50	4.2 X10-4 M	BG	VISUAL SPCTR CHNGE PNPN	WEIL STIR	59004	T L		
	60	8. X10-4 M	XE	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L		
	90	1.0 X10-3 M	XD	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L		
	UNK	2.0 X10-2 D	XG	VISUAL SPCTR CHNGE PNPN	DEMC	61031	T L		
		5.80 X10-4 M				M			
5. E O H	DIOXANE	40	1.27 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
1.0 E 1 H	DIOXANE	40	2.0 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
1.5 E 1 H	DIOXANE	40	2.8 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
2.0 E 1 H	DIOXANE	40	3.54 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
2.5 E 1 H	DIOXANE	40	4.3 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
3.0 E 1 H	DIOXANE	40	5.0 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	SHIR MATU	65020	L 3	
. E 0	NA CL	50			GRAPH DATA NOT RETRIEVED	LANG	53005	R	
. E 0	NA CL	50			THEORETICALLY ESTIMATED	PRIN HERM	56011	R	
1. E 1 I	NA CL	50	5.4 X10-4 M	CC	EQUIV CONDCTNCE GRAPH	LANG	53005	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2.	E 1 I	NA CL	50	5.35 X10-4 M	CB	EQUIV CONDCTNCE GRAPH	LANG	53005	T L
3.	E O M	UREA	45	6.4 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	64020	T L
6.	E O M	UREA	45	7.9 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	64020	T L
.	E O	0001	50			GRAPH DATA NOT RETRIEVED	LANG	53005	R
1.	E 1 C	0001	50	6.0 X10-4 M	CC	EQUIV CONDCTNCE GRAPH	LANG	53005	T L
2.	E 1 C	0001	50	6.5 X10-4 M	CC	EQUIV CONDCTNCE GRAPH	LANG	53005	T L
34 ENTRIES FOR COMPOUND									
COMPOUND NO =	6	MOL WGT -	260.2	SODIUM ALPHA SULFOPELARGONIC ACID					
	UNK		1.00 X10 0 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	60008	T L	
			3.843X10-2 M					M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	7	MOL WGT -	372.4	SODIUM OCTYL ALPHA SULFOPELARGONATE					
	UNK		2.1 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
	UNK		8. X10-2 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	60008	T L	
			2.1 X10-3 M					M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO =	8	MOL WGT -	484.6	DISODIUM ALPHA SULFOPHENYLSTEARATE					
	UNK		5.6 X10-2 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	60008	T L	
			1.15 X10-3 M					M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	9	MOL WGT -	386.5	SODIUM ALPHA SULFOSTEARIC ACID					
	25			QUESTIONABLE CRITERION		WEIL STIR	60008	R	
	25			QUESTIONABLE CRITERION		WEIL STIR	63013	R	
2 ENTRIES FOR COMPOUND									
COMPOUND NO =	10	MOL WGT -	408.5	DISODIUM ALPHA SULFOSTEARATE					
	25			QUESTIONABLE CRITERION		WEIL STIR	60008	R	
	25			QUESTIONABLE CRITERION		WEIL STIR	63013	R	
2 ENTRIES FOR COMPOUND									
COMPOUND NO =	11	MOL WGT -	462.6	SODIUM ALPHA SULFO PHENYL STEARIC ACID					
	UNK		5. X10-3 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	60008	T L	
			1.0 X10-4 M					M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	12	MOL WGT -	455.4	SODIUM 9,10 DICHLORO ALPHA SULFOSTEARIC ACID					
	UNK		1.7 X10-2 D	FG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	60008	T L	
			3.73 X10-4 M					M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	13	MOL WGT -	477.3	DISODIUM 9,10 DICHLORO ALPHA SULFOSTEARATE					
	UNK		1.5 X10-1 D	FG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	60008	T L	
			3.14 X10-3 M					M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	14	MOL WGT -	440.5	DISODIUM 9,10 DIHYDROXY ALPHA SULFOSTEARATE					
	UNK		5.9 X10-1 D	FG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	60008	T L	
			1.33 X10-2 M					M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	15	MOL WGT -	260.3	SODIUM DECYL 2 SULFATE					
	10		5.15 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	15		4.92 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	20		4.70 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	25		4.56 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	30		4.52 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	35		4.50 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	40		4.50 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	40.0		4.95 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T L	
	45		4.52 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	50		4.57 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	
	55		4.65 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L 3	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/g; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation		
	60	4.79 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	65	4.95 X10-2 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
13 ENTRIES FOR COMPOUND										
COMPOUND NO = 16 MOL WGT -	316.4	SODIUM TETRADECYL 2 SULFATE								
	25	3.27 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2		
	30	3.28 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2		
	35	3.31 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	2		
	40	3.3 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	WINS	48008	L	L		
	40.0	3.30 X10-3 M	BB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T	L		
	40	3.38 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	45	3.48 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	50	3.64 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	55	3.83 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	60	3.75 X10-3 M	CB	UNSPECIFIED CONDUCTANCE	WINS	48008	L	L		
	60	4.04 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	65	4.29 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	70	4.60 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	75	5.00 X10-3 M	BA	AVER SP EQUIV COND	FLOC	61007	L	3		
	RM	3.26 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	WINS	48008	T	L		
15 ENTRIES FOR COMPOUND										
COMPOUND NO = 17 MOL WGT -	316.4	SODIUM TETRADECYL 4 SULFATE								
	25	5.12 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	30	5.05 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	35	5.04 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	40	5.2 X10-3 M	CB	UNSPECIFIED CONDUCTANCE	WINS	48008	L	L		
	40.0	5.15 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	EVAN	56006	T	3		
	40	5.12 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	45	5.23 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	50	5.38 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	55	5.57 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	60	6.10 X10-3 M	CB	UNSPECIFIED CONDUCTANCE	WINS	48008	L	L		
	60	5.85 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	65	6.21 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	70	6.62 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	75	7.11 X10-3 M	CA	AVER SP EQUIV COND	FLOC	61007	L	3		
	RM	4.76 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	WINS	48008	T	L		
15 ENTRIES FOR COMPOUND										
COMPOUND NO = 18 MOL WGT -	292.4	OCTYL BETA D GLUCOSIDE								
	20	2.6 X10-2 W	BD	SURFACE TENSION LOG PLOT	BURY BROW	52011	T	L		
	25	61008		VALUES FRM REF IN CMC	SHIN YAMA	59013	T	R		
	25	2.5 X10-2 M	BB	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	3		
	30	2.3 X10-2 W	BC	SURFACE TENSION LOG PLOT	BURY BROW	52011	T	L		
9.3 E-1 N	CA CL2	25	1.7 X10-2 M	BB	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	L	
4.7 E-1 N	NA CL	25	1.7 X10-2 M	BB	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	L	
9.3 E-1 N	NA CL	25	1.2 X10-2 M	BC	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	L	
9.3 E-1 N	NA2 SO4	25	9. X10-3 M	BC	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	L	
8.9 E O C	0019	25	1.25 X10-2 M	BB	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	L	
3.4 E-1 C	0020	25	1.5 X10-2 M	BC	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	L	
10 ENTRIES FOR COMPOUND										
COMPOUND NO = 19 MOL WGT -	320.4	DECYL BETA D GLUCOSIDE								
	25	2.2 X10-3 M	BB	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	3		
0018				SEE CMPD NMBR IN ADDITV	SHIN YAMA	61008	T	X		
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 20 MOL WGT -	348.5	DODECYL BETA D GLUCOSIDE								
0018			25	1.9 X10-4 M	BB	SURFACE TENSION LOG PLOT	SHIN YAMA	61008	T	3
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 21 MOL WGT -	229.4	DIMETHYL DODECYL AMINE OXIDE								
	62005			VALUES FRM REF IN CMC	HERR	64006		R		
01.0	2.84 X10-3 M	BB		TURBIDITY PLT LITE SCATR	HERR	62005	T	L		
	2. X10-3 M	BE		HEAT OF DILUTION	BENJ	64016	L	L		
	26.5	4.3 X10-1 P	BC	HEAT OF DILUTION	BENJ	66012	L	L		
	1.87 X10-2 S					M				
	27.0	2.10 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HERR	62005	T	3		
	30	4.5 X10-2 P	BC	DENSITY	BENJ	66040	T	L		
	1.96 X10-3 S					M				

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; F – wt % solvent; G – mol % surfactant; H – normality; I – mol % surfactant mixture; J – normality; K – counterions; L – molar; M – normal; N – normal; O – wt % surfactant; P – wt % surfactant mixture; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives			Temp. °C	CMC	Qual. Meth.	Method	Authors	Reference	Source	Evaluation
2.5	E 0 B	BUTANOL-1	40.0 50.0 26.5	1.83 X10-3 M 1.75 X10-3 M 2.2 X10-1 P	BB BB BC	TURBIDITY PLT LITE SCATR TURBIDITY PLT LITE SCATR HEAT OF DILUTION	HERR HERR BENJ	62005 62005 66012	T T L	L
5.	E 0 B	BUTANOL-1	26.5	9.59 X10-3 S	BD	HEAT OF DILUTION	BENJ	66012	L	L
7.	E 0 B	BUTANOL-1	26.5	5.23 X10-3 S 6. X10-2 P 2.6 X10-3 S	BE	HEAT OF DILUTION	BENJ	66012	L	L
5.	E 0 B	ETHANOL	26.5	1.2 X10-1 P 1.70 X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
1.0	E 1 B	ETHANOL	26.5	3.4 X10-1 P 1.48 X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
1.0	E 1 B	METHANOL	26.5	4.5 X10-1 P 1.96 X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
1.5	E 1 B	METHANOL	26.5	4.0 X10-1 P 1.74 X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
2.	E-1 M	NA CL	27	3.4 X10-2 D 1.48 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HERR	62005	T	L
5.	E 0 B	PROPANOL-1	26.5	2.8 X10-1 P 1.22 X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
1.0	E 1 B	PROPANOL-1	26.5	1.7 X10-1 P 7.41 X10-3 S	BD	HEAT OF DILUTION	BENJ	66012	L	L
2.0	E 1 B	PROPANOL-1	26.5	1.1 X10-1 P 4.79 X10-3 S	BD	HEAT OF DILUTION	BENJ	66012	L	L
2.	E 0 M	UREA	26.5	5.4 X10-1 P 2.35 X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
4.	E 0 M	UREA	26.5	6.9 X10-1 P 3.00 X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
6.5	E 0 M	UREA	26.5	1.04 X10 0 P 4.533X10-2 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
1.	E 0 M	GUANIDINIUM CL	26.5	4.8 X10-1 P	BC	HEAT OF DILUTION	BENJ	66012	L	L
7.5	E 0	PH OF SOLUTION		2.09 X10-2 S					M	
2.	E 0 M	GUANIDINIUM CL	26.5	6.2 X10-1 P	BC	HEAT OF DILUTION	BENJ	66012	L	L
7.5	E 0	PH OF SOLUTION		2.70 X10-2 S					M	
3.	E 0 M	GUANIDINIUM CL	26.5	7.8 X10-1 P	BC	HEAT OF DILUTION	BENJ	66012	L	L
7.5	E 0	PH OF SOLUTION		3.40 X10-2 S					M	
4.	E 0 M	GUANIDINIUM CL	26.5	8.8 X10-1 P	BC	HEAT OF DILUTION	BENJ	66012	L	L
7.5	E 0	PH OF SOLUTION		3.83 X10-2 S					M	
2.	E 0 N	GUANIDINIUM CO3	26.5	1.9 X10-1 P 8.28 X10-3 S	BC	HEAT OF DILUTION	BENJ	66012	L	L
1.	E-1 M	NA CL	25	3.50 X10-3 M	BC	FOTOMTR SOLUBLZTN YLOB	TOKI OHKI	66027	G	L
2.0	E 0	PH OF SOLUTION								
1.	E-1 M	NA CL	25	3.10 X10-3 M	BC	FOTOMTR SOLUBLZTN YLOB	TOKI OHKI	66027	G	L
3.2	E 0	PH OF SOLUTION								
1.	E-1 M	NA CL	25	1.95 X10-3 M	BC	FOTOMTR SOLUBLZTN YLOB	TOKI OHKI	66027	G	L
4.2	E 0	PH OF SOLUTION								
1.	E-1 M	NA CL	25	1.25 X10-3 M	BC	FOTOMTR SOLUBLZTN YLOB	TOKI OHKI	66027	G	L
5.1	E 0	PH OF SOLUTION								
1.	E-1 M	NA CL	25	1.01 X10-3 M	BC	FOTOMTR SOLUBLZTN YLOB	TOKI OHKI	66027	G	L
6.1	E 0	PH OF SOLUTION								
1.	E-1 M	NA CL	25	1.00 X10-3 M	BC	FOTOMTR SOLUBLZTN YLOB	TOKI OHKI	66027	G	L
7.1	E 0	PH OF SOLUTION								
33 ENTRIES FOR COMPOUND										

COMPOUND NO = 22 MOL WGT -

265.9 DIMETHYL DODECYL AMINE OXIDE HYDROCHLORIDE

1.	E-2 M	H CL	25	5. X10-3 M	BE	VALUES FRM REF IN CMC	HERR	64006	R
1.	E-3 M	H CL	27	1.9 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HERR	64016	L
1.	E-2 M	H CL	27	7.14 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HERR	62005	T
1.	E-3 M	H CL	27	1.8 X10-1 D	BB	TURBIDITY PLT LITE SCATR	HERR	62005	M
2.	E-2 M	NA CL	27	6.76 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HERR	62005	T
1.	E-3 M	H CL	27	3.4 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HERR	62005	L
1.	E-1 M	NA CL	27	1.27 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HERR	62005	M
1.	E-1 M	NA CL	27	4.8 X10-2 D	BB	TURBIDITY PLT LITE SCATR	HERR	62005	T
1.	E-1 M	NA CL	27	1.80 X10-3 M	BB	TURBIDITY PLT LITE SCATR	HERR	62005	L
6 ENTRIES FOR COMPOUND									

COMPOUND NO = 23 MOL WGT -

373.2 SILVER DODECYL 1 SULFATE

1.	E-2 M	AG NO3	35	7.3 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	2
1.	E-2 M	NA NO3	35	4.7 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	3
4 ENTRIES FOR COMPOUND			35	5.4 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T	3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 24 MOL WGT -			570.8	CALCIUM DODECYL 1 SULFATE					
1.0 E-4 N CA CL2		54	1.3 X10-3 M	BC	SURFACE TENSION LOG PLOT	MIYA	60029	T L	
3.0 E-4 N CA CL2		54	1.3 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	MIYA	60029	T L	
1. E-3 N CA CL2		70	3.4 X10-3 N	BB	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T 3	
5. E-3 N CA CL2		70	2.9 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
2.5 E-2 N CA CL2		70	1.45 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
1. E-3 M NA CL		70	2.85 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
4. E-3 M NA CL		70	2.8 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
1. E-2 M NA CL		70	2.7 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
2. E-2 M NA CL		70	2.6 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
3. E-2 M NA CL		70	2.6 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
5. E-2 M NA CL		70	2.4 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
1.0 E-1 M NA CL		70	1.7 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
2.0 E-1 M NA CL		70	1.25 X10-3 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
5.0 E-1 M NA CL		70	7.3 X10-4 N	CG	VISUAL SPCTR CHNGE PNCN	LANG	51005	L L	
1. E-2 M NA NO3		70	3.3 X10-3 N	BB	SPECFC CONDCTNCE GRAPH	CORK GOOD	62006	T 3	
0001		70			SEE CMPD NMBR IN ADDITV	LANG	51005	X	
20 ENTRIES FOR COMPOUND									
COMPOUND NO = 25 MOL WGT -			288.3	SODIUM ETHYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	9.80 X10-2 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 26 MOL WGT -			330.4	SODIUM AMYL ALPHA SULFOPELARGONATE					
2 ENTRIES FOR COMPOUND		UNK	1.56 X10-2 M	CG	VISUAL SPCTR CHNGE PNCN	MAUR STIR	64002	T L	
		UNK	1.60 X10-2 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 27 MOL WGT -			344.4	SODIUM HEXYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	7.0 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 28 MOL WGT -			358.4	SODIUM HEPTYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	4.5 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 29 MOL WGT -			386.5	SODIUM NONYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	1.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 30 MOL WGT -			400.5	SODIUM DECYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	5. X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 31 MOL WGT -			372.4	SODIUM 2 OCTYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	3.6 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 32 MOL WGT -			372.4	SODIUM 2 ETHYLHEXYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	1.9 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 33 MOL WGT -			400.5	SODIUM /OXO/ DECYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	5. X10-4 M	HG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 34 MOL WGT -			428.6	SODIUM DODECYL ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	1. X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	
COMPOUND NO = 35 MOL WGT -			574.3	SODIUM H/CF2CF2/3CH2 ALPHA SULFOPELARGONATE					
1 ENTRIES FOR COMPOUND		UNK	2.4 X10-3 M	HG	VISUAL SPCTR CHNGE PNCN	STIR WEIL	62008	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(0 or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
		RM 1.4 X10-2 D 3.75 X10-4 M	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L
		RM 1.5 X10-2 D 4.02 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	56008	T L
4 ENTRIES FOR COMPOUND							M
COMPOUND NO = 37 MOL WGT -	193.7	DECYLAMMONIUM CHLORIDE					
	20	3.2 X10-2 M	CC	EQUIV CONDCTNCE GRAPH	RALS HOER	42002	P L
	25	4. X10-2 M	CE	REFRACTIVE INDEX	KLEV	48005	T L
	25	4.8 X10-2 M	XC	REFRACTIVE INDEX	KLEV	53010	T L
	25.0	5.40 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	HOYE GREE	57004	T L
	40	3.8 X10-2 M	CC	EQUIV CONDCTNCE GRAPH	RALS HOER	42002	P L
	60	3.8 X10-2 M	CC	EQUIV CONDCTNCE GRAPH	RALS HOER	42002	P L
	RM	4. X10-2 M	BD	PH OR HYDROLYSIS	VEIS HOER	60001	T L
7 ENTRIES FOR COMPOUND							
COMPOUND NO = 38 MOL WGT -	221.8	DODECYL AMMONIUM CHLORIDE					
	59016			VALUES FRM REF IN CMC	BOTR CRES	60024	R
	50008			VALUES FRM REF IN CMC	CORR HARK	46004	R
	47006			VALUES FRM REF IN CMC	CORR HARK	46005	R
	48016			VALUES FRM REF IN CMC	KOLT STRI	49005	R
	15	1.56 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T 3
	18.0	1.45 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	HOYE GREE	57004	T L
	20	1.5 X10-2 M	XG	VISCOSITY MINIMUM	SATA TYUZ	53006	T L
	20	1.56 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T L
	20	1.50 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	RALS EGGE	48027	P 2
	23.5	2.95 X10-1 D	BB	REFRACTIVE INDEX	KLEV	46012	T L
		1.330X10-2 M					M
	25	1.43 X10-2 M	BG	EQUIV COND 1ST DEVIATION	BROW GRIE	49014	T L
	25	1.46 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	P 1
	25	1.28 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
	25	1.52 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T L
	25	1.34 X10-2 W	BB	SPECFC CONDCTNCE GRAPH	CZER	65031	T L
	25.6	1.31 X10-2 M	CG	FOTOMTR SPCTR CHNGE INPX	KLEV	47004	T L
	25.0	1.34 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G L
	25	1.47 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	RALS EGGE	48027	P 1
	25.0	1.38 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	HOYE GREE	57004	T L
	26	1.27 X10-2 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47006	T L
	26	1.36 X10-2 M	BG	VISUAL SPCTR CHNGE EOSN	CORR HARK	47006	T L
	26	1.30 X10-2 M	BG	VISUAL SPCTR CHNGE FL	CORR HARK	47006	T L
	26	1.24 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	47006	T L
	26	1.31 X10-2 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L
	26	1.31 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	47010	T L
	27	1.50 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T 3
	30	3.23 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L
		1.456X10-2 M					M
	30	1.6 X10-2 M	XG	VISCOSITY MINIMUM	SATA TYUZ	53006	T L
	30	1.46 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS BROO	49013	T L
	30	1.65 X10-2 W	XB	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
	30	1.3 X10-2 M	CD	REFRACTIVE INDEX	KLEV	48005	T L
	30	1.4 X10-2 M	XG	VISUAL SPCTR CHNGE	KLEV	53010	T L
	30	1.47 X10-2 M	BB	DENSITY	CART ANAC	60005	K 1
	30	1.29 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	RALS HOER	46001	T L
	30	1.34 X10-2 M	CB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L
	30	1.47 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K 1
	30	1.45 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48021	K L
	30	1.44 X10-2 N	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T L
	30	1.50 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T 1
	30	1.48 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	RALS EGGE	48027	P 1
	30	1.34 X10-2 M	CB	FOTOMTR SOLUBLZTN AZBZ	KOLT STRI	49005	T L
	30	1.34 X10-2 M	CB	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L
	30	1.48 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	GK 1
	35	1.36 X10-2 W	BB	SPECFC CONDCTNCE GRAPH	CZER	65031	T L
	35.8	1.25 X10-2 M	CG	FOTOMTR SPCTR CHNGE INPX	KLEV	47004	T L
	35.0	1.33 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	HOYE GREE	57004	T L
	40	1.49 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T L
	40	1.50 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	RALS EGGE	48027	P 2
	45.0	1.21 X10-2 M	CG	FOTOMTR SPCTR CHNGE INPX	KLEV	47004	T L
	45	1.43 X10-2 W	BB	SPECFC CONDCTNCE GRAPH	CZER	65031	T L
	50	1.30 X10-2 M	HB	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L
	50	1.4 X10-2 W	XD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
	50	1.34 X10-2 M	CB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L
	50	1.65 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T L
	50	1.58 X10-2 M	BA	SPECFC CONDCTNCE GRAPH	RALS EGGE	48027	P 3

A-mol %; B-vol % solvent; C-mol % surfactant; D-wt/vol %; E-% saturation; F-wt % solvent; G-mol % surfactant; H-normality counterions;

M-molar; N-normal; P-wt %; Q-wt % surfactant; R-varied; S-mol/kg; T-wt % surfactant; U-mol/(l or kg); W-molal; Y-atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C.	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 36 MOL WGT -	372.4	SODIUM METHYL ALPHA SULFOPALMITATE						
	28	1.2 X10-2 D	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		3.22 X10-4 M					M	
UNK	4.	X10-4 M	CG	VISUAL SPCTR CHNGE PNCC	STIR WEIL	62008	T L	
50	1.37 X10-2 M	CB	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L		
50	1.34 X10-2 M	CB	FOTOMTR SOLUBLZTN AZBZ	KOLT STRI	49005	T L		
55.0	1.20 X10-2 M	CG	FOTOMTR SPCTR CHNGE INPX	KLEV	47004	T L		
55	1.50 X10-2 W	BB	SPECFC CONDCTNCE GRAPH	CZER	65031	T L		
60	1.3 X10-2 M	CC	EQUIV CONDCTNCE GRAPH	RALS HOER	42002	T L		
60	1.80 X10-2 M	BC	SPECFC CONDCTNCE GRAPH	EGGE HARW	51006	T L		
60	1.71 X10-2 M	BB	SPECFC CONDCTNCE GRAPH	RALS EGGE	48027	P 3		
UNK	1.4 X10-2 M	CC	UNSPECIFIED CONDUCTANCE	BOTR CRES	59016	T L		
UNK	1.4 X10-2 M	CD	ELECTROMOTIVE FORCE	BOTR CRES	59016	T L		
RM	1.3 X10-2 M	BD	PH OR HYDROLYSIS	VEIS HOER	60001	T L		
UNK	1.31 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L		
UNK	1.4 X10-2 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L		
3.66 E 0 A	ACETONITRILE	30	1.61 X10-2 N	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T 3
3.27 E 0 A	ACETONE	30	1.30 X10-2 N	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T 3
7.06 E 0 A	ACETONE	30	1.51 X10-2 N	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T L
9.95 E 0 A	ACETONE	30	4.54 X10-2 N	BD	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T L
7.67 E-3 M	BA CL2	26	1.05 X10-2 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L
1.27 E-2 M	BA CL2	26	8.66 X10-3 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T L
1.76 E-2 M	BA CL2	26	8.01 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	47010	T L
2.19 E-2 M	BA CL2	26	7.48 X10-3 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L
3.16 E-2 M	BA CL2	26	6.21 X10-3 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L
3.45 E-2 M	BA CL2	26	5.89 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	47010	T L
4.07 E-2 M	BA CL2	26	5.56 X10-3 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L
2.5 E-3 N	BA CL2	30	1.36 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
5. E-3 N	BA CL2	30	1.30 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
1. E-2 N	BA CL2	30	1.13 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
2. E-2 N	BA CL2	30	9.4 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
1.00 E 2 E	BENZENE	30	1.12 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48021	K L
1.55 E-1 M	BUTANOL-1	25	9.03 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.42 E-1 M	BUTANOL-1	25	5.96 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
4.64 E-1 M	BUTANOL-1	25	4.06 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
2.5 E-3 N	CA ACETATE	30	1.40 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
5. E-3 N	CA ACETATE	30	1.28 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
1. E-2 N	CA ACETATE	30	1.09 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
2. E-2 N	CA ACETATE	30	9.2 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
2. E-2 N	CA FORMATE	30	9.4 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
1.00 E 2 E	CYCLOHEXANE	30	1.25 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48021	K L
3.7 E 0 H	DIOXANE	UNK	1.36 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
7.5 E 0 H	DIOXANE	UNK	1.42 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
1.63 E 1 H	DIOXANE	UNK	1.51 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
9.02 E-5 M	DECANOL-1	25	1.09 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.43 E-4 M	DECANOL-1	25	9.43 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.88 E-4 M	DECANOL-1	25	8.27 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
2.00 E-4 M	DECANOL-1	25	7.91 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
2.31 E-4 M	DECANOL-1	25	7.61 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.9 E 0 H	ETHYLENE GLYCOL	UNK	1.33 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
7.9 E 0 H	ETHYLENE GLYCOL	UNK	1.36 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
1.54 E 1 H	ETHYLENE GLYCOL	UNK	1.40 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
5.69 E-1 M	ETHANOL	25	1.26 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
8.68 E-1 M	ETHANOL	25	1.20 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.918E 0 M	ETHANOL	25	1.06 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
2.627E 0 M	ETHANOL	25	9.73 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.163E 0 M	ETHANOL	25	8.75 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
2.65 E 0 A	ETHANOL	30	1.77 X10-2 N	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T 3
4.17 E 0 A	ETHANOL	30	1.38 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	RALS HOER	46001	T L
5.77 E 0 A	ETHANOL	30	3.00 X10-2 N	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T L
8.80 E 0 A	ETHANOL	30	1.52 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	RALS HOER	46001	T L
1.437E 1 A	ETHANOL	30	2.1 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	RALS HOER	46001	T L
2.068E 1 A	ETHANOL	30	4.3 X10-2 M	CD	SPECFC CONDCTNCE GRAPH	RALS HOER	46001	T L
2.812E 1 A	ETHANOL	30	1.03 X10-1 M	CD	SPECFC CONDCTNCE GRAPH	RALS HOER	46001	T L
8.6 E 0 H	GLYCEROL	UNK	1.33 X10-Q M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
1.30 E 1 H	GLYCEROL	UNK	1.35 X10-2 M	CG	VISUAL SPCTR CHNGE DCFL	CORR HARK	46004	G L
2. E-2 N	ACETIC ACID	30	1.49 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
2.5 E-3 M	HEXYL AMMONIUM CL	30	1.43 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K 3
5. E-3 M	HEXYL AMMONIUM CL	30	1.26 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K 3
1.0 E-2 M	HEXYL AMMONIUM CL	30	1.16 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K L
2.5 E-2 M	HEXYL AMMONIUM CL	30	8.8 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K L
5.0 E-2 M	HEXYL AMMONIUM CL	30	6.1 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K L
2.5 E-3 N	H CL	30	1.39 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
2.50 E-3 W	H CL	30.0	1.306X10-2 W	BB	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L
5. E-3 N	H CL	30	1.27 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
1. E-2 N	H CL	30	1.15 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.003E-2 W H CL	30.0	1.165X10-2 W	BB	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L	
2. E-2 N H CL	30	9.2 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
5.008E-2 W H CL	30.0	5.54 X10-3 W	BC	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L	
1.002E-1 W H CL	30.0	2.0 X10-3 W	BD	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L	
2.51 E-3 W H CL	40.0	1.162X10-2 W	BB	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L	
1.004E-2 W H CL	40.0	1.000X10-2 W	BB	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L	
5.012E-2 W H CL	40.0	5.04 X10-3 W	BE	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L	
1.002E-1 W H CL	40.0	1.9 X10-3 W	BD	ELECTROMOTIVE FORCE	HUTC WINS	57021	T L	
2. E-2 N FORMIC ACID	30	1.45 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
1.00 E 2 E N-HEXANE	30	1.30 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48021	K L	
2.5 E-3 N H NO3	30	1.20 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
5. E-3 N H NO3	30	1.02 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
1. E-2 N H NO3	30	7.3 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
2. E-2 N H NO3	30	4.9 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
5.0 E-3 M HEXANOL-1	25	1.17 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
6.6 E-3 M HEXANOL-1	25	1.13 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
8.2 E-3 M HEXANOL-1	25	1.06 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.18 E-2 M HEXANOL-1	25	1.01 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.46 E-2 M HEXANOL-1	25	9.5 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.64 E-2 M HEXANOL-1	25	9.4 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.68 E-2 M HEXANOL-1	25	9.1 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.96 E-2 M HEXANOL-1	25	8.4 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
2.5 E-3 N SUCCINIC ACID	30	1.48 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
2. E-2 N TARTARIC ACID	30	1.83 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
1.90 E-3 M HEPTANOL-1	25	1.15 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
2.74 E-3 M HEPTANOL-1	25	1.11 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
3.59 E-3 M HEPTANOL-1	25	1.09 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
4.94 E-3 M HEPTANOL-1	25	9.97 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
5.96 E-3 M HEPTANOL-1	25	9.02 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
6.43 E-3 M HEPTANOL-1	25	8.65 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
7.07 E-3 M HEPTANOL-1	25	8.55 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
7.73 E-3 M HEPTANOL-1	25	7.80 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
7.5 E-3 M K CL	25	1.01 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.50 E-2 M K CL	25	8.06 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
3.00 E-2 M K CL	25	6.62 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
6.00 E-2 M K CL	25	4.70 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
5. E-2 N K CL	50	6 X10-3 M	CE	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	49005	T L	
1. E-1 N K CL	50	3.5 X10-3 M	CB	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	49005	T L	
7.10 E-3 M LA CL3	26	1.15 X10-2 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L	
1.27 E-2 M LA CL3	26	8.12 X10-3 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L	
1.72 E-2 M LA CL3	26	6.95 X10-3 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L	
1.619E 1 H METHANOL	25	3.1 X10-2 M	BF	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T L	
1.619E 1 H METHANOL	25	1.77 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	P 3	
2.465E 1 H METHANOL	25	2.37 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3	
2.465E 1 H METHANOL	25	5.2 X10-2 M	BF	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T L	
4.82 E 0 A METHANOL	30	1.64 X10-2 N	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T 3	
1.271E 1 A METHANOL	30	2.50 X10-2 N	BD	EQUIV CONDCTNCE GRAPH	RALS EGGE	48023	T L	
2.5 E 1 B METHANOL	30	2.50 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
2.5 E-3 N NA ACETATE	30	1.39 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
5. E-3 N NA ACETATE	30	1.29 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
1. E-2 N NA ACETATE	30	1.10 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
2. E-2 N NA ACETATE	30	8.6 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
8.02 E-3 M NA CL	26	1.16 X10-2 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L	
1.71 E-2 M NA CL	26	9.88 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	47010	T L	
2.75 E-2 M NA CL	26	7.93 X10-3 M	BG	VISUAL SPCTR CHNGE SKYB	CORR HARK	47010	T L	
4.96 E-2 M NA CL	26	7.16 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	47010	T L	
9.23 E-2 M NA CL	26	5.13 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	47010	T L	
1. E-3 N NA CL	30	1.42 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
2.5 E-3 N NA CL	30	1.39 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
5. E-3 N NA CL	30	1.30 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
1. E-2 M NA CL	30	2.72 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L	
		1.226X10-2 M				M		
1. E-2 N NA CL	30	1.13 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
2. E-2 N NA CL	30	8.9 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3	
2. E-2 M NA CL	30	2.27 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L	
		1.023X10-2 M				M		
4. E-2 M NA CL	30	1.65 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L	
		7.439X10-3 M				M		
5. E-2 M NA CL	30	6.7 X10-3 M	BB	DENSITY	CART ANAC	60005	K 3	
5. E-2 N NA CL	30	6.0 X10-3 M	CB	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L	
5. E-2 N NA CL	30	6.0 X10-3 M	CB	FOTOMTR SOLUBLZTN AZBZ	KOLT STRI	49005	T L	
5. E-2 N NA CL	30	6.2 X10-3 M	CB	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	49005	T L	
6. E-2 M NA CL	30	1.45 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L	
		6.537X10-3 M				M		
8. E-2 M NA CL	30	1.18 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L	
		5.320X10-3 M				M		

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality

counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation	
1.0	E-1	M	NA CL	30	1.09 X10-1 D 4.914X10-3 M	CD	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T M
1.5	E-1	M	NA CL	30	1.01 X10-1 D 4.553X10-3 M	CD	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T M
2.0	E-1	M	NA CL	30	9.3 X10-2 D 4.19 X10-3 M	CE	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T M
2.5	E-1	M	NA CL	30	9.3 X10-2 D 4.19 X10-3 M	CE	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T M
3.0	E-1	M	NA CL	30	8.5 X10-2 D 3.83 X10-3 M	CE	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T M
5.	E-2	N	NA CL	50	6.5 X10-3 M	CB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
5.	E-2	N	NA CL	50	6.0 X10-3 M	CB	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L
1.	E-1	N	NA CL	50	3.6 X10-3 M	CB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
1.	E-1	N	NA CL	50	3.5 X10-3 M	CB	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L
2.	E-1	N	NA CL	50	2.6 X10-3 M	CC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
2.	E-2	N	NA HCO ₂ FORMATE	30	9.1 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
1.	E-3	N	NA N03	30	1.37 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
2.5	E-3	N	NA N03	30	1.23 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
5.	E-3	N	NA N03	30	9.9 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
1.	E-2	N	NA N03	30	7.6 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
2.	E-2	N	NA N03	30	4.6 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	G 3
2.5	E-3	N	NA SUCCINATE	30	6.1 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L
3.85	E-1	M	PROPANOL-1	25	1.09 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.31	E-1	M	PROPANOL-1	25	8.99 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
9.88	E-1	M	PROPANOL-1	25	7.03 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.54	E-1	M	PROPANOL-2	25	9.53 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.046E 0	M	PROPANOL-2	25	7.62 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
8.00	E 0	H	PROPANOL-2	25	9.2 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3
1.619E 1	H	PROPANOL-2	25	1.04 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3	
5.	E 2	Y	PRESSURE	25.0	1.39 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G L
1.	E 3	Y	PRESSURE	25.0	1.41 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G L
2.	E 3	Y	PRESSURE	25.0	1.42 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G L
4.84	E-5	M	UNDECANOL-1	25	1.06 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.39	E-5	M	UNDECANOL-1	25	9.43 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
8.31	E-5	M	UNDECANOL-1	25	9.19 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.01	E-4	M	UNDECANOL-1	25	7.47 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.28	E-4	M	UNDECANOL-1	25	7.07 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
5.66	E 0	H	TERTIARY BUTANOL	25	7.9 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3
7.88	E 0	H	TERTIARY BUTANOL	25	7.4 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T 3
6.39	E-3	M	TRIETHYL CARBINOL	25	1.18 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.22	E-2	M	TRIETHYL CARBINOL	25	1.11 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
2.22	E-2	M	TRIETHYL CARBINOL	25	1.01 X10-2 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.08	E-2	M	TRIETHYL CARBINOL	25	9.32 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.72	E-2	M	TRIETHYL CARBINOL	25	8.44 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
4.27	E-2	M	TRIETHYL CARBINOL	25	7.76 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
4.80	E-2	M	TRIETHYL CARBINOL	25	7.27 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
2.5	E-3	M	0039	30	2.5 X10-3 M	BE	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K L
1.	E-1	K	CL- ION	UNK	3.50 X10-3 M	CB	SURFACE TENSION LOG PLOT	ROE BRAS	54013	T L
2.0	E 0		PH OF SOLUTION							
2.52	E-5	M	DECANOL-1	25	4.43 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.00	E-2	M	K CL							
4.53	E-5	M	DECANOL-1	25	3.98 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.00	E-2	M	K CL							
4.96	E-5	M	DECANOL-1	25	6.00 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.00	E-2	M	K CL							
5.52	E-5	M	DECANOL-1	25	6.73 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.50	E-2	M	K CL							
6.32	E-5	M	DECANOL-1	25	3.70 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.00	E-2	M	K CL							
6.81	E-5	M	DECANOL-1	25	8.63 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
7.5	E-3	M	K CL							
8.05	E-5	M	DECANOL-1	25	3.53 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.00	E-2	M	K CL							
8.23	E-5	M	DECANOL-1	25	4.98 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.00	E-2	M	K CL							
8.86	E-5	M	DECANOL-1	25	3.11 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.00	E-2	M	K CL							
9.42	E-5	M	DECANOL-1	25	5.74 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
1.50	E-2	M	K CL							
1.01	E-4	M	DECANOL-1	25	2.95 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
6.00	E-2	M	K CL							
1.03	E-4	M	DECANOL-1	25	4.14 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.00	E-2	M	K CL							
1.10	E-4	M	DECANOL-1	25	6.95 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
7.5	E-3	M	K CL							
1.26	E-4	M	DECANOL-1	25	3.82 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L
3.00	E-2	M	K CL							

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.44 E-4 M DECANOL-1 3.00 E-2 M K CL	25	3.48 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.51 E-4 M DECANOL-1 1.50 E-2 M K CL	25	4.61 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.56 E-4 M DECANOL-1 7.5 E-3 M K CL	25	6.57 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.63 E-4 M DECANOL-1 3.00 E-2 M K CL	25	3.21 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.71 E-4 M DECANOL-1 1.50 E-2 M K CL	25	4.17 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.86 E-4 M DECANOL-1 1.50 E-2 M K CL	25	3.79 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.97 E-4 M DECANOL-1 7.5 E-3 M K CL	25	6.23 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
2.18 E-4 M DECANOL-1 7.5 E-3 M K CL	25	5.53 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
2.44 E-4 M DECANOL-1 7.5 E-3 M K CL	25	5.15 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
3.46 E-3 M HEXANOL-1 1.50 E-2 M K CL	25	7.79 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
4.03 E-3 M HEXANOL-1 7.5 E-3 M K CL	25	9.41 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
4.78 E-3 M HEXANOL-1 6.00 E-2 M K CL	25	3.86 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
4.88 E-3 M HEXANOL-1 3.00 E-2 M K CL	25	5.70 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
7.54 E-3 M HEXANOL-1 6.00 E-2 M K CL	25	3.05 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
7.88 E-3 M HEXANOL-1 3.00 E-2 M K CL	25	4.61 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
8.20 E-3 M HEXANOL-1 7.5 E-3 M K CL	25	8.70 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
9.79 E-3 M HEXANOL-1 1.50 E-2 M K CL	25	6.02 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
9.96 E-3 M HEXANOL-1 6.00 E-2 M K CL	25	2.68 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.10 E-2 M HEXANOL-1 3.00 E-2 M K CL	25	4.30 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CCRR HARK	50008	T L	
1.17 E-2 M HEXANOL-1 6.00 E-2 M K CL	25	2.36 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.20 E-2 M HEXANOL-1 7.5 E-3 M K CL	25	7.61 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.29 E-2 M HEXANOL-1 6.00 E-2 M K CL	25	2.09 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.36 E-2 M HEXANOL-1 3.00 E-2 M K CL	25	3.97 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.37 E-2 M HEXANOL-1 1.50 E-2 M K CL	25	5.62 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.41 E-2 M HEXANOL-1 6.00 E-2 M K CL	25	1.90 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.44 E-2 M HEXANOL-1 3.00 E-2 M K CL	25	3.38 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.60 E-2 M HEXANOL-1 7.5 E-3 M K CL	25	6.79 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.65 E-2 M HEXANOL-1 1.50 E-2 M K CL	25	5.08 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.69 E-2 M HEXANOL-1 3.00 E-2 M K CL	25	3.29 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.91 E-2 M HEXANOL-1 1.50 E-2 M K CL	25	4.71 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
1.93 E-2 M HEXANOL-1 7.5 E-3 M K CL	25	6.13 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
2.23 E-2 M HEXANOL-1 7.5 E-3 M K CL	25	5.69 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
2.54 E-2 M HEXANOL-1 7.5 E-3 M K CL	25	5.39 X10-3 M	BG	VISUAL SPCTR CHNGE INPX	CORR HARK	50008	T L	
2.564E 1 H METHANOL 4.4 E 0 I LAURYL ALCOHOL	25	1.69 X10-2 M	BG	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T L	
2.564E 1 H METHANOL 1.11 E 1 I LAURYL ALCOHOL	25	1.08 X10-2 M	BG	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T L	
2.564E 1 H METHANOL 2.61 E 1 I LAURYL ALCOHOL	25	8.3 X10-3 M	BG	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T L	
2.564E 1 H METHANOL 6.51 E 1 I LAURYL ALCOHOL	25	6.9 X10-3 M	BG	EQUIV CONDCTNCE GRAPH	BROW GRIE	49014	T L	
2.5 E 1 B METHANOL 2. E-2 N NA CL	30	1.82 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	49008	K L	
293 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 39 MOL WGT -	249.9	TETRADECYL AMMONIUM CHLORIDE						
25.0 2.8 X10-3 M	CC	SPECFC CONDCTNCE GRAPH	HOYE GREE	57004	T L			
30 3.7 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	RALS EGGE	48020	K L			
40 3.1 X10-3 M	CC	REFRACTIVE INDEX	KLEV	48005	T L			
40 3.1 X10-3 M	XC	REFRACTIVE INDEX	KLEV	53010	T L			
60 4.5 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	RALS HOER	42002	P L			
0038		SEE CMPD NMBR IN ADDITV	RALS EGGE	48020	X			
6 ENTRIES FOR COMPOUND								
COMPOUND NO = 40 MOL WGT -	288.4	POTASSIUM DODECYL 1 SULFONATE						
25 9.00 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
25 9.0 X10-3 M	CC	REFRACTIVE INDEX	LIN	57005	T L			
30 9.10 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
35 9.20 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
40 9.30 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
40 1.1 X10-2 M	BC	SURFACE TENSION LOG PLOT	V VO	61026	TL L			
45 9.50 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
50 9.75 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
55 1.000X10-2 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
60 1.040X10-2 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L			
5. E-2 D BENZENE	25	8.65 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.0 E-1 D BENZENE	25	8.25 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.5 E-1 D BENZENE	25	7.60 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
5. E-2 D BENZENE	30	8.80 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.0 E-1 D BENZENE	30	8.45 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.5 E-1 D BENZENE	30	7.95 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
5. E-2 D BENZENE	35	9.00 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.0 E-1 D BENZENE	35	8.75 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.5 E-1 D BENZENE	35	8.35 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
5. E-2 D BENZENE	40	9.10 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.0 E-1 D BENZENE	40	8.90 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
1.5 E-1 D BENZENE	40	8.65 X10-3 M	CA	SPECFC CONDCTNCE GRAPH	LIN	55008	T L	
2. E-2 K K NO3	20	4.47 X10-3 M	BB	SURFACE TENSION LOG PLOT	V VO	61026	TL L	
4. E-2 K K NO3	20	3.55 X10-3 M	BB	SURFACE TENSION LOG PLOT	V VO	61026	TL L	
24 ENTRIES FOR COMPOUND								
COMPOUND NO = 41 MOL WGT -	263.9	DODECYL TRIMETHYL AMMONIUM CHLORIDE						
23 5.70 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L			
2.159X10-2 M						M		
25 1.72 X10-2 M	BB	UNSPECIFIED CONDUCTANCE	HOYE MARM	61002	T L			
25 2.0 X10-2 M	BC	SURFACE TENSION LOG PLOT	WEIN ZOGR	65026	T L			
25.0 2.03 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3			
30 2.28 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS BROO	49013	T L			
30 1.6 X10-2 M	XG	VISUAL SPCTR CHNGE	KLEV	53010	T L			
UNK 2.28 X10-2 M	BC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T L			
25 7.5 X10-3 M	BC	SURFACE TENSION LOG PLOT	WEIN ZOGR	65026	T L			
23 4.20 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L			
1.591X10-2 M						M		
4. E-2 M NA CL	23	3.10 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L	
1.174X10-2 M						M		
1.0 E-1 M NA CL	23	1.90 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L	
7.199X10-3 M						M		
5. E 2 Y PRESSURE	25.0	2.09 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
1. E 3 Y PRESSURE	25.0	2.11 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
1.5 E 3 Y PRESSURE	25.0	2.04 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
2. E 3 Y PRESSURE	25.0	1.98 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
3. E 3 Y PRESSURE	25.0	1.87 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
4. E 3 Y PRESSURE	25.0	1.83 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
5. E 3 Y PRESSURE	25.0	1.81 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3	
2.59 E 1 C 0042	25	1.14 X10-2 M	BB	UNSPEC SOLUBLZTN SDN 4	HOYE MARM	61002	T L	
5.00 E 1 C 0042	25	7.47 X10-3 M	BB	UNSPEC SOLUBLZTN SDN 4	HOYE MARM	61002	T L	
7.59 E 1 C 0042	25	5.01 X10-3 M	BB	UNSPEC SOLUBLZTN SDN 4	HOYE MARM	61002	T L	
4.96 E-1 W NA CL	31.5	3.8 X10-3 M	BC	DEBYE PLT LIGHT SCATTER	ANAC GHOS	63016	T L	
3.8 E-3 M NA BR								
22 ENTRIES FOR COMPOUND								
COMPOUND NO = 42 MOL WGT -	292.0	TETRADECYL TRIMETHYL AMMONIUM CHLORIDE						
23 1.20 X10-1 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L			
4.109X10-3 M						M		
25 3.0 X10-3 M	HE	EQUIV CONDCTNCE GRAPH	BRAD MCBA	48009	T L			
25 4.47 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	HOYE MARM	61002	T 3			
40 3.6 X10-3 M	XG	VISUAL SPCTR CHNGE	KLEV	53010	T L			

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality

counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation		
2.	E-2 M	NA CL	40 UNK 23 2.39 X10-3 M	4.0 X10-3 M 5.3 X10-3 M 7.0 X10-2 D 1.36 X10-3 M	XC CC CC	REFRACTIVE INDEX UNSPECIFIED CONDUCTANCE TURBIDITY PLT LITE SCATR	KLEV CELL EGGE KUSH PARK	53010 52001 57006	T L T L T L		
4.	E-2 M	NA CL	23	4.0 X10-2 D	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L		
1.0	E-1 M	NA CL	23	1.36 X10-3 M 3.0 X10-2 D 1.02 X10-3 M	CC	TURBIDITY PLT LITE SCATR	KUSH PARK	57006	T L		
		0041				SEE CMPD NMBR IN ADDITV	HOYE MARM	61002	X		
10 ENTRIES FOR COMPOUND											
COMPOUND NO = 43 MOL WGT - 88.1 BUTYRIC ACID											
		0	1.50 X10 1 P 1.702X10 0 S	AB	PARTIAL VOLUME	GRIN BURY	29001	T L			
		0	1.1 X10 0 W	AE	FREEZING POINT	JONE BURY	27002	T L			
		0	1.5 X10 0 W	AE	FREEZING POINT	JONE BURY	27002	P L			
		0	1.82 X10 0 M	AD	UNSPEC SOLUBLZTN PDMAB	MOUL BENS	59007	T L			
		12	1.37 X10 1 P 1.555X10 0 S	AB	PARTIAL VOLUME	GRIN BURY	29001	T L			
		18	1.30 X10 1 P 1.475X10 0 S	AB	PARTIAL VOLUME	GRIN BURY	29001	T L			
		25	1.23 X10 1 P 1.396X10 0 S	AB	PARTIAL VOLUME	GRIN BURY	29001	T L			
		25	1.75 X10 0 M	AD	UNSPEC SOLUBLZTN PDMAB	MOUL BENS	59007	T L			
		35	1.20 X10 1 P 1.362X10 0 S	AB	PARTIAL VOLUME	GRIN BURY	29001	T L			
		40	1.58 X10 0 M	AD	UNSPEC SOLUBLZTN PDMAB	MOUL BENS	59007	T L			
		60	1.62 X10 0 M	AD	UNSPEC SOLUBLZTN PDMAB	MOUL BENS	59007	T L			
		UNK	1.2 X10 0 M	XG	METHOD NOT CITED	KLEV RAIS	54004	T L			
12 ENTRIES FOR COMPOUND											
COMPOUND NO = 44 MOL WGT - 182.3 POTASSIUM OCTANOATE											
		25	4.7 X10-1 M	BC	GRAPH DATA NOT RETRIEVED	KLEV	46007	R			
		25	3.9 X10-1 M	DC	PARTIAL VOLUME	DAVI BURY	30001	T L			
		25	3.95 X10-1 M	CG	REFRACTIVE INDEX	KLEV	48005	T L			
		25	4.00 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	T L			
		25	3.9 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		45	4.5 X10-1 M	DC	UNSPEC SPCTR CHNGE PNCN	KLEV	58011	T L			
		UNK	3.9 X10-1 M	XG	REFRACTIVE INDEX	KLEV	48005	T L			
		25	3.84 X10-1 M	CG	METHOD NOT CITED	KLEV RAIS	54004	T L			
		25	3.76 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	T L			
		25	3.09 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	2.65 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	2.16 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	1.75 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	1.52 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	1.11 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	9.1 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	8.5 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		25	8.0 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L			
		2.	E 0 I	K OH	VISUAL SPCTR CHNGE PNCN	SHIN	55004	S L			
		3.3	E-2 W	K OH	4.00 X10-1 M	BB	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
		3.3	E-2 W	K OH	3.72 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3	
		4.2	E-2 W	K OH	3.55 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59008	T L	
		3.3	E-2 W	K OH	4.3 X10-1 W	BB	VAPR PRESSURE LOWERING	WHIT BENS	59008	T L	
		25	3.45 X10-1 W	BA	VAPR PRESSURE LOWERING	WHIT BENS	59012	E 3			
		25	4.25 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	60002	T L			
		25	3.8 X10-1 M	BG	SPECFC CONDCTNCE EQUATNS	SHIN	54005	G L			
		30	3.30 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3			
		30	4.2 X10-1 W	BB	VAPR PRESSURE LOWERING	WHIT BENS	59008	T L			
		35	3.13 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3			
		35	3.92 X10-1 W	BA	SPECIFIC HEAT	WHIT BENS	60002	T L			
		40	3.05 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3			
		45	3.10 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3			
		45	3.93 X10-1 W	BA	SPECIFIC HEAT	WHIT BENS	60002	T L			
		50	3.18 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3			
		55	3.31 X10-1 W	BA	SPECFC CONDCTNCE EQUATNS	WHIT BENS	59012	E 3			
		55	4.42 X10-1 W	BA	SPECIFIC HEAT	WHIT BENS	60002	T L			
		25	3.40 X10-1 M	DG	SPECFC CONDCTNCE EQUATNS	HERZ	52015	T L			
		25	3.88 X10-1 M	DG	SPECFC CONDCTNCE EQUATNS	HERZ	52015	T L			
		25	4.01 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52015	T L			
		25	4.02 X10-1 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52015	T L			
		25			SEE CMPD NMBR IN ADDITV	SHIN	54005	X			

Concentration units: A - mol %; B - vol % solvent; C - mol % surfactant mixture; D - wt/vol %; E - % saturation; H - wt % solvent; I - mol % surfactant; K - normality;

counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molar; Y – atm. Details on page 222

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
0.64 E-2 M BUTANOL-1 2. E O I K OH	10	3.80 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.20 E-1 M BUTANOL-1 2. E O I K OH	10	3.55 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.17 E-1 M BUTANOL-1 2. E O I K OH	10	3.22 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.21 E-1 M BUTANOL-1 2. E O I K OH	10	2.73 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.64 E-1 M BUTANOL-1 2. E O I K OH	10	2.29 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.10 E-4 M DECANOL-1 2. E O I K OH	10	3.91 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.13 E-4 M DECANOL-1 2. E O I K OH	10	3.76 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
6.4 E-1 M ETHANOL 2. E O I K OH	10	3.59 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.16 E 0 M ETHANOL 2. E O I K OH	10	3.32 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.82 E 0 M ETHANOL 2. E O I K OH	10	2.93 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.50 E 0 M ETHANOL 2. E O I K OH	10	2.57 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.11 E 0 M ETHANOL 2. E O I K OH	10	2.20 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
7.0 E-3 M HEXANOL-1 2. E O I K OH	10	3.79 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.32 E-2 M HEXANOL-1 2. E O I K OH	10	3.55 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.03 E-2 M HEXANOL-1 2. E O I K OH	10	3.27 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.94 E-2 M HEXANOL-1 2. E O I K OH	10	2.97 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.90 E-2 M HEXANOL-1 2. E O I K OH	10	2.63 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.08 E-3 M HEPTANOL-1 2. E O I K OH	10	3.83 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.95 E-3 M HEPTANOL-1 2. E O I K OH	10	3.65 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
7.42 E-3 M HEPTANOL-1 2. E O I K OH	10	3.41 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.43 E-2 M HEPTANOL-1 2. E O I K OH	10	3.28 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
7.1 E-4 M OCTANOL-1 2. E O I K OH	10	3.84 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.36 E-3 M OCTANOL-1 2. E O I K OH	10	3.71 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.73 E-3 M OCTANOL-1 2. E O I K OH	10	3.54 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.54 E-3 M OCTANOL-1 2. E O I K OH	10	3.47 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.28 E-1 M PROPANOL-1 2. E O I K OH	10	3.73 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.43 E-1 M PROPANOL-1 2. E O I K OH	10	3.53 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.37 E-1 M PROPANOL-1 2. E O I K OH	10	3.18 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
7.05 E-1 M PROPANOL-1 2. E O I K OH	10	2.92 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
9.42 E-1 M PROPANOL-1 2. E O I K OH	10	2.48 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.27 E 0 M PROPANOL-1 2. E O I K OH	10	2.21 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.63 E-4 M NONANOL-1 2. E O I K OH	10	3.88 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.18 E-4 M NONANOL-1 2. E O I K OH	10	3.80 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.65 E-4 M NONANOL-1 2. E O I K OH	10	3.69 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
6.36 E-4 M NONANOL-1 2. E O I K OH	10	3.63 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.1 E 0 C 0090 2. E O I K OH	25	3.4 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
5.6 E 0 C 0090 2. E O I K OH	25	3.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
9.4 E 0 C 0090 2. E O I K OH	25	2.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
1.36 E 1 C 0090 2. E O I K OH	25	2.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.19 E 1 C 0090 2. E O I K OH	25	2.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.51 E 1 C 0090 2. E O I K OH	25	2.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.32 E 1 C 0090 2. E O I K OH	25	1.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
4.30 E 1 C 0090 2. E O I K OH	25	1.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
5.91 E 1 C 0090 2. E O I K OH	25	1.3 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
7.58 E 1 C 0090 2. E O I K OH	25	1.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.0 E 0 C 0091 2. E O I K OH	25	2.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.3 E 0 C 0091 2. E O I K OH	25	2.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
4.4 E 0 C 0091 2. E O I K OH	25	1.4 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
7.7 E 0 C 0091 2. E O I K OH	25	1.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.21 E 1 C 0091 2. E O I K OH	25	8.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.00 E 1 C 0091 2. E O I K OH	25	6.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.74 E 1 C 0091 2. E O I K OH	25	4.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
5.62 E 1 C 0091 2. E O I K OH	25	3.3 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
E 0 0092 2. E O I K OH	25		GRAPH DATA NOT RETRIEVED		SHIN	54005	R	
1.2 E 0 C 0092 2. E O I K OH	25	7.3 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.9 E 0 C 0092 2. E O I K OH	25	5.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.6 E 0 C 0092 2. E O I K OH	25	4.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
4.6 E 0 C 0092 2. E O I K OH	25	3.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.67 E 1 C 0092 2. E O I K OH	25	2.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.61 E 1 C 0092 2. E O I K OH	25	1.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
6.18 E 1 C 0092 2. E O I K OH	25	1.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
0296 2. E O I K OH	25		SEE CMPD NMBR IN ADDITV		SHIN	54005	X	
2.3 E 0 C 0297 2. E O I K OH	25	3.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.8 E 0 C 0297 2. E O I K OH	25	2.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
6.1 E 0 C 0297 2. E O I K OH	25	2.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.09 E 1 C 0297 2. E O I K OH	25	1.7 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.73 E 1 C 0297 2. E O I K OH	25	1.3 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.94 E 1 C 0297 2. E O I K OH	25	9.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
5.01 E 1 C 0297 2. E O I K OH	25	6.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
6.87 E 1 C 0297 2. E O I K OH	25	5.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.68 E-2 M 3-METHYL BUTANOL-1 2. E O I K OH	10	3.80 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.47 E-2 M 3-METHYL BUTANOL-1 2. E O I K OH	10	3.53 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
9.04 E-2 M 3-METHYL BUTANOL-1 2. E O I K OH	10	3.20 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.42 E-1 M 3-METHYL BUTANOL-1 2. E O I K OH	10	2.88 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.92 E-1 M 3-METHYL BUTANOL-1 2. E O I K OH	10	2.27 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.68 E-1 M 3-METHYL BUTANOL-1 2. E O I K OH	10	2.00 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
 D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality;
 counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg;
 T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
9.7 E O C 0090 5.61 E 1 C 0091 2. E O I K OH	25	3.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.40 E 1 C 0090 3.61 E 1 C 0091 2. E O I K OH	25	4.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.65 E 1 C 0090 2.49 E 1 C 0091 2. E O I K OH	25	5.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.81 E 1 C 0090 1.75 E 1 C 0091 2. E O I K OH	25	6.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.92 E 1 C 0090 1.24 E 1 C 0091 2. E O I K OH	25	8.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.01 E 1 C 0090 8.6 E O C 0091 2. E O I K OH	25	9.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.07 E 1 C 0090 5.7 E O C 0091 2. E O I K OH	25	1.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.12 E 1 C 0090 3.3 E O C 0091 2. E O I K OH	25	1.4 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.16 E 1 C 0090 1.5 E O C 0091 2. E O I K OH 0188	25	1.7 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
127 ENTRIES FOR COMPOUND				SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
COMPOUND NO = 45 MOL WGT -	334.4	SODIUM P 1	METHYL DECYL BENZENE SULFONATE					
	19.0	2.45 X10-3 W	BC	KRAFFT POINT SOLUBILITY	GERS	57012	T L	
	35	2.53 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 2	
	40	1.90 X10-3 W	BC	SURFACE TENSION LOG PLOT	GERS	57012	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 46 MOL WGT -	362.4	SODIUM P 1	METHYL DODECYL BENZENE SULFONATE					
	27.7	7.1 X10-4 W	CC	KRAFFT POINT SOLUBILITY	GERS	57012	T L	
	35	7.2 X10-4 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
	40	6.2 X10-4 W	CD	SURFACE TENSION MINIMUM	GERS	57012	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 47 MOL WGT -	390.5	SODIUM P 1	METHYL TETRADECYL BENZENE SULFONATE					
	32.6	5.0 X10-4 W	BC	KRAFFT POINT SOLUBILITY	GERS	57012	T L	
	40	2.2 X10-4 W	BC	SURFACE TENSION LOG PLOT	GERS	57012	T L	
	40	3.1 X10-4 W	BB	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 48 MOL WGT -	418.6	SODIUM P 1	METHYL HEXADECYL BENZENE SULFONATE					
	45.5	1.4 X10-4 W	CD	KRAFFT POINT SOLUBILITY	GERS	57012	T L	
	50	1.3 X10-4 W	CC	SPECFC CONDCTNCE GRAPH	GERS	57012	T L	
	50	1.4 X10-4 W	CD	SURFACE TENSION MINIMUM	GERS	57012	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 49 MOL WGT -	292.3	SODIUM P OCTYL BENZENE SULFONATE						
	25	1.11 X10-2 W	CD	SURFACE TENSION MINIMUM	GERS	57012	T L	
	25	1.1 X10-2 M	CE	EQUIV COND 1ST DEVIATION	LING TART	43001	T L	
	25	1.23 X10-2 W	CB	EQUIV CONDCTNCE GRAPH	LING TART	43001	P L	
	35	1.47 X10-2 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
	40	1.2 X10-2 M	CE	EQUIV COND 1ST DEVIATION	LING TART	43001	T L	
	40	1.32 X10-2 W	CB	EQUIV CONDCTNCE GRAPH	LING TART	43001	P L	
	60	1.63 X10-2 W	CB	EQUIV CONDCTNCE GRAPH	LING TART	43001	P L	
	60	1.5 X10-2 M	CE	EQUIV COND 1ST DEVIATION	LING TART	43001	T L	
9 ENTRIES FOR COMPOUND	UNK	1.12 X10-2 M	CG	METHOD NOT CITED	KLEV CARR	56001	T L	
COMPOUND NO = 50 MOL WGT -	320.4	SODIUM P DECYL BENZENE SULFONATE						
	50	1.47 X10-2 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3	
	50	3.14 X10-3 W	CD	SURFACE TENSION MINIMUM	GERS	57012	T L	
	UNK	3.5 X10-3 M	CG	METHOD NOT CITED	KLEV CARR	56001	T L	
3 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 51	MOL WGT –	348.4	SODIUM P DODECYL BENZENE SULFONATE					
		23	1.6 X10-3 M	HG	STREAMING CURRENT	CARD	66011	T L
		60	1.20 X10-3 W	CA	SPECFC CONDCTNCE GRAPH	GERS	57012	T 3
		60	1.26 X10-3 W	CB	EQUIV CONDCTNCE GRAPH	LING TART	43001	P L
		60	1.2 X10-3 M	CE	EQUIV COND 1ST DEVIATION	LING TART	43001	T L
		UNK	2.8 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	FAVA EYRI	56016	T L
		UNK	2.8 X10-3 M	CG	VISUAL SPCTR CHNGE RHD6	FAVA EYRI	56016	T L
		UNK	1.0 X10-3 M	CB	SURFACE TENSION LOG PLOT	FAVA EYRI	56016	T L
7 ENTRIES FOR COMPOUND								
COMPOUND NO = 52	MOL WGT –	388.5	SODIUM HEXADECYL MONO OXYETHYLENE SULFATE					
		25	2.2 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		25	2.1 X10-4 M	CC	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		25	2.34 X10-4 M	CC	EQUIV CONDCTNCE GRAPH	WEIL BIST	58003	T L
		50	2.4 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 53	MOL WGT –	432.6	SODIUM HEXADECYL DI OXYETHYLENE SULFATE					
		25	1.4 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		25	1.2 X10-4 M	CC	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		25	1.34 X10-4 M	CC	EQUIV CONDCTNCE GRAPH	WEIL BIST	58003	T L
		50	1.4 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 54	MOL WGT –	476.6	SODIUM HEXADECYL TRI OXYETHYLENE SULFATE					
		25	1.0 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		25	7.0 X10-5 M	CD	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		25	1.23 X10-4 M	CC	EQUIV COND MAX BEGINING	WEIL BIST	58003	T L
		50	1.95 X10-4 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L
		50	1.2 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
5 ENTRIES FOR COMPOUND								
COMPOUND NO = 55	MOL WGT –	520.7	SODIUM HEXADECYL TETRA OXYETHYLENE SULFATE					
		25	8.0 X10-5 M	CD	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		25	1.0 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		50	1.01 X10-4 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L
		50	1.2 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 56	MOL WGT –	416.6	SODIUM OCTADECYL MONO OXYETHYLENE SULFATE					
		25	1.1 X10-4 M	BG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		25	1.9 X10-4 M	BD	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		50	9.0 X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 57	MOL WGT –	460.6	SODIUM OCTADECYL DI OXYETHYLENE SULFATE					
		25	7.0 X10-5 M	BG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		25	8.0 X10-5 M	BD	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		50	1.00 X10-4 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L
		50	7.0 X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 58	MOL WGT –	504.7	SODIUM OCTADECYL TRI OXYETHYLENE SULFATE					
		25	5.0 X10-5 M	CD	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		25	5.0 X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		50	6.98 X10-5 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L
		50	7.0 X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 59	MOL WGT –	548.7	SODIUM OCTADECYL TETRA OXYETHYLENE SULFATE					
		25	4.0 X10-5 M	CD	SURFACE TENSION LOG PLOT	WEIL BIST	58003	T L
		25	4.0 X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL BIST	58003	T L
		50	3.97 X10-5 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L
		50	7.0 X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 60	MOL WGT –	471.7	TRIETHANOL AMMONIUM HEXADECYL SULFATE					
		50	3.4 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L
1 ENTRIES FOR COMPOUND								

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – mola; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 61 MOL WGT -	370.5 25 3.0 X10-4 M 50 3.50 X10-3 M 50 2.9 X10-4 M	SODIUM OLEYL/CIS 9 OCTADECENOYL/	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L		
				GOTT	60018	G L		
				WEIL STIR	59004	T L		
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 62 MOL WGT -	370.5 50 1.8 X10-4 M	SODIUM ELAIDYL/TRANS 9 OCTADECENOYL/SULFATE	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L		
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 63 MOL WGT -	441.4 25 3.1 X10-4 M 50 2.6 X10-4 M	SODIUM 9 10 DICHLORO OCTADECYL SULFATE	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L		
				WEIL STIR	59004	T L		
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 64 MOL WGT -	372.5 53005 40.0 1.65 X10-4 M 40.0 3.0 X10-4 M 50 1.7 X10-4 M 50 2.31 X10-4 M 50 1.9 X10-4 M 50 2.3 X10-4 M 50 1.1 X10-4 M 50 1.1 X10-4 M UNK 1.0 X10-2 D E 0 0001 11 ENTRIES FOR COMPOUND	SODIUM OCTADECYL 1 SULFATE	VALUES FRM REF IN CMC SPECFC CONDCTNCE GRAPH EQUIV CONDCTNCE GRAPH EQUIV CONDCTNCE GRAPH SURFACE TENSION UNSPEC INTERFACIAL TENSION LOGM SURFACE TENSION LOG PLOT VISUAL SPCTR CHNGE PNCN VISUAL SPCTR CHNGE PNCN VISUAL SPCTR CHNGE PNCN GRAPH DATA NOT RETRIEVED	KLIN LANG EVAN EVAN LANG GOTT KLIN LANG KLIN LANG WEIL STIR WEIL STIR DEMC	57022 56006 56006 53005 60018 57022 57022 59004 63013 61031 M	R T L KL L T L G L T L T L T L T L T L R		
	50			LANG	53005			
COMPOUND NO = 65 MOL WGT -	499.7 50 7.0 X10-5 M	TRIETHANOLAMMONIUM OCTADECYL 1 SULFATE	VISUAL SPCTR CHNGE PNCN	WEIL STIR	59004	T L		
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 66 MOL WGT -	232.2 40.0 1.80 X10-1 M	SODIUM OCTYL 2 SULFATE	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3		
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 67 MOL WGT -	288.3 55 2.82 X10-1 D 2.22 E 1 Q N-C10 GLYCEROL ETHER 55 9.781X10-3 M 55 1.76 X10-1 D 6.104X10-3 M	SODIUM DODECYL 2 SULFATE	VISUAL SPCTR CHNGE PNCN VISUAL SPCTR CHNGE PNCN	SCHI FOWK SCHI FOWK	57014 57014	T L T L M		
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 68 MOL WGT -	302.3 40.0 6.50 X10-3 M	SODIUM TRIDEDECYL 2 SULFATE	SPECFC CONDCTNCE GRAPH	EVAN	56006	T 3		
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 69 MOL WGT -	330.4 40.0 1.71 X10-3 M	SODIUM PENTADECYL 2 SULFATE	SPECFC CONDCTNCE GRAPH	EVAN	56006	T L		
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 70 MOL WGT -	358.5 40.0 4.9 X10-4 M	SODIUM HEPTADECYL 2 SULFATE	SPECFC CONDCTNCE GRAPH	EVAN	56006	T L		
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 71 MOL WGT -	372.5 40.0 2.6 X10-4 M	SODIUM OCTADECYL 2 SULFATE	SPECFC CONDCTNCE GRAPH	EVAN	56006	T L		
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 72 MOL WGT -	274.3 40.0 2.89 X10-2 M RM 3.88 X10-2 M	SODIUM UNDECYL 3 SULFATE	SPECFC CONDCTNCE GRAPH VISUAL SPCTR CHNGE PNCN	EVAN WINS	56006 48008	T 3 T L		
2 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 73	MOL WGT —	316.4	SODIUM TETRADECYL 3 SULFATE						
		40	4.3 X10-3 M	CB UNSPECIFIED CONDUCTANCE		WINS	48008	L	L
		40.0	4.30 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
		60	4.85 X10-3 M	CB UNSPECIFIED CONDUCTANCE		WINS	48008	L	L
		RM	4.53 X10-3 M	CG VISUAL SPCTR CHNGE PNCN		WINS	48008	T	L
4 ENTRIES FOR COMPOUND									
COMPOUND NO = 74	MOL WGT —	330.4	SODIUM PENTADECYL 3 SULFATE						
		40.0	2.20 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 75	MOL WGT —	344.4	SODIUM HEXADECYL 4 SULFATE						
		40.0	1.72 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 76	MOL WGT —	372.5	SODIUM OCTADECYL 4 SULFATE						
		40.0	4.5 X10-4 M	CC SPECFC CONDCTNCE GRAPH		EVAN	56006	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 77	MOL WGT —	316.4	SODIUM TETRADECYL 5 SULFATE						
		40	6.9 X10-3 M	CB UNSPECIFIED CONDUCTANCE		WINS	48008	L	L
		40.0	6.75 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	L
		60	7.95 X10-3 M	CB UNSPECIFIED CONDUCTANCE		WINS	48008	L	L
		RM	7.95 X10-3 M	CG VISUAL SPCTR CHNGE PNCN		WINS	48008	T	L
4 ENTRIES FOR COMPOUND									
COMPOUND NO = 78	MOL WGT —	330.4	SODIUM PENTADECYL 5 SULFATE						
		40.0	3.40 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 79	MOL WGT —	386.5	SODIUM NONADECYL 5 SULFATE						
		40.0	3.3 X10-4 M	CC SPECFC CONDCTNCE GRAPH		EVAN	56006	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 80	MOL WGT —	274.3	SODIUM UNDECYL 6 SULFATE						
		40.0	8.3 X10-2 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 81	MOL WGT —	344.4	SODIUM HEXADECYL 6 SULFATE						
		40.0	2.35 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 82	MOL WGT —	372.5	SODIUM OCTADECYL 6 SULFATE						
		40.0	7.2 X10-4 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 83	MOL WGT —	302.3	SODIUM TRIDECYL 7 SULFATE						
		40.0	1.93 X10-2 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 84	MOL WGT —	316.4	SODIUM TETRADECYL 7 SULFATE						
		40	9.7 X10-3 M	CB UNSPECIFIED CONDUCTANCE		WINS	48008	L	L
		40.0	9.70 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	L
		60	1.15 X10-2 M	CB UNSPECIFIED CONDUCTANCE		WINS	48008	L	L
		RM	1.58 X10-2 M	CG VISUAL SPCTR CHNGE PNCN		WINS	48008	T	L
4 ENTRIES FOR COMPOUND									
COMPOUND NO = 85	MOL WGT —	330.4	SODIUM PENTADECYL 8 SULFATE						
		40.0	6.65 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 86	MOL WGT —	344.4	SODIUM HEXADECYL 8 SULFATE						
		40.0	4.25 X10-3 M	CB SPECFC CONDCTNCE GRAPH		EVAN	56006	T	3
1 ENTRIES FOR COMPOUND									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 87	MOL WGT -	358.5 40.0	SODIUM 2.35 X10-3 M	HEPTADECYL 9 CB	SULFATE SPECFC CONDCTNCE GRAPH	EVAN	56006	T L	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 88	MOL WGT -	386.5 40.0	SODIUM 9.4 X10-4 M	1 NONYL DECYL CB	SULFATE SPECFC CONDCTNCE GRAPH	EVAN	56006	T L	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 89	MOL WGT -	526.8 40.0	SODIUM 8. X10-5 M	1 TETRADECYL PENTADECYL CG	SULFATE EQUIV CONDCTNCE MAXIMUM	EVAN	56006	T L	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 90	MOL WGT -	210.3	POTASSIUM DECANOATE						
0.22 E-1 P	BENZENE	25	9.9 X10-2 M	DB	GRAPH DATA NOT RETRIEVED REFRACTIVE INDEX	KLEV	46007	R	
1.05 E-2 M	K CL	25	1.00 X10-1 M	CG	VISUAL SPCTR CHNGE PNCCN	KLEV	48005	T L	
2.29 E-2 M	K CL	25	9.98 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	50003	T L	
3.79 E-2 M	K CL	25	9.5 X10-2 M	BG	UNSPEC SPCTR CHNGE PNCCN	KLEV	52015	T L	
5.59 E-2 M	K CL	25	9.98 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	58011	T L	
7.87 E-2 M	K CL	25	1.06 X10-1 M	DB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	52017	T L	
1.10 E-1 M	K CL	45	9.8 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48016	T L	
1.51 E-1 M	K CL	45	1.18 X10-1 M	DB	REFRACTIVE INDEX	KLEV	48005	T L	
2.27 E-1 M	K CL	50	1.05 X10-1 M	DB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L	
3.44 E-1 M	K CL	50	48016		VALUES FRM REF IN CMC	KOLT STRI	49005	R	
3.86 E-1 M	K CL	50	9.5 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	CORR HARK	46015	T L	
1. E O N	K CL	25	9.3 X10-2 M	CG	VISUAL SPCTR CHNGE PNCCN	KLEV	50003	T L	
. E O	K OH	25	9.49 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
2. E O I	K OH	25	9.17 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
4.4 E-2 N	K OH	25	8.85 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
4.4 E-2 N	K OH	25	8.38 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
4.4 E-2 N	K OH	25	7.87 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
2. E O I	K OH	25	7.31 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
. E O	K OH	25	6.45 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
2.27 E-1 M	K CL	25	5.68 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
3.44 E-1 M	K CL	25	4.69 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
3.86 E-1 M	K CL	25	4.29 X10-2 M	DG	VISUAL SPCTR CHNGE PNCCN	HERZ	52017	T L	
1. E-1 N	K2 SO4	50	2.2 X10-2 M	DC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L	
2. E-1 N	K2 SO4	50	1.09 X10-1 M	BG	VAPR PRESSURE LOWERING	HUFF MCBA	51004	T L	
2. E-1 N	K2 SO4	50	7. X10-2 M	XG	VISCOSITY MINIMUM	SHIN	55004	S L	
1. E-1 N	K2 SO4	20	5.7 X10-2 M	CE	DIFFUSION COEFFICIENT	TYUZ	53006	T L	
2. E-1 N	K2 SO4	20	1.06 X10-1 M	CE	DIFFUSION COEFFICIENT	TYUZ	61025	T L	
0.044		25	1.00 X10-1 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	54005	G L	
0.091		30	9.8 X10-2 W	CB	VAPR PRESSURE LOWERING	HUFF MCBA	51004	T L	
0.091		50	1.15 X10-1 W	CB	VAPR PRESSURE LOWERING	HUFF MCBA	51004	T L	
0.092		50	8.2 X10-2 M	DB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L	
0.092		50	6.5 X10-2 M	DB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L	
0.188		50	4.5 X10-2 M	DC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L	
3.63 E-2 M	BUTANOL-1	25	SEE CMPD NMBR IN ADDITV			SHIN	54005	X	
2. E O I	K OH	10	1.01 X10-1 M	BG	SEE CMPD NMBR IN ADDITV	CORR HARK	46015	X	
6.74 E-2 M	BUTANOL-1	10	5.2 X10-2 M	DC	REFRACTIVE INDEX	KLEV	48005	T L	
2. E O I	K OH	10	9.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
9.52 E-2 M	BUTANOL-1	10	1.5 X10-2 M	DD	REFRACTIVE INDEX	KLEV	48005	T L	
2. E O I	K OH	10	1.2 X10-2 M	DD	REFRACTIVE INDEX	KLEV	48005	T L	
1.44 E-1 M	BUTANOL-1	10	SEE CMPD NMBR IN ADDITV			SHIN	54005	X	
2. E O I	K OH	10	8.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
1.80 E-1 M	BUTANOL-1	10	7.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
2. E O I	K OH	10	6.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
2.29 E-1 M	BUTANOL-1	10	1.06 X10-1 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
2. E O I	K OH	10	1.02 X10-1 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
6.4 E-5 M	DECANOL-1	10	1.01 X10-1 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
2. E O I	K OH	10	9.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
1.20 E-4 M	DECANOL-1	10	1.01 X10-1 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
2. E O I	K OH	10	9.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
1.64 E-4 M	DECANOL-1	10	1.01 X10-1 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
2. E O I	K OH	10	9.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
5.44 E-1 M	ETHANOL	10	9.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCCN	SHIN	55004	T L	
2. E O I	K OH	10				SHIN	55004	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
9.9	E-1 M ETHANOL 2. E O I K OH	10	8.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.37	E O M ETHANOL 2. E O I K OH	10	8.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.91	E O M ETHANOL 2. E O I K OH	10	6.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.33	E O M ETHANOL 2. E O I K OH	10	6.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.82	E O M ETHANOL 2. E O I K OH	10	5.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
9.0	E-3 M HEXANOL-1 2. E O I K OH	10	9.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.60	E-2 M HEXANOL-1 2. E O I K OH	10	8.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.12	E-2 M HEXANOL-1 2. E O I K OH	10	7.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.85	E-2 M HEXANOL-1 2. E O I K OH	10	6.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.70	E-3 M HEPTANOL-1 2. E O I K OH	10	9.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.73	E-3 M HEPTANOL-1 2. E O I K OH	10	8.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
8.35	E-3 M HEPTANOL-1 2. E O I K OH	10	7.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
7.8	E-4 M OCTANOL-1 2. E O I K OH	10	1.02 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.49	E-3 M OCTANOL-1 2. E O I K OH	10	9.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.11	E-3 M OCTANOL-1 2. E O I K OH	10	9.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.70	E-3 M OCTANOL-1 2. E O I K OH	10	8.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.30	E-1 M PROPANOL-1 2. E O I K OH	10	9.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.35	E-1 M PROPANOL-1 2. E O I K OH	10	8.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.51	E-1 M PROPANOL-1 2. E O I K OH	10	8.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.82	E-1 M PROPANOL-1 2. E O I K OH	10	7.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
6.34	E-1 M PROPANOL-1 2. E O I K OH	10	6.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
8.25	E-1 M PROPANOL-1 2. E O I K OH	10	5.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.91	E-4 M NONANOL-1 2. E O I K OH	10	1.02 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.64	E-4 M NONANOL-1 2. E O I K OH	10	9.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.84	E-4 M NONANOL-1 2. E O I K OH	10	9.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
5.70	E-4 M NONANOL-1 2. E O I K OH 0044	10	9.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.9	E O C 0091 2. E O I K OH	25		SEE CMPD NMBR IN ADDITV	SHIN	54003	X		
3.8	E O C 0091 2. E O I K OH	25	9.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
9.0	E O C 0091 2. E O I K OH	25	8.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.55	E 1 C 0091 2. E O I K OH	25	7.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.34	E 1 C 0091 2. E O I K OH	25	6.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.80	E 1 C 0091 2. E O I K OH	25	5.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.58	E 1 C 0091 2. E O I K OH	25	4.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
4.75	E 1 C 0091 2. E O I K OH	25	4.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
6.80	E 1 C 0091 2. E O I K OH	25	3.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.0	E O C 0092 2. E O I K OH	25	3.3 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.7	E O C 0092 2. E O I K OH	25	8.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.5	E O C 0092 2. E O I K OH	25	7.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
		25	5.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
4.2	E O C 0092	25	4.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
6.9	E O C 0092	25	3.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
1.13	E 1 C 0092	25	2.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
1.78	E 1 C 0092	25	2.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
3.18	E 1 C 0092	25	1.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
	0296	25			SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
2.	E O I K OH								
4.8	E O C 0297	25	9.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
1.15	E 1 C 0297	25	8.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
1.78	E 1 C 0297	25	8.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
2.53	E 1 C 0297	25	7.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
3.28	E 1 C 0297	25	7.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
4.39	E 1 C 0297	25	6.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
5.36	E 1 C 0297	25	6.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
6.65	E 1 C 0297	25	5.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
8.18	E 1 C 0297	25	5.3 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.	E O I K OH								
9.6	E-3 M 3-METHYL BUTANOL-1	10	1.06 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.	E O I K OH								
1.99	E-2 M 3-METHYL BUTANOL-1	10	9.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.	E O I K OH								
3.03	E-2 M 3-METHYL BUTANOL-1	10	9.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.	E O I K OH								
4.60	E-2 M 3-METHYL BUTANOL-1	10	8.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.	E O I K OH								
5.79	E-2 M 3-METHYL BUTANOL-1	10	8.3 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.	E O I K OH								
8.14	E-2 M 3-METHYL BUTANOL-1	10	7.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.	E O I K OH								
	0044	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
5.9	E O C 0091	25	3.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
6.4	E O C 0092								
2.	E O I K OH								
1.24	E 1 C 0091	25	3.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
5.9	E O C 0092								
2.	E O I K OH								
1.94	E 1 C 0091	25	3.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
5.5	E O C 0092								
2.	E O I K OH								
2.74	E 1 C 0091	25	3.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
4.9	E O C 0092								
2.	E O I K OH								
3.62	E 1 C 0091	25	3.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
4.3	E O C 0092								
2.	E O I K OH								
4.59	E 1 C 0091	25	3.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
3.7	E O C 0092								
2.	E O I K OH								
5.68	E 1 C 0091	25	2.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
2.9	E O C 0092								
2.	E O I K OH								
6.92	E 1 C 0091	25	2.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
2.1	E O C 0092								
2.	E O I K OH								
8.35	E 1 C 0091	25	2.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	G L	
1.1	E O C 0092								
2.	E O I K OH								
2.3	E O C 0297	25	9.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.2	E O C 0091								
2.	E O I K OH								
4.9	E O C 0297	25	8.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.5	E O C 0091								
2.	E O I K OH								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives			Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
8.0 E O C 0297			25	7.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
4.1 E O C 0091										
2. E O I K OH										
1.17 E 1 C 0297			25	7.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
6.0 E O C 0091										
2. E O I K OH										
1.61 E 1 C 0297			25	6.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
8.2 E O C 0091										
2. E O I K OH										
2.14 E 1 C 0297			25	5.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.09 E 1 C 0091										
2. E O I K OH										
2.82 E 1 C 0297			25	5.3 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.44 E 1 C 0091										
2. E O I K OH										
3.75 E 1 C 0297			25	4.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.91 E 1 C 0091										
2. E O I K OH										
4.96 E 1 C 0297			25	4.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.52 E 1 C 0091										
2. E O I K OH										
130 ENTRIES FOR COMPOUND										
COMPOUND NO = 91 MOL WGT -	238.4	POTASSIUM DODECANOATE								
	47005				VALUES FRM REF IN CMC		KLEV	48005	R	
	47006				VALUES FRM REF IN CMC		CORR HARK	46005	R	
					GRAPH DATA NOT RETRIEVED		KLEV	46007	R	
	48016				VALUES FRM REF IN CMC		KOLT STRI	49005	R	
	15.0 2.35 X10-2 M	DG			FOTOMTR SPCTR CHNGE PNCN		KLEV	47004	T L	
	23.5 6.15 X10-1 D	DB			REFRACTIVE INDEX		KLEV	46012	T L	
	2.579X10-2 M								M	
	25 9. X10-1 P	BD			DENSITY		BURY PARR	35008	T L	
	3.7 X10-2 S								M	
	25 2.55 X10-2 M	XG			VISUAL SPCTR CHNGE		KLEV	53010	T L	
	25 2.4 X10-2 M	CG			VISUAL SPCTR CHNGE PNCN		KLEV	50003	T L	
	25.6 2.3 X10-2 M	DG			FOTOMTR SPCTR CHNGE PNCN		CORR HARK	46002	T L	
	25.6 2.25 X10-2 M	DG			FOTOMTR SPCTR CHNGE PNCN		KLEV	47004	T L	
	25 2.34 X10-2 M	DG			VISUAL SPCTR CHNGE PNCN		HERZ	52015	T L	
	25 2.55 X10-2 M	DC			REFRACTIVE INDEX		KLEV	47005	T L	
	25 2.5 X10-2 M	BG			UNSPEC SPCTR CHNGE PNCN		KLEV	58011	T L	
	26 2.35 X10-2 M	DG			VISUAL SPCTR CHNGE PNCN		CORR HARK	47006	T L	
	26 2.34 X10-2 M	DG			VISUAL SPCTR CHNGE RH6		CORR HARK	47006	T L	
	26 2.30 X10-2 M	DG			VISUAL SPCTR CHNGE PNCN		CORR HARK	47010	T L	
	30 2.60 X10-2 M	DB			REFRACTIVE INDEX		KLEV	48005	T L	
	30 2.35 X10-2 M	DB			FOTOMTR SOLUBLZTN PDMAB		KOLT STRI	48016	T L	
	30 2.35 X10-2 M	DB			FOTOMTR SOLUBLZTN OROT		KOLT STRI	48016	T L	
	30 2.3 X10-2 M	DC			UNSPEC SOLUBLZTN PDMAB		KOLT JOHN	48025	T L	
	30 2.2 X10-2 M	DC			ELECTROMOTIVE FORCE		KOLT JOHN	48025	T L	
	30 2.47 X10-2 M	DB			FOTOMTR SOLUBLZTN AZBZ		KOLT STRI	49005	T L	
	35 8. X10-1 P	BD			DENSITY		BURY PARR	35008	T L	
	3.3 X10-2 S								M	
	35.8 2.13 X10-2 M	DG			FOTOMTR SPCTR CHNGE PNCN		KLEV	47004	T L	
	35 1.9 X10-2 M	XG			VELOCITY OF SOUND		KUPP SURY	65028	T L	
	35 2.70 X10-2 M	DC			REFRACTIVE INDEX		KLEV	47005	T L	
	45.0 2.08 X10-2 M	DG			FOTOMTR SPCTR CHNGE PNCN		KLEV	47004	T L	
	45 3.05 X10-2 M	DC			REFRACTIVE INDEX		KLEV	47005	T L	
	50 2.45 X10-2 M	DB			FOTOMTR SOLUBLZTN PDMAB		KOLT STRI	48016	T L	
	50 2.85 X10-2 M	DB			FOTOMTR SOLUBLZTN OROT		KOLT STRI	48016	T L	
	50 2.14 X10-2 M	DG			UNSPEC SPCTR CHNGE PNCN		RAIS	52016	T L	
	50 2.1 X10-2 M	DC			FOTOMTR SOLUBLZTN AZBZ		KOLT STRI	49005	T L	
	55.0 2.08 X10-2 M	DG			FOTOMTR SPCTR CHNGE PNCN		KLEV	47004	T L	
	55 3.50 X10-2 M	DC			REFRACTIVE INDEX		KLEV	47005	T L	
	60 2.6 X10-2 M	DG			SOLUBLZTN TOLUENE		DEMC DUMA	60032	T L	
	60 5.8 X10-1 D	DG			SOLUBLZTN TOLUENE		DEMC	60034	T L	
	2.43 X10-2 M								M	
	60 2.3 X10-2 M	DC			FOTOMTR SOLUBLZTN OROT		MERR GETT	48024	T L	
	65 4.20 X10-2 M	DC			REFRACTIVE INDEX		KLEV	47005	T L	
	UNK 6.2 X10-1 D	DG			VISUAL SPCTR CHNGE PNCN		DEMC	60034	T L	
	2.60 X10-2 M								M	
	UNK 2.33 X10-2 M	DG			VISUAL SPCTR CHNGE PNCN		CORR HARK	46015	G L	
	UNK 6.0 X10-1 D	DG			METHOD NOT CITED		DEMC	62037	T L	
	2.51 X10-2 M								M	
	UNK 2.40 X10-2 M	DG			METHOD NOT CITED		HARK MITT	49006	T L	
	UNK 2.55 X10-2 M	DG			METHOD NOT CITED		ARRI PATT	53003	T L	
	25 2.0 X10-2 M	CG			VISUAL SPCTR CHNGE PNCN		KLEV	50003	T L	
2.2 E-1 P BENZENE										
4.4 E-2 P BENZENE										
6.1 E-2 P BENZENE										
1.00 E-1 P BENZENE										

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C.	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.39 E-1 P	BENZENE	UNK	2.29 X10-2 M	DG	METHOD NOT CITED	HARK MITT	49006	T L
1.50 E-1 P	BENZENE	UNK	2.30 X10-2 M	DG	METHOD NOT CITED	HARK MITT	49006	T L
2.65 E-2 M	K BR	25	1.55 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
1.735E-1 M	K BR	25	7.9 X10-3 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
2.52 E-2 M	K CL	25	1.59 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
6.70 E-2 M	K CL	25	1.19 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
1.005E-1 M	K CL	25	1.02 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
2.124E-1 M	K CL	25	7.1 X10-3 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
3.825E-1 M	K CL	25	4.8 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L
5.014E-1 M	K CL	25	4.1 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L
8.380E-3 M	K CL	26	1.97 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.321E-2 M	K CL	26	1.76 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.680E-2 M	K CL	26	1.69 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
3.872E-2 M	K CL	26	1.29 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
4.217E-2 M	K CL	26	1.25 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
4.263E-2 M	K CL	26	1.23 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
5.471E-2 M	K CL	26	1.12 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
6.466E-2 M	K CL	26	1.07 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
6.631E-2 M	K CL	26	1.01 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
7.466E-2 M	K CL	26	9.92 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.038E-1 M	K CL	26	8.44 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.062E-1 M	K CL	26	8.30 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.199E-1 M	K CL	26	7.96 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.485E-1 M	K CL	26	7.48 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.739E-1 M	K CL	26	5.77 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
2.411E-1 M	K CL	26	5.47 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
2.617E-1 M	K CL	26	5.27 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
3.142E-1 M	K CL	26	4.58 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
3.335E-1 M	K CL	26	4.64 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
4.338E-1 M	K CL	26	3.53 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
5.927E-1 M	K CL	26	3.48 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1. E O N	K CL	30	2.1 X10-3 M	DC	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L
1. E O N	K CL	30	2.3 X10-3 M	DC	FOTOMTR SOLUBLZTN AZBZ	KOLT STRI	49005	T L
1. E O N	K CL	30	2.4 X10-3 M	DC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
5. E-2 N	K CL	50	1.77 X10-2 M	DB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
1. E-1 N	K CL	50	1.1 X10-2 M	DD	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
5. E-1 N	K CL	50	4. X10-3 M	DE	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
1. E O N	K CL	50	2.6 X10-3 M	DC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
1. E O N	K CL	50	3. X10-3 M	DE	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L
1. E O N	K CL	50	2.3 X10-3 M	DC	FOTOMTR SOLUBLZTN AZBZ	KOLT STRI	49005	T L
. E O	K CL	UNK		GRAPH DATA NOT RETRIEVED	DEM C	62037	R	
1.66 E-2 M	K I	25	1.81 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
2.615E-1 M	K I	25	6.3 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L
3.06 E-2 M	K NO3	25	1.54 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
1.38 E-1 M	K NO3	25	8.8 X10-3 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
3.33 E-1 M	K NO3	25	5.3 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L
1.0 E-1 M	K NO3	30	1.2 X10-2 M	DD	UNSPEC SOLUBLZTN PDMAB	KOLT JOHN	48025	T L
1.0 E-1 M	K NO3	30	1.2 X10-2 M	DD	ELECTROMOTIVE FORCE	KOLT JOHN	48025	T L
5.0 E-1 M	K NO3	30	5.5 X10-3 M	DC	ELECTROMOTIVE FORCE	KOLT JOHN	48025	T L
5.0 E-1 M	K NO3	30	5. X10-3 M	DD	UNSPEC SOLUBLZTN PDMAB	KOLT JOHN	48025	T L
1. E-1 N	K NO3	50	1.17 X10-2 M	DB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
1. E O N	K NO3	50	1.17 X10-2 M	DC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
. E O	K NO3	50	2.5 X10-3 M	DC	GRAPH DATA NOT RETRIEVED	DEM C	62037	R
. E O	K OH	0	3.75 X10-2 W	CB	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
4. E O I	K OH	0	3.5 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L
4. E O I	K OH	0	3.3 X10-2 W	CD	FREEZING POINT	FINE MCBA	48011	K L
2. E O I	K OH	10	2.91 X10-2 M	BG	VISUAL SPCTR CHNGE PNCC	SHIN	55004	S L
2. E O I	K OH	25	2.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCC	SHIN	54005	G L
4. E O I	K OH	25	2.4 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L
4. E O I	K OH	25	2.4 X10-2 M	CB	EQUIV CONDCTNCE GRAPH	BRAD MCBA	48009	T L
4. E O I	K OH	25	1.7 X10-2 W	CG	VISUAL SPCTR CHNGE RHD6	FINE MCBA	48011	T L
4. E O I	K OH	25	2.5 X10-2 W	CD	VAPR PRESURE LOWERING	FINE MCBA	48011	T L
. E O	K OH	30	2.05 X10-2 W	CB	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
1. E O N	K OH	30	2.6 X10-3 M	CC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L
1. E O N	K OH	30	2.7 X10-3 M	CC	FOTOMTR SOLUBLZTN OROT	KOLT STRI	49005	T L
. E O	K OH	50	2.05 X10-2 W	CB	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
4. E O I	K OH	50	2.0 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L
4. E O I	K OH	70	2.2 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L
4. E O I	K OH	90	3.1 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L
1.87 E-2 M	K4 P207 PYRO	25	8.8 X10-3 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
5.07 E-2 M	K4 P207 PYRO	25	6.9 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L
7.42 E-2 M	K4 P207 PYRO	25	5.3 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L
1.88 E-2 M	K2 S04	25	1.41 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
5.25 E-2 M	K2 S04	25	1.03 X10-2 M	DB	REFRACTIVE INDEX	KLEV	48005	T L
1.38 E 0 M	K2 S04	25	5.9 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L
7.65 E-3 M	K2 S04	26	1.74 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
1.76 E-2 M	K2 S04	26	1.33 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L
2.92 E-2 M	K2 S04	26	1.10 X10-2 M	DG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l (or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
4.66 E-2 M K2 S04	26	9.12 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
6.57 E-2 M K2 S04	26	7.70 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
1.04 E-1 M K2 S04	26	5.94 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
1.42 E-1 M K2 S04	26	4.84 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
1.87 E-1 M K2 S04	26	4.25 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
1. E-1 N K2 S04	50	1.1 X10-2 M	DD	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L	
5. E-1 N K2 S04	50	4.5 X10-3 M	DC	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	49005	T L	
. E O K2 S04	UNK	GRAPH DATA NOT RETRIEVED	DEMC			62037	R	
4. E 2 I NA CL	UNK	3.2 X10-1 D	DG	METHOD NOT CITED	DEMC	62037	T L	
		1.34 X10-2 M					M	
5.18 E-3 M NA4 P207 PYRO	26	1.65 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
1.39 E-2 M NA4 P207 PYRO	26	1.10 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
2.15 E-2 M NA4 P207 PYRO	26	8.56 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	47010	T L	
3.05 E-2 M NA4 P207 PYRO	26	7.47 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HAR2	47010	T L	
4. E 2 I NA2 S04	UNK	2.6 X10-1 D	DG	METHOD NOT CITED	DEMC	62037	T L	
		1.09 X10-2 M					M	
2.5 E-2 M NA2 SI03 META	60	1.3 X10-2 M	HD	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	K L	
1.25 E-2 M SI02/NA20=1.60	60	1.8 X10-2 M	HD	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	T L	
2.5 E-2 M SI02/NA20 = 1.60	60	1.4 X10-2 M	HD	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	T L	
1.0 E 1 PH OF SOLUTION	UNK	2.5 X10-2 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L	
1. E-5 M PINACYANOL CL (DYE)	25.8	2.17 X10-2 M	DG	FOTOMTR SPCTR CHNGE PNCN	CORR KLEV	46010	T L	
1. E-4 M PINACYANOL CL (DYE)	25.8	2.35 X10-2 M	DG	FOTOMTR SPCTR CHNGE PNCN	CORR KLEV	46010	T L	
		0001		SEE CMPD NMBR IN ADDITV	CORR HARK	46015	X	
		0044		SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
		0090		SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
7.0 E 0 C 0090	UNK	2.39 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
1.18 E 1 C 0090	UNK	2.48 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
2.51 E 1 C 0090	UNK	2.71 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
2.79 E 1 C 0090	UNK	2.83 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
3.29 E 1 C 0090	UNK	2.92 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
3.95 E 1 C 0090	UNK	3.12 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
4.92 E 1 C 0090	UNK	3.41 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
5.67 E 1 C 0090	UNK	3.61 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
6.20 E 1 C 0090	UNK	3.83 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
6.90 E 1 C 0090	UNK	4.20 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
7.68 E 1 C 0090	UNK	4.82 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
8.14 E 1 C 0090	UNK	5.2 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
8.63 E 1 C 0090	UNK	5.9 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
9.32 E 1 C 0090	UNK	7.1 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G L	
1.52 E 1 C 0092	25	1.4 X10-2 M	DD	REFRACTIVE INDEX	KLEV	48005	T L	
3.01 E 1 C 0092	25	1.1 X10-2 M	DD	REFRACTIVE INDEX	KLEV	48005	T L	
4.52 E 1 C 0092	25	1.0 X10-2 M	DD	REFRACTIVE INDEX	KLEV	48005	T L	
6.45 E 1 C 0092	25	7.8 X10-3 M	DC	REFRACTIVE INDEX	KLEV	48005	T L	
0.92 E-2 M BUTANOL-1	10	2.46 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
1.65 E-1 M BUTANOL-1	10	2.22 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
2.42 E-1 M BUTANOL-1	10	1.95 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
3.54 E-1 M BUTANOL-1	10	1.58 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
4.20 E-1 M BUTANOL-1	10	1.21 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
4.96 E-1 M BUTANOL-1	10	1.00 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
4.9 E-5 M DECANOL-1	10	2.81 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
8.1 E-5 M DECANOL-1	10	2.76 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
1.56 E-4 M DECANOL-1	10	2.62 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
4.01 E-1 M ETHANOL	10	2.74 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
7.62 E-1 M ETHANOL	10	2.61 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
1.14 E 0 M ETHANOL	10	2.32 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
1.67 E 0 M ETHANOL	10	2.14 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
2.24 E 0 M ETHANOL	10	1.92 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
7.1 E-3 M HEXANOL-1	10	2.60 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
9.3 E-3 M HEXANOL-1	10	2.53 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								
1.31 E-2 M HEXANOL-1	10	2.40 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.99 E-2 M HEXANOL-1 2. E O I K OH	10	2.18 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.70 E-2 M HEXANOL-1 2. E O I K OH	10	1.85 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.61 E-2 M HEXANOL-1 2. E O I K OH	10	1.58 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.17 E-3 M HEPTANOL-1 2. E O I K OH	10	2.73 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.12 E-3 M HEPTANOL-1 2. E O I K OH	10	2.58 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.00 E-2 M HEPTANOL-1 2. E O I K OH	10	1.88 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1. E-1 K K ION 1.00 E 1 PH OF SOLUTION	UNK	1.37 X10-2 M	CB	SURFACE TENSION LOG PLOT	ROE BRAS	54013	T L	
4. E O I K OH 1.4 E-1 I N-HEXANE	0	2.8 X10-2 W	CD	FREEZING POINT	FINE MCBA	48011	T L	
4. E-2 K NA CL 1.05 E 1 PH OF SOLUTION	20	1.7 X10-2 M	BC	INTERFACIAL TENSION LOGM	V VO	60025	T L	
7.15 E-4 M OCTANOL-1 2. E O I K OH	10	2.65 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
9.30 E-4 M OCTANOL-1 2. E O I K OH	10	2.58 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.27 E-3 M OCTANOL-1 2. E O I K OH	10	2.40 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.84 E-3 M OCTANOL-1 2. E O I K OH	10	2.32 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.02 E-1 M PROPANOL-1 2. E O I K OH	10	2.70 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.96 E-1 M PROPANOL-1 2. E O I K OH	10	2.59 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.56 E-1 M PROPANOL-1 2. E O I K OH	10	2.44 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
5.03 E-1 M PROPANOL-1 2. E O I K OH	10	2.28 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
6.95 E-1 M PROPANOL-1 2. E O I K OH	10	2.09 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
8.06 E-1 M PROPANOL-1 2. E O I K OH	10	1.82 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
9.80 E-1 M PROPANOL-1 2. E O I K OH	10	1.67 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.39 E-4 M NONANOL-1 2. E O I K OH	10	2.79 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.70 E-4 M NONANOL-1 2. E O I K OH	10	2.69 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.88 E-4 M NONANOL-1 2. E O I K OH 0296	10	2.61 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH 0297	25		SEE CMPD NMBR IN ADDITV	SHIN	54005	X		
2. E O I K OH	25		SEE CMPD NMBR IN ADDITV	SHIN	54003	X		
2.76 E-2 M 3-METHYL BUTANOL-1 2. E O I K OH	10	2.67 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
6.20 E-2 M 3-METHYL BUTANOL-1 2. E O I K OH	10	2.31 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.02 E-1 M 3-METHYL BUTANOL-1 2. E O I K OH	10	1.91 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.38 E-1 M 3-METHYL BUTANOL-1 2. E O I K OH	10	1.61 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.71 E-1 M 3-METHYL BUTANOL-1 2. E O I K OH 0044 0090 0090	10	1.45 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
217 ENTRIES FOR COMPOUND			SEE CMPD NMBR IN ADDITV	SHIN	54003	X		
SEE CMPD NMBR IN ADDITV	25		SEE CMPD NMBR IN ADDITV	SHIN	54003	X		
SEE CMPD NMBR IN ADDITV	25		SEE CMPD NMBR IN ADDITV	SHIN	54003	X		

COMPOUND NO = 92 MOL WGT = 266.5 POTASSIUM TETRADECANOATE

47005	VALUES FRM REF IN CMC	KLEV	48005	R
25 6.3 X10-3 M	CG VISUAL SPCTR CHNGE PNCN	KLEV	50003	T L
25.6 6.0 X10-3 M	DG FOTOMTR SPCTR CHNGE PNCN	CORR HARK	46002	T L
25.8 6.4 X10-3 M	DG FOTOMTR SPCTR CHNGE PNCN	CORR KLEV	46010	T L
25.6 6.4 X10-3 M	DG FOTOMTR SPCTR CHNGE PNCN	KLEV	47004	T L
25 5.9 X10-3 M	DG VISUAL SPCTR CHNGE PNCN	HERZ	52015	T L
25 6.6 X10-3 M	DC REFRACTIVE INDEX	KLEV	47005	T L
25 5.8 X10-3 M	BG UNSPEC SPCTR CHNGE PNCN	KLEV	58011	T L
30 7.0 X10-3 M	DB FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L
35.8 5.7 X10-3 M	DG FOTOMTR SPCTR CHNGE PNCN	CORR HARK	46002	T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
8. E-3 P BENZENE	35.8	5.7 X10-3 M	DG	FOTOMTR SPCTR CHNGE PNCN	KLEV	47004	T	L
1.5 E-2 P BENZENE	35	7.0 X10-3 M	DC	REFRACTIVE INDEX	KLEV	47005	T	L
2.7 E-2 P BENZENE	45.0	5.5 X10-3 M	DG	FOTOMTR SPCTR CHNGE PNCN	KLEV	47004	T	L
4.0 E-2 P BENZENE	45	7.4 X10-3 M	DC	REFRACTIVE INDEX	KLEV	47005	T	L
5.4 E-2 P BENZENE	50	7.2 X10-3 M	DB	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T	L
6.1 E-2 P BENZENE	50	5. X10-3 M	DE	FOTOMTR SOLUBLZTN PDMAB	KOLT JOHN	46006	T	L
8.5 E-2 P BENZENE	50	48016		VALUES FRM REF IN CMC	KOLT STRI	49005	R	
8.7 E-2 P BENZENE	55.0	5.5 X10-3 M	DG	FOTOMTR SPCTR CHNGE PNCN	KLEV	47004	T	L
1.04 E-1 P BENZENE	55	7.9 X10-3 M	DC	REFRACTIVE INDEX	KLEV	47005	T	L
1.08 E-1 P BENZENE	65	8.6 X10-3 M	DC	REFRACTIVE INDEX	KLEV	47005	T	L
1.31 E-1 P BENZENE	UNK	3. X10-3 M	DE	ELECTROMOTIVE FORCE	CARR JOHN	47013	T	L
1.55 E-1 P BENZENE	UNK	5. X10-3 M	DE	FOTOMTR SOLUBLZTN PDMAB	CARR JOHN	47013	T	L
1.82 E-1 P BENZENE	UNK	6.4 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	CORR HARK	46015	G	L
2.20 E-1 P BENZENE	UNK	6.6 X10-3 M	DG	METHOD NOT CITED	ARRI PATT	53003	T	L
2.81 E-1 P BENZENE	25	6.2 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	G	L
3.35 E-1 P BENZENE	25	6.1 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	G	L
3.93 E-1 P BENZENE	25	5.9 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	G	L
5. E-2 N K CL	25	5.7 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	G	L
5. E-1 N K CL	25	5.9 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	G	L
2. E O I K OH	18	7.04 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	SHIN	55004	T	L
2. E O I K OH	25	5.6 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G	L
4. E O I K OH	25	5.7 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	G	L
. E O K OH	25	5.6 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	KLEV	50003	G	L
. E O K OH	30	4. X10-3 W	CE	VAPR PRESURE LOWERING	HUFF MCBA	51004	T	L
1. E-3 N K OH	50	5. X10-3 W	CE	VAPR PRESURE LOWERING	HUFF MCBA	51004	T	L
1. E-2 N K OH	UNK	7. X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T	L
1. E-1 N K OH	UNK	6. X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T	L
1.0 E 1 PH OF SOLUTION	UNK	4.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T	L
0001	UNK	7.0 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T	L
. E O 0001	SEE CMPD NMBR IN ADDITV				KLEV	48005	X	
. E O 0001	GRAPH DATA NOT RETRIEVED				KLEV	46007	R	
. E O 0040	SEE CMPD NMBR IN ADDITV				CORR HARK	46015	X	
. E O 0044	GRAPH DATA NOT RETRIEVED				KLEV	46007	R	
. E O 0044	25				SHIN	54005	X	
. E O 0090					KLEV	48005	X	
. E O 0090	25				SHIN	54005	X	
. E O 0091					KLEV	48005	X	
. E O 0091	0091				GRAPH DATA NOT RETRIEVED	KLEV	46007	R
. E O 0188	25				SEE CMPD NMBR IN ADDITV	SHIN	54005	X
1.9 E 1 C 0372	UNK	8. X10-3 M	DG	METHOD NOT CITED	ARRI PATT	53003	G	L
3.0 E 1 C 0372	UNK	8.2 X10-3 M	DG	METHOD NOT CITED	ARRI PATT	53003	G	L
5.6 E 1 C 0372	UNK	1.3 X10-2 M	DG	METHOD NOT CITED	ARRI PATT	53003	G	L
8.5 E 1 C 0372	UNK	2.1 X10-2 M	DG	METHOD NOT CITED	ARRI PATT	53003	G	L
9.4 E 1 C 0372	UNK	2.9 X10-2 M	DG	METHOD NOT CITED	ARRI PATT	53003	G	L
1.20 E-1 M BUTANOL-1	18	5.70 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
2. E O I K OH	18	4.75 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
2.02 E-1 M BUTANOL-1	18	3.68 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
3.18 E-1 M BUTANOL-1	18	3.02 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
2. E O I K OH	18	2.59 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
3.92 E-1 M BUTANOL-1	18	6.78 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
2. E O I K OH	18	6.10 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
4.48 E-1 M BUTANOL-1	18	5.76 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
2. E O I K OH	18	6.51 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T	L
4.5 E-5 M DECANOL-1					SHIN	55004	T	L
2. E O I K OH					SHIN	55004	T	L
8.9 E-5 M DECANOL-1					SHIN	55004	T	L
2. E O I K OH					SHIN	55004	T	L
1.91 E-4 M DECANOL-1					SHIN	55004	T	L
2. E O I K OH					SHIN	55004	T	L
5.5 E-1 M ETHANOL					SHIN	55004	T	L
2. E O I K OH					SHIN	55004	T	L

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality

counterions: M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l (or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.00 E O M ETHANOL	18	5.98 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	5.1 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.74 E O M ETHANOL	18	4.52 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	3.97 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.44 E O M ETHANOL	18	5.64 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	5.08 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.18 E-2 M HEXANOL-1	18	4.58 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	3.90 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.08 E-2 M HEXANOL-1	18	5.91 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	5.09 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2.50 E-3 M HEPTANOL-1	18	4.03 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	4.03 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
5.52 E-3 M HEPTANOL-1	18	4.03 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	4.03 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1. E-3 N K OH	UNK	5.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L	
5. E-2 N K CL	UNK	4.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L	
1. E-3 N K OH	UNK	3. X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L	
1. E-1 N K CL	UNK	1.5 X10-3 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L	
1. E-3 N K OH	UNK	8. X10-4 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L	
1. E O N K CL	18	6.47 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
5.5 E-4 M OCTANOL-1	18	5.82 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	5.36 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.39 E-3 M OCTANOL-1	18	4.80 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	5.94 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
3.08 E-1 M PROPANOL-1	18	5.26 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	4.71 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
7.28 E-1 M PROPANOL-1	18	4.18 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	3.52 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.25 E O M PROPANOL-1	18	2.97 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	6.62 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.65 E-4 M NONANOL-1	18	6.17 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	5.90 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
4.41 E-4 M NONANOL-1	18	5.55 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
6.82 E-4 M NONANOL-1	18	0296		SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
2. E O I K OH	25	5.12 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
5.95 E-2 M 3-METHYL BUTANOL-1	18	4.17 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	3.48 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.29 E-1 M 3-METHYL BUTANOL-1	18	2.64 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
2. E O I K OH	18	2.64 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55004	T L	
1.83 E-1 M 3-METHYL BUTANOL-1	18	0090		SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
2. E O I K OH								
113 ENTRIES FOR COMPOUND								

COMPOUND NO = 93 MOL WGT = 252.2 OCTYL TRIMETHYL AMMONIUM BROMIDE
 20 2.82 X10-1 M BB INTERFACIAL TENSION LOGM HAYD TAYL 62004 L L
 25 1.4 X10-1 M CD REFRACTIVE INDEX KLEV 48005 T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
 D—wt/vol %; E—% saturation; F—wt % solvent; G—mol % surfactant; H—normality
 counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg;
 T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.25 E-2 M	K BR	25	1.3 X10-1 M	CE	EQUIV CONDCTNCE GRAPH	SCOT TART	43003	K	L
2.5 E-2 M	K BR	30	2.24 X10-1 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
1.0 E-1 M	NA CL	40	1.5 X10-1 M	CE	EQUIV CONDCTNCE GRAPH	SCOT TART	43003	K	L
2.5 E-1 M	NA CL	60	1.3 X10-1 M	CE	EQUIV CONDCTNCE GRAPH	SCOT TART	43003	K	L
5.0 E-1 M	NA CL	30	2.26 X10-1 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
11 ENTRIES FOR COMPOUND		30	2.20 X10-1 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
COMPOUND NO = 94 MOL WGT -		20	2.69 X10-1 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	L
1.25 E-2 M	K BR	20	2.54 X10-1 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	L
2 ENTRIES FOR COMPOUND		20	2.34 X10-1 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	L
COMPOUND NO = 95 MOL WGT -		266.3	NONYL TRIMETHYL AMMONIUM BROMIDE						
1.25 E-2 M	K BR	30	1.43 X10-1 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
2 ENTRIES FOR COMPOUND		30	1.40 X10-1 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
49006			VALUES FRM REF IN CMC			CORR HARK	46004	R	
47006			VALUES FRM REF IN CMC			CORR HARK	46005	R	
25	6.8 X10-2 M	CC	REFRACTIVE INDEX			KLEV	48005	T	L
25	6.5 X10-2 M	XB	UNSPECIFIED CONDUCTANCE			KLEV	53010	T	L
25	7.0 X10-2 W	CC	EQUIV CONDCTNCE GRAPH			SCOT TART	43003	P	L
25	6.46 X10-2 W	BA	SPECFC CONDCTNCE GRAPH			TUDD ALEX	62035	T	2
26	6.43 X10-2 M	BG	VISUAL SPCTR CHNGE SKYB			CORR HARK	47006	T	L
26	6.35 X10-2 M	BG	VISUAL SPCTR CHNGE EOSN			CORR HARK	47006	T	L
26	6.10 X10-2 M	BG	VISUAL SPCTR CHNGE FL			CORR HARK	47006	T	L
26	6.02 X10-2 M	BG	VISUAL SPCTR CHNGE INPX			CORR HARK	47006	T	L
26	6.36 X10-2 M	BG	VISUAL SPCTR CHNGE			CORR HARK	47010	T	L
30	6.3 X10-2 M	CC	TURBIDITY PLT LITE SCATR			TRAP HERM	55009	T	L
40	7.0 X10-2 W	CC	EQUIV CONDCTNCE GRAPH			SCOT TART	43003	P	L
60	7.5 X10-2 M	CC	REFRACTIVE INDEX			KLEV	48005	T	L
60	8.0 X10-2 W	CC	EQUIV CONDCTNCE GRAPH			SCOT TART	43003	P	L
UNK	7.00 X10-2 M	BC	DEBYE PLT LIGHT SCATTER			DEBY	49001	T	L
UNK	6.4 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB			HARK MITT	49006	G	L
UNK	6.86 X10-2 M	BC	METHOD NOT CITED			WASI HUBB	64043	T	L
2.15 E-2 M	BA CL2	26	5.84 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L
5.64 E-2 M	BA CL2	26	5.11 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L
1.00 E-1 M	BA CL2	26	4.54 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L
1.43 E-1 M	BA CL2	26	3.88 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L
2.42 E-1 M	BA CL2	26	3.29 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L
1.9 E-1 M	BUTANOL-1	UNK	4.5 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
3.9 E-1 M	BUTANOL-1	UNK	3.0 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
5.4 E-1 M	BUTANOL-1	UNK	2.1 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
3.8 E-1 M	DIOXANE	UNK	6.5 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
9.6 E-1 M	DIOXANE	UNK	6.7 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
1.96 E 0 M	DIOXANE	UNK	6.9 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
1.25 E-1 M	ETHYLENE GLYCOL	UNK	6.4 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
1.38 E 0 M	ETHYLENE GLYCOL	UNK	6.5 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
2.83 E 0 M	ETHYLENE GLYCOL	UNK	6.6 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
1.21 E 0 M	ETHANOL	UNK	6.4 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
2.29 E 0 M	ETHANOL	UNK	5.6 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
3.08 E 0 M	ETHANOL	UNK	5.5 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
3.63 E 0 M	ETHANOL	UNK	5.7 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
4.08 E 0 M	ETHANOL	UNK	5.8 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
4.62 E 0 M	ETHANOL	UNK	6.0 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
1.25 E-2 M	K BR	30	5.9 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
2.5 E-2 M	K BR	30	5.0 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
5.0 E-2 M	K BR	30	4.5 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L
1.30 E-2 M	K BR	UNK	6.72 X10-2 M	BC	DEBYE PLT LIGHT SCATTER	DEBY	49001	T	L
1.4 E-1 M	METHANOL	UNK	6.3 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
2.9 E-1 M	METHANOL	UNK	6.1 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
3.6 E-1 M	METHANOL	UNK	5.9 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
8.8 E-1 M	METHANOL	UNK	5.9 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
1.88 E 0 M	METHANOL	UNK	6.3 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
2.88 E 0 M	METHANOL	UNK	6.5 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
4.04 E 0 M	METHANOL	UNK	6.8 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
5.86 E 0 M	METHANOL	UNK	8.2 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G	L
7. E-2 M	NA CL	20	6.30 X10-2 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	L
1.0 E-1 M	NA CL	20	5.95 X10-2 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	L
2.5 E-1 M	NA CL	20	4.62 X10-2 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	L
5.0 E-1 M	NA CL	20	3.54 X10-2 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	62004	L	L
4.96 E-2 M	NA CL	26	5.69 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L
1.32 E-1 M	NA CL	26	5.05 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L
2.19 E-1 M	NA CL	26	4.38 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T	L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/
kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
3.22 E-1 M NA CL	26	3.70 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T L	
5.65 E-1 M NA CL	26	3.15 X10-2 M	BG	VISUAL SPCTR CHNGE	CORR HARK	47010	T L	
4.3 E-1 M PROPANOL-1	UNK	5.1 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G L	
9.4 E-1 M PROPANOL-1	UNK	3.9 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G L	
1.46 E 0 M PROPANOL-1	UNK	3.1 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G L	
2.04 E 0 M PROPANOL-1	UNK	2.1 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G L	
2.37 E 0 M PROPANOL-1	UNK	1.9 X10-2 M	CG	VISUAL SPCTR CHNGE SKYB	HARK MITT	49006	G L	
5.00 E 2 Y PRESSURE	25	6.70 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3	
1.000E 3 Y PRESSURE	25	6.70 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3	
1.500E 3 Y PRESSURE	25	6.50 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3	
3.000E 3 Y PRESSURE	25	5.56 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3	
68 ENTRIES FOR COMPOUND								
COMPOUND NO = 96 MOL WGT -	294.3	UNDECYL TRIMETHYL AMMONIUM BROMIDE						
1.25 E-2 M K BR	30	3.6 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L	
2.5 E-2 M K BR	30	3.1 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L	
3 ENTRIES FOR COMPOUND	30	2.7 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L	
COMPOUND NO = 97 MOL WGT -	308.4	DODECYL TRIMETHYL AMMONIUM BROMIDE						
01.0 1.6 X10-2 M	BD	SURFACE TENSION LOG PLOT	SCHI	63026	L L			
UN 1.48 X10-2 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L			
10.0 1.5 X10-2 M	BD	SURFACE TENSION LOG PLOT	SCHI	63026	L L			
20 1.59 X10-2 M	BB	INTERFACIAL TENSION LOGM	HAUD PHIL	58012	L D			
25 4.48 X10-1 D	BB	TURBIDITY PLT LITE SCATR	ANAC JOHN	64017	K D			
1.452X10-2 M					M			
25 1.44 X10-2 N	BA	EQUIV CONDCTNCE GRAPH	VOEK TART	55006	T D			
25 1.64 X10-2 M	CC	REFRACTIVE INDEX	KLEV	48005	T L			
25 1.40 X10-2 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L			
25 1.58 X10-2 W	CB	SPECFC CONDCTNCE GRAPH	SCOT TART	43003	P L			
25.0 1.4 X10-2 M	BD	SURFACE TENSION LOG PLOT	SCHI	63026	L L			
25 1.564X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T D			
25 1.42 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	BRUN HOLT	61016	T D			
30 1.47 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L			
40 1.65 X10-2 W	CB	SPECFC CONDCTNCE GRAPH	SCOT TART	43003	P L			
40.0 1.5 X10-2 M	BD	SURFACE TENSION LOG PLOT	SCHI	63026	L L			
50 1.73 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L			
55.0 1.6 X10-2 M	BD	SURFACE TENSION LOG PLOT	SCHI	63026	L L			
60 1.9 X10-2 M	CD	REFRACTIVE INDEX	KLEV	48005	T L			
60 1.90 X10-2 W	CC	SPECFC CONDCTNCE GRAPH	SCOT TART	43003	P L			
70 1.94 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L			
UNK 4.1 X10-2 D	HG	SURFACE TENSION UNSPEC	WAN	66018	T L			
UNK 1.51 X10-2 M	BC	DEBYE PLT LIGHT SCATTER	DEBY	49001	T L			
25 4.62 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3			
6.25 E-3 M CA BR2	UNK 1.13 X10-2 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L		
1.0 E 0 M H NO3	25 6.0 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
1.0 E 0 M H NO3	25 9.5 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
	3.08 X10-6 M				M			
1.25 E-2 M K BR	30 1.08 X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L		
2.5 E-2 M K BR	30 9.1 X10-3 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L		
3.403E-2 M K BR	30 8.48 X10-3 M	BC	DEBYE PLT LIGHT SCATTER	DEBY	49001	T L		
5.0 E-2 M K BR	30 7.0 X10-3 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L		
3.403E-2 M K BR	45 9.31 X10-3 M	BC	DEBYE PLT LIGHT SCATTER	DEBY	49001	T L		
3.403E-2 M K BR	60 1.201X10-2 M	BC	DEBYE PLT LIGHT SCATTER	DEBY	49001	T L		
1.25 E-2 M K BR	UNK 1.16 X10-2 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L		
1.30 E-2 M K BR	UNK 1.05 X10-2 M	BC	DEBYE PLT LIGHT SCATTER	DEBY	49001	T L		
1.0 E 0 M K CL	25 4.0 X10-4 M	CE	SURFACE TN SN LINEAR PLOT	COLI	50012	T L		
1.0 E 0 M K CL	25 3.2 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
	1.03 X10-5 M				M			
1.0 E 0 M K CL	25 5.3 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
	1.71 X10-5 M				M			
1.0 E 0 M K NO3	25 9.3 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
	3.01 X10-6 M				M			
1.0 E 0 M K NO3	25 3.8 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
	1.23 X10-5 M				M			
1.0 E 0 M K OH	25 3.5 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
	1.13 X10-6 M				M			
1.0 E 0 M K OH	25 3.8 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L		
	1.23 X10-5 M				M			
4.17 E-3 M LA BR3	UNK 1.17 X10-2 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L		
2. E-1 M NA BR	10.0 2.7 X10-3 M	BC	SURFACE TENSION LOG PLOT	SCHI	63026	L L		
1.00 E-1 M NA BR	25 1.38 X10-1 D	BB	TURBIDITY PLT LITE SCATR	ANAC JOHN	64017	K 3		
	4.474X10-3 M				M			

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2.	E-1 M	NA BR	25	2.0 X10-3 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.	E-1 M	NA BR	25.0	2.5 X10-3 M	BC	SURFACE TENSION LOG PLOT	SCHI	63026	L L
4.	E-1 M	NA BR	25	1.0 X10-3 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.02	E-1 M	NA BR	25	6.2 X10-2 D 2.01 X10-3 M	BB	TURBIDITY PLT LITE SCATR	ANAC JOHN	64017	K 3 M
8.	E-1 M	NA BR	25	6.3 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.02	E-1 W	NA BR	31.5	1.9 X10-3 M	BD	DEBYE PLT LIGHT SCATTER	ANAC GHOS	63016	T L
2.	E-1 M	NA BR	40.0	3.0 X10-3 M	BC	SURFACE TENSION LOG PLOT	SCHI	63026	L L
2.	E-1 M	NA BR	55.0	3.2 X10-3 M	BC	SURFACE TENSION LOG PLOT	SCHI	63026	L L
7.	E-2 M	NA CL	20	8.7 X10-3 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
1.0	E-1 M	NA CL	20	7.5 X10-3 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
2.5	E-1 M	NA CL	20	4.25 X10-3 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
5.0	E-1 M	NA CL	20	2.57 X10-3 M	BB	INTERFACIAL TENSION LOGM	HAYD TAYL	60011	L L
5.00	E 2 Y	PRESSURE	25	1.61 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
1.000E 3 Y	PRESSURE		25	1.616X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
1.500E 3 Y	PRESSURE		25	1.56 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
3.000E 3 Y	PRESSURE		25	1.272X10-2 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
5.	E-1 M	UREA	25	1.56 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	BRUN HOLT	61016	T 3
2.0	E 0 M	UREA	25	2.04 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	BRUN HOLT	61016	T 3
6.0	E 0 M	UREA	25	4.54 X10-2 M	BA	EQUIV CONDCTNCE GRAPH	BRUN HOLT	61016	T 3
1.0	E 1 C	0115	25	6.3 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.5	E 1 C	0115	25	2.5 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 1 C	0115	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
1.0	E 1 C	0116	25	6.3 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.5	E 1 C	0116	25	2.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 1 C	0116	25	8.0 X10-5 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
1.0	E 1 C	0325	25	8.0 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.5	E 1 C	0325	25	6.0 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 1 C	0325	25	4.0 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
7.5	E 1 C	0325	25	2.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
1.0	E 1 C	0327	25	2.0 X10-3 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.5	E 1 C	0327	25	2.5 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 1 C	0327	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 0 M	NH3	25	3.0 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
1.	E-1 M	NH4 CL		9.72 X10-6 M				M	
5.00	E 2 Y	PRESSURE	25	5.00 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
1.	E-1 W	PHENOL		5.03 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
1.000E 3 Y	PRESSURE		25	4.88 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
1.	E-1 W	PHENOL		3.88 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	TUDD ALEX	62035	T 3
3.000E 3 Y	PRESSURE		25	3.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
1.0	E 1 C	0327	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.	E-1 M	NA BR	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
1.0	E 1 C	0327	25	4. E-1 M NA BR	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
1.0	E 1 C	0327	25	1.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
8.	E-1 M	NA BR	25	2.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.5	E 1 C	0327	25	1.25 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.	E-1 M	NA BR	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.5	E 1 C	0327	25	4. E-1 M NA BR	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.5	E 1 C	0327	25	1.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
8.	E-1 M	NA BR	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 1 C	0327	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
2.	E-1 M	NA BR	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 1 C	0327	25	1.25 X10-4 M	BC	SURFACE TENSION LOG PLOT	SCHI	66025	L L
5.0	E 1 C	0327	25	1.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
8.	E-1 M	NA BR	25	1.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	SCHI	66025	L L
90	ENTRIES FOR COMPOUND								
COMPOUND NO =	98	MOL WGT -	336.4	TETRADECYL TRIMETHYL AMMONIUM BROMIDE					
1.25	E-2 M	K BR	30	3.51 X10-3 M	BB	SURFACE TENSION LOG PLOT	VENA NAUM	64001	T L
2.5	E-2 M	K BR	30	3.6 X10-3 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L
5.0	E-2 M	K BR	50	4.2 X10-3 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L
1.30	E-2 M	K BR	70	4.2 X10-3 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L
5.	E-2 M	NA BR	UNK	8.2 X10-2 D	HG	SURFACE TENSION UNSPEC	WAN	66018	T L
10	ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; F—wt % solvent; G—mol % surfactant; H—normality; I—counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/l; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Meth.	Method	Authors	Reference	Source	Evaluation	
COMPOUND NO = 99 MOL WGT - 364.5 HEXADECYL TRIMETHYLAMMONIUM BROMIDE										
					QUESTIONABLE CRITERION	DEBY ANAC	51001	R		
					GRAPH DATA NOT RETRIEVED	COHE VASS	61027	R		
		25	8.0 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	66025	L	L	
		25	9.8 X10-4 M	CC	SPECFC CONDCTNCE GRAPH	SCOT TART	43003	P	L	
		25	5. X10-4 W	HG	VISUAL SPCTR CHNGE	FINE MCBA	48011	T	L	
		25	9.20 X10-4 W	BA	SPECFC CONDCTNCE GRAPH	CZER	66030	T	2	
		25	3.3 X10-2 D	HE	METHOD NOT CITED	GINN HARR	61014	T	L	
		25	3.3 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T	L	
		30	4. X10-4 M	CE	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L	
		30	3.0 X10-2 D	BB	REFRACTIVE INDEX	STEI COHE	65012	T	L	
			8.23 X10-4 M					M		
		35	9.5 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	HART COLL	36001	P	D	
		35	9.8 X10-4 M	BB	SPECFC CONDCTNCE GRAPH	HART COLL	36001	P	D	
		35	9. X10-4 M	BD	EQUIV CONDCTNCE GRAPH	HART COLL	36001	T	L	
		35	1.020 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	CZER	66030	T	D	
		35	9.1 X10-4 M	XG	VELOCITY OF SOUND	KUPP SURY	65028	T	L	
		45	1.155 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	CZER	66030	T	3	
		50	1.5 X10-3 M	CD	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L	
		55	1.320 X10-3 W	BA	SPECFC CONDCTNCE GRAPH	CZER	66030	T	3	
		60	1.0 X10-3 M	CD	REFRACTIVE INDEX	KLEV	48005	T	L	
		70	1.5 X10-3 M	CD	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T	L	
		UNK	5.1 X10-2 D	HG	SURFACE TENSION UNSPEC	WAN	66018	T	L	
.	E 0	NH4 BR	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
.	E 0	H BR	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
1.0	E 0 M	H N03	25	7.0 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
			1.92 X10-6 M					M		
1.0	E 0 M	H N03	25	3.8 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	' COLI	50012	T	L
			1.04 X10-5 M					M		
3.	E-3 M	K BR	30	8.2 X10-4 M	BD	TURBIDITY PLT LITE SCATR	TART	59010	T	L
	E 0	K BR	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
1.0	E 0 M	K CL	25	6.0 X10-5 M	CE	SURFACE TNSN LINEAR PLOT	COLI	50012	T	L
1.0	E 0 M	K CL	25	2.0 X10-4 M	CE	SURFACE TENSION MINIMUM	COLI	50012	T	L
1.0	E 0 M	K CL	25	7.4 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
			2.03 X10-6 M					M		
1.0	E 0 M	K CL	25	3.5 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
			9.60 X10-6 M					M		
1.0	E 0 M	K N03	25	7.4 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
1.0	E 0 M	K N03	25	6.5 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
			1.78 X10-6 M					M		
			2.03 X10-6 M					M		
1.0	E 0 M	K N03	25	4.2 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
1.0	E 0 M	K N03	25	2.3 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
			6.31 X10-6 M					M		
			1.15 X10-5 M					M		
1.0	E 0 M	K OH	25	2.1 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
1.0	E 0 M	K OH	25	5.76 X10-7 M				M		
			1.3 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L	
			3.56 X10-6 M					M		
.	E 0	K OH	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
.	E 0	(C4H9)4 N BR /NORMAL	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
.	E 0	(C2H5)4 N BR	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
.	E 0	(CH3) N BR	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
.	E 0	(C3H7)4 N BR /NORMAL	30		QUESTIONABLE CRITERION	STEI COHE	65012	R		
1.0	E 1 C	0117	25	8.0 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
2.5	E 1 C	0117	25	4.5 X10-5 M	CC	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
5.0	E 1 C	0117	25	1.4 X10-5 M	CD	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
.	E 0	0427			GRAPH DATA NOT RETRIEVED	LANG	53005	R		
1.0	E 1 C	0535	25	8.0 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
2.5	E 1 C	0535	25	3.0 X10-5 M	CD	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
5.0	E 1 C	0535	25	1.3 X10-5 M	CD	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
5.0	E 0 M	NH3	25	8.0 X10-5 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
1.	E-1 M	NH4 CL			2.19 X10-6 M			M		
5.0	E 0 M	NH3	25	1.5 X10-4 D	CG	POLAROGRAPHIC MAXIMUM	COLI	50012	T	L
1.	E-1 M	NH4 CL			4.11 X10-6 M			M		
1.00	E 0	PH OF SOLUTION	30	6. X10-5 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T	L
1.50	E-5 M	BROMPHENOL BLUE								
1.00	E 0	PH OF SOLUTION	30	6. X10-5 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T	L
1.50	E-5 M	BROMPHENOL BLUE								
1.00	E 0	PH OF SOLUTION	30	1.2 X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T	L
3.00	E-5 M	BROMPHENOL BLUE								
1.00	E 0	PH OF SOLUTION	30	1.1 X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T	L
3.00	E-5 M	BROMPHENOL BLUE								
1.00	E 0	PH OF SOLUTION	30	1.2 X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T	L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l (or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC		Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.00 E 0	PH OF SOLUTION	30	1.1	X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
3.00 E-5 M	BROMPHENOL BLUE									
3.00 E-5 M	BROMPHENOL BLUE	30	1.2	X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
1.00 E 0	PH OF SOLUTION	30	1.1	X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
3.00 E-5 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	1.1	X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
1.50 E-5 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	1.2	X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
1.50 E-5 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	1.2	X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
1.50 E-5 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	1.1	X10-4 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
1.50 E-5 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	2.5	X10-5 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
3.75 E-6 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	3.5	X10-5 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
3.75 E-6 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	6.8	X10-5 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
7.50 E-6 M	BROMPHENOL BLUE									
6.50 E 0	PH OF SOLUTION	30	5.0	X10-5 M	CG	FOTOMTR SPCTR CHNGE BRPB	COLI	51008	T L	
7.50 E-6 M	BROMPHENOL BLUE									
66 ENTRIES FOR COMPOUND										
COMPOUND NO = 100 MOL WGT -		272.2	OCTYL PYRIDINIUM BROMIDE							
20	2.3	X10-1 W	CC	SURFACE TENSION LOG PLOT		BURY BROW	52011	T L		
30	1.93	X10-1 M	CC	TURBIDITY PLT LITE SCATR		TRAP HERM	55009	T L		
5.0 E-2 M K BR		30	1.89	X10-1 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L	
3 ENTRIES FOR COMPOUND										
COMPOUND NO = 101 MOL WGT -		314.3	UNDECYL PYRIDINIUM BROMIDE							
30	4.2	X10-2 M	CC	TURBIDITY PLT LITE SCATR		TRAP HERM	55009	T L		
5.0 E-2 M K BR		30	3.1	X10-2 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L	
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 102 MOL WGT -		356.4	TETRADECYL PYRIDINIUM BROMIDE							
14.0	3.2	X10-3 M	BC	KRAFFT POINT SOLUBILITY		ADDI FURM	56019	T L		
18.5	3.1	X10-3 M	BC	INTERFACIAL TNSN UNSPEC		ADDI FURM	56019	T L		
25	2.9	X10-3 M	CC	SPECFC CONDCTNCE GRAPH		BENT SPAR	66038	T L		
30	2.57	X10-3 M	BB	SURFACE TENSION LOG PLOT		VENA NAUM	64001	T L		
30	4.1	X10-3 M	CC	TURBIDITY PLT LITE SCATR		TRAP HERM	55009	T L		
2.5 E-2 M K BR		30	2.0	X10-3 M	CC	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L	
5.0 E-2 M K BR		30	1.5	X10-3 M	CD	TURBIDITY PLT LITE SCATR	TRAP HERM	55009	T L	
5. E-2 M NA BR		30	2.4	X10-4 M	BC	SURFACE TENSION LOG PLOT	VENA NAUM	64001	TA L	
8 ENTRIES FOR COMPOUND										
COMPOUND NO = 103 MOL WGT -		234.3	HEXYL /OXYETHYLENE/ 3 ALCOHOL							
HOMOGENEOUS HEAD GROUP										
15	1.07	X10-3 M	BB	SURFACE TENSION LOG PLOT		CORK GOOD	64003	T 3		
20	7.5	X10-2 M	BE	SURFACE TENSION LOG PLOT		MULL METC	62015	T L		
25	1.00	X10-1 M	BB	SURFACE TENSION LOG PLOT		CORK GOOD	64003	T 3		
		35	7.8	X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
4 ENTRIES FOR COMPOUND										
COMPOUND NO = 104 MOL WGT -		262.4	OCTYL /OXYETHYLENE/ 3 ALCOHOL							
HOMOGENEOUS HEAD GROUP										
15	9.3	X10-3 M	BB	SURFACE TENSION LOG PLOT		CORK GOOD	64003	T 3		
		25	7.5	X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 105 MOL WGT -		394.6	OCTYL /OXYETHYLENE/ 6 ALCOHOL							
HOMOGENEOUS HEAD GROUP										
15	1.19	X10-2 M	BB	SURFACE TENSION LOG PLOT		CORK GOOD	64003	T 3		
18	7.6	X10-3 M	BC	UNSPEC LIGHT SCATTER		GOOD OTTE	61004	T L		
18	4.45	X10-1 D	BB	SURFACE TENSION LOG PLOT		CORK GOOD	64023	T L		
		1.127X10-2 M								
20	3.8	X10-1 D	BD	DENSITY		FLOR	66020	T L		
		9.63	X10-3 M							
25	9.9	X10-3 M	BB	SURFACE TENSION LOG PLOT		CORK GOOD	64003	T 3		
25	9.8	X10-3 M	BB	SURFACE TENSION LOG PLOT		GOOD OTTE	61004	T L		

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
	30	3.50 X10-1 D 8.869X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64023	T L M	
	35	7.7 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	40	2.85 X10-1 D 7.222X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64023	T L M	
	45	6.7 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T L	
10 ENTRIES FOR COMPOUND								
COMPOUND NO = 106 MOL WGT - HOMOGENEOUS HEAD GROUP	526.7	OCTYL /OXYETHYLENE/ 9 ALCOHOL						
	15	1.6 X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	25	1.3 X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	35	1.1 X10-2 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 107 MOL WGT - HOMOGENEOUS HEAD GROUP	290.4	DECYL /OXYETHYLENE/ 3 ALCOHOL						
	15	7.3 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	25	6.0 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	35	5.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 108 MOL WGT - HOMOGENEOUS HEAD GROUP	422.6	DECYL /OXYETHYLENE/ 6 ALCOHOL						
	15	1.14 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	20	9.2 X10-4 M	BD	SURFACE TENSION LOG PLOT	CARL CHAL	64009	T L	
	20	9.6 X10-4 M	BC	REFRACTIVE INDEX	DONB JAN	63021	T L	
	23.5	9.5 X10-4 M	BG	FOTOMTR SPCTR CHNG I2	CARL CHAL	64009	T L	
	25	9.0 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	25	3.80 X10-2 D 8.991X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64023	T L M	
	35	6.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	35	2.79 X10-2 D 6.601X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64023	T L M	
	45	6.4 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T L	
	45	1.83 X10-2 D 4.330X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64023	T P M	
10 ENTRIES FOR COMPOUND								
COMPOUND NO = 109 MOL WGT - HOMOGENEOUS HEAD GROUP	554.8	DECYL /OXYETHYLENE/ 9 ALCOHOL						
	15	1.4 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	25	1.3 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	35	1.1 X10-3 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 110 MOL WGT - HOMOGENEOUS HEAD GROUP	450.7	DODECYL /OXYETHYLENE/ 6 ALCOHOL						
	15	1.08 X10-4 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	15	1.08 X10-4 M	BB	SURFACE TENSION LOG PLOT	GOOD OTTE	61004	T L	
	20	8.2 X10-5 M	BC	SURFACE TENSION LOG PLOT	CARL CHAL	64009	T L	
	20	7.2 X10-5 M	BG	FOTOMTR SPCTR CHNG I2	CARL CHAL	64009	T L	
	20	1.00 X10-4 M	BC	REFRACTIVE INDEX	DONB JAN	63021	T L	
	25	8.7 X10-5 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	25	8.7 X10-5 M	BB	SURFACE TENSION LOG PLOT	GOOD OTTE	61004	T L	
	35	7.2 X10-5 M	BB	SURFACE TENSION LOG PLOT	CORK GOOD	64003	T 3	
	35	7.2 X10-5 M	BB	SURFACE TENSION LOG PLOT	GOOD OTTE	61004	T L	
9 ENTRIES FOR COMPOUND								
COMPOUND NO = 111 MOL WGT -	272.3	LITHIUM DODECYL 1 SULFATE						
6.75 E-2 M NA CL	10	7.0 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
1. E-5 M RHODAMINE 6GPC	25	7.9 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
5. E-6 M RHODAMINE 6GPC	25	8.77 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2	
	25	8.93 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2	
	40	1.05 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	MEGU KOND	56020	T L	
	45	8.0 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
	50	6.9 X10-3 M	DG	VISUAL SPCTR CHNGE PNCC	RAIS	52016	T L	
	40	2.5 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	56020	T L	
	40	9.7 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	56020	T L	
	40	9.6 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	56020	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; F—wt % solvent; G—mol % surfactant; H—normality; I—mol % surfactant mixture; J—molar; K—normal; L—wt %; M—wt % surfactant; N—wt % counterions; O—varied; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l (or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
3. E O M UREA	10	8.0 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	10	1.0 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
1. E O M UREA	25	8.0 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
2. E O M UREA	25	8.2 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	25	8.8 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
4.5 E O M UREA	25	9.4 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	25	1.0 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	45	8.9 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	45	1.05 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
19 ENTRIES FOR COMPOUND								
COMPOUND NO = 112 MOL WGT -	339.5	TETRAMETHYL AMMONIUM DODECYL 1 SULFATE						
3. E O M UREA	10	4.8 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	25	5.50 X10-3 M	DA	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L	
1. E O M UREA	25	4.8 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
2. E O M UREA	25	5.41 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2	
3. E O M UREA	25	5.52 X10-3 M	BA	SPECFC CONDCTNCE GRAPH	MYSE PRIN	59002	TL 2	
4.5 E O M UREA	30	5.7 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	MEGU KOND	59026	T L	
6. E O M UREA	30	4.5 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	59026	T L	
3. E O M UREA	45	6.3 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	10	6.6 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
1. E O M UREA	10	1.0 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
2. E O M UREA	25	5.0 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	25	5.8 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
4.5 E O M UREA	25	6.9 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	25	7.9 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	45	1.0 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	45	7.9 X10-3 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
17 ENTRIES FOR COMPOUND	45	1.08 X10-2 M	BB	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
COMPOUND NO = 113 MOL WGT -	420.5	SODIUM DODECYL TRI-OXYETHYLENE SULFATE						
3. E O M UREA	01.0	1.25 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
6. E O M UREA	10	1.1 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
1. E O M UREA	25	1.0 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
2. E O M UREA	45	1.2 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	50	1.97 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G P	
6. E O M UREA	55.0	1.4 X10-4 M	CD	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
3. E O M UREA	10	1.55 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	10	2.85 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	25	1.5 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	25	2.5 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	45	1.93 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	45	2.70 X10-4 M	CC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
12 ENTRIES FOR COMPOUND								
COMPOUND NO = 114 MOL WGT -	1,059.4	SODIUM DODECYL /OXYETHYLENE/17.5 SULFATE						
3. E O M UREA	01.0	7.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
6. E O M UREA	10	6.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	25	6.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	45	4.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	55.0	4.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
6. E O M UREA	10	1.3 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	10	2.48 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	25	1.2 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	25	2.2 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
3. E O M UREA	45	7.02 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
6. E O M UREA	45	1.41 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
11 ENTRIES FOR COMPOUND								
COMPOUND NO = 115 MOL WGT -	494.7	DODECYL /OXYETHYLENE/7 ALCOHOL REDUCED POLYDISPERSION OF HEAD GROUPS						
	62019	VALUES FRM REF IN CMC						
	01.0	1.00 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
	05.0	9.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
	05.0	9.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
	10.0	8.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L L	
	10	8.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
	25	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	66025	L L	
	25.0	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
	25	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T L	
	25.0	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/l; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
3. E O M UREA	40.0	3.2 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
6. E O M UREA	40.0	62019		VALUES FRM REF IN CMC	SCHI	63026		R
3. E O M UREA	45	2.8 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6. E O M UREA	45.0	2.8 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
3. E O M UREA	55.0	2.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T	L
6. E O M UREA	10	1.2 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6. E O M UREA	10	2.08 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3. E O M UREA	25	6.25 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6. E O M UREA	25	1.25 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3. E O M UREA	45	3.4 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6. E O M UREA	45	5.6 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
2.5 E 1 I 0001	05.0	9.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
5.0 E 1 I 0001	05.0	9.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
7.5 E 1 I 0001	05.0	1.5 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
9.0 E 1 I 0001	05.0	3.0 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
2.5 E 1 I 0001	25.0	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
5.0 E 1 I 0001	25.0	5.1 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
7.5 E 1 I 0001	25.0	7.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
9.0 E 1 I 0001	25.0	2.0 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
2.5 E 1 I 0001	45.0	2.8 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
5.0 E 1 I 0001	45.0	3.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
7.5 E 1 I 0001	45.0	4.2 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
9.0 E 1 I 0001	45.0	1.3 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
0097	25			SEE CMPD NMBR IN ADDITV	SCHI	66025		X
34 ENTRIES FOR COMPOUND								
COMPOUND NO = 116 MOL WGT - 1,508.1 DODECYL /OXYETHYLENE/30 ALCOHOL								
REDUCED POLYDISPERSION OF HEAD GROUPS								
64020				VALUES FRM REF IN CMC	SCHI	63026		R
62019				VALUES FRM REF IN CMC	SCHI	63026		R
62019				VALUES FRM REF IN CMC	SCHI ATLA	62020		R
01.0	1.20	X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
01.0	1.20	X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
05.0	1.00	X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
05.0	1.00	X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
05.0	1.0	X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
10.0	9.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
10	9.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
25	8.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
25.0	8.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T	L
25	8.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
25.0	8.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
40.0	5.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
40.0	5.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
45	4.8	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
45.0	4.8	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
55.0	4.0	X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T	L
8.6 E-1 M LI CL	25	4.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
4.3 E-1 M NA CL	01.0	9.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CL	01.0	7.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
4.3 E-1 M NA CL	05.0	7.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CL	05.0	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
4.3 E-1 M NA CL	10.0	5.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CL	10.0	3.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
4.3 E-1 M NA CL	25	3.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
8.6 E-1 M NA CL	25	2.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
4.3 E-1 M NA CL	40.0	2.0 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CL	40.0	1.2 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
4.3 E-1 M NA CL	55.0	1.3 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CL	55.0	8. X10-6 M	EE	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 N NA2 SO4	01.0	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 N NA2 SO4	05.0	4.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 N NA2 SO4	10.0	2.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
4.3 E-1 M NA2 SO4	25	1.2 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
8.6 E-1 N NA2 SO4	40.0	6. X10-6 M	EE	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 N NA2 SO4	55.0	4. X10-6 M	EE	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CNS	01.0	9.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CNS	05.0	8.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CNS	10.0	6.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CNS	25	5.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
8.6 E-1 M NA CNS	40.0	4.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M NA CNS	55.0	3.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
8.6 E-1 M (CH3)4 N CL	25	3.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3. E O M UREA	10	3.6 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6. E O M UREA	10	7.2 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
3.	E O M	UREA	25	1.6 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA	25	2.5 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA	45	6.7 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA	45	9.45 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
2.5	E O C	0001	05.0	1.0 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
5.0	E O C	0001	05.0	1.6 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
7.5	E O C	0001	05.0	2.5 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
9.0	E O C	0001	05.0	7.0 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
2.5	E O C	0001	25.0	8.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
5.0	E O C	0001	25.0	9.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
7.5	E O C	0001	25.0	1.6 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
9.0	E O C	0001	25.0	4.6 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
2.5	E O C	0001	45.0	4.8 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
5.0	E O C	0001	45.0	6.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
7.5	E O C	0001	45.0	9.6 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L	L
9.0	E O C	0001	45.0	2.4 X10-4 M	ED	SURFACE TENSION LOG PLOT SEE CMPD NMBR IN ADDITV	SCHI MANN	66001	L	L
		0097		25			SCHI	66025	X	
8.6	E-1 M	LI CL	25	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA								
8.6	E-1 M	LI CL	25	8.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA								
4.3	E-1 M	NA CL	25	4.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA								
4.3	E-1 M	NA CL	25	1.0 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA								
8.6	E-1 M	NA CL	25	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA								
8.6	E-1 M	NA CL	25	1.2 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA								
4.3	E-1 M	NA2 SO4	25	2.9 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA								
4.3	E-1 M	NA2 SO4	25	3.6 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA								
8.6	E-1 M	NA CNS	25	8.7 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA								
8.6	E-1 M	NA CNS	25	1.1 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA								
8.6	E-1 M	(CH3)4 N CL	25	5.7 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA								
8.6	E-1 M	(CH3)4 N CL	25	1.0 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA								
76	ENTRIES FOR COMPOUND									
COMPOUND NO =	117	MOL WGT -	1,564.2	HEXADECYL /OXYETHYLENE/30 ALCOHOL						
		REDUCED POLYDISPERSION OF HEAD GROUPS								
			01.0	3.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
			05.0	2.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
			10.0	2.0 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
			10	2.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
			25	1.4 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	66025	L	L
			25.0	1.1 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
			25	1.1 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
			40.0	6. X10-6 M	EE	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
			45	5.0 X10-6 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
			55.0	4. X10-6 M	EE	SURFACE TENSION LOG PLOT	SCHI	63026	L	L
3.	E O M	UREA	10	3.2 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA	10	4.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA	25	1.6 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA	25	2.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
3.	E O M	UREA	45	6.35 X10-6 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
6.	E O M	UREA	45	7.9 X10-6 M	EC	SURFACE TENSION LOG PLOT	SCHI	64020	T	L
		0099	25			SEE CMPD NMBR IN ADDITV	SCHI	66025	X	
17	ENTRIES FOR COMPOUND									
COMPOUND NO =	118	MOL WGT -	215.3	OCTYL N BETAINE						
			21	2.17 X10-1 M	BC	TURBIDITY PLT LITE SCATR	TORI NAKA	63009	T	L
			23	1.7 X10-1 M	CD	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T	L
			23	1.7 X10-1 M	CD	FOTOMTR SPCTR CHNGE BRPB	BECK WOOD	63015	T	L
			27	2.50 X10-1 M	BB	SURFACE TENSION LOG PLOT	TORI NAKA	63009	T	L
			27	2.50 X10-1 M	BC	DENSITY	TORI NAKA	63009	T	L
5	ENTRIES FOR COMPOUND									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 119 MOL WGT -	243.4	DECYL N BETAINE					
20	1.68 X10-2 M	BB	SURFACE TENSION LOG PLOT	MOLY RHOD	65003	GL L	
20.2	2.1 X10-2 M	CD	REFRACTIVE INDEX	BECK WOOD	63015	T L	
23	1.8 X10-2 M	CD	FOTOMTR SPCTR CHNGE BRPB	BECK WOOD	63015	T L	
23	2.0 X10-2 M	CD	FOTOMTR SPCTR CHNG I2	BECK WOOD	63015	T L	
23	1.8 X10-2 M	CD	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T L	
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 120 MOL WGT -	257.4	UNDECYL N BETAINE					
20	7.1 X10-3 M	BB	SURFACE TENSION LOG PLOT	MOLY RHOD	65003	GL L	
20.2	6.4 X10-3 M	CC	REFRACTIVE INDEX	BECK WOOD	63015	T L	
23	6.0 X10-3 M	CC	FOTOMTR SPCTR CHNGE BRPB	BECK WOOD	63015	T L	
23	6.4 X10-3 M	CC	FOTOMTR SPCTR CHNG I2	BECK WOOD	63015	T L	
23	6.6 X10-3 M	CC	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T L	
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 121 MOL WGT -	271.4	DODECYL N BETAINE					
10.2	2.0 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
20.2	2.1 X10-3 M	CD	REFRACTIVE INDEX	BECK WOOD	63015	T L	
20	2.00 X10-3 M	BB	SURFACE TENSION LOG PLOT	MOLY RHOD	65003	GL L	
20.2	2.1 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
23	1.8 X10-3 M	CD	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T L	
23	1.6 X10-3 M	CD	FOTOMTR SPCTR CHNGE BRPB	BECK WOOD	63015	T L	
23	1.8 X10-3 M	CD	FOTOMTR SPCTR CHNG I2	BECK WOOD	63015	T L	
25.2	2.2 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
30.0	2.3 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
35.7	2.4 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
45.1	2.6 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
49.7	2.7 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
57.0	2.8 X10-3 M	BC	REFRACTIVE INDEX	MOLY RHOD	65003	GL L	
13 ENTRIES FOR COMPOUND							
COMPOUND NO = 122 MOL WGT -	299.5	TETRADECYL N BETAINE					
20	1.78 X10-4 M	BB	SURFACE TENSION LOG PLOT	MOLY RHOD	65003	GL L	
20.2	1.7 X10-4 M	CD	REFRACTIVE INDEX	BECK WOOD	63015	T L	
23	1.8 X10-4 M	CD	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T L	
23	1.5 X10-4 M	CD	FOTOMTR SPCTR CHNGE BRPB	BECK WOOD	63015	T L	
23	1.5 X10-4 M	CD	FOTOMTR SPCTR CHNG I2	BECK WOOD	63015	T L	
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 123 MOL WGT -	327.6	HEXADECYL N BETAINE					
20.2	1.6 X10-5 M	CD	REFRACTIVE INDEX	BECK WOOD	63015	T L	
20	2.52 X10-5 M	BB	SURFACE TENSION LOG PLOT	MOLY RHOD	65003	GL L	
23	2.0 X10-5 M	CD	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T L	
23	1.8 X10-5 M	CD	FOTOMTR SPCTR CHNGE BRPB	BECK WOOD	63015	T L	
23	1.8 X10-5 M	CD	FOTOMTR SPCTR CHNG I2	BECK WOOD	63015	T L	
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 124 MOL WGT -	307.9	DODECYL N BETAINE HYDROCHLORIDE					
23	1.98 X10-3 M	CC	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 125 MOL WGT -	336.0	TETRADECYL N BETAINE HYDROCHLORIDE					
23	1.96 X10-3 M	CC	SURFACE TENSION LOG PLOT	BECK WOOD	63015	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 126 MOL WGT -	355.3	DODECYL TRIMETHYLAMMONIUM IODIDE	QUESTIONABLE CRITERION		BECK WOOD	63015	R
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 127 MOL WGT -	403.3	DODECYL TRIMETHYL AMMONIUM IODATE					
4.95 E-1 W NA IO3	31.5	5.1 X10-3 M	BC	TURBIDITY PLT LITE SCATR	ANAC GHOS	63016	T L
5.1 E-3 M NA BR							
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 128 MOL WGT -	273.5	DODECYL TRIMETHYL AMMONIUM FORMATE					
4.94 E-1 W NA HCO2 FORMATE	31.5	6.0 X10-3 M	BC	DEBYE PLT LIGHT SCATTER	ANAC GHOS	63016	T L
6.0 E-3 M NA BR							
1 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp, °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
5. E-1 M	NA CL	60	1.20 X10-3 M	BB	SURFACE TENSION LOG PLOT	TORI NAKA	63008	T L
5. E-1 M	NA CL	27	7.5 X10-4 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
5. E-1 M	NA CL	7	9.6 X10-4 M	BB	SURFACE TENSION LOG PLOT	TORI NAKA	63008	T L
1.0 E O M	NA CL	27	6.8 X10-4 M	BB	SURFACE TENSION LOG PLOT	TORI NAKA	63008	T L
1.0 E O M	NA CL	27	5.3 X10-4 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
1.5 E O M	NA CL	27	4.8 X10-4 M	BB	SURFACE TENSION LOG PLOT	TORI NAKA	63008	T L
2.0 E O M	NA CL	27	3.4 X10-4 M	BB	SURFACE TENSION LOG PLOT	TORI NAKA	63008	T L
14 ENTRIES FOR COMPOUND								
COMPOUND NO = 135 MOL WGT -		265.8	OCTYL C	BETAINE HYDROCHLORIDE				
1. E-1 M	NA CL	27	6.0 X10-2 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
2 ENTRIES FOR COMPOUND		27	4.2 X10-2 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
COMPOUND NO = 136 MOL WGT -		237.3	SODIUM ALPHA DIMETHYL AMINO CAPRATE					
1. E-1 M	NA CL	27	9.6 X10-2 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
2 ENTRIES FOR COMPOUND		27	7.0 X10-2 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
COMPOUND NO = 137 MOL WGT -		251.8	ALPHA DIMETHYLAMINO CAPRIC ACID HYDROCHLORIDE					
1. E-1 M	NA CL	27	8.0 X10-2 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
2 ENTRIES FOR COMPOUND		27	4.2 X10-2 M	BC	FOTOMTR SOLUBLZTN SDN 4	TORI NAKA	63008	T L
COMPOUND NO = 138 MOL WGT -		306.3	SODIUM NONYL BENZENE SULFONATE					
BRANCHED HYDROCARBON CHAIN								
		25	1.57 X10-2 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L
		50	1.71 X10-2 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L
		75	2.30 X10-2 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 139 MOL WGT -		348.4	SODIUM DODECYL BENZENE SULFONATE					
BRANCHED HYDROCARBON CHAIN								
		25	4.90 X10 3 M	HC	QUESTIONABLE CRITERION	YANG FOST	53015	R
		25	4.80 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
		25	5.0 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
		25	1.53 X10-1 D	HC	SPECFC CONDCTNCE GRAPH	GINN HARR	60010	G L
		25	1.23 X10-1 D	HC	SURFACE TENSION UNSPEC	GINN HARR	58008	T L
		25	7. X10-2 D	HG	VISUAL SPCTR CHNGE PNCCN	GINN HARR	58008	T L
		25	4.1 X10-2 D	HG	FOTOMTR SPCTR CHNGE PNCCN	GINN HARR	58008	T L
		25	1.17 X10-1 D	HE	METHOD NOT CITED	GINN HARR	61014	T L
		25	1.32 X10-1 D	HG	VISUAL SPCTR CHNGE PNCCN	GINN HARR	58008	T L
		25	1.10 X10-1 D	HG	FOTOMTR SPCTR CHNGE PNCCN	GINN HARR	58008	T L
		25	1.32 X10-1 D	HC	SPECFC CONDCTNCE GRAPH	GINN HARR	58008	T L
		25	1.15 X10-1 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
		26	1.23 X10-1 D	HB	SURFACE TENSION LOG PLOT	MANK	64010	T L
		26	3.53 X10-3 M	HB	SURFACE TENSION LOG PLOT	MANK	66021	T L
		35	1.3 X10-1 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
		50	7.5 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L
		60	1.4 X10-1 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
		60	7.0 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
		75	8.0 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L
		75	1.75 X10-1 D	HE	METHOD NOT CITED	GINN HARR	61014	T L
2.1 E O Q	C12 DIETHANOLAMIDE	25	5.2 X10-2 D	HG	FOTOMTR SPCTR CHNGE PNCCN	GINN HARR	58008	T L
2.1 E O Q	C12 DIETHANOLAMIDE	25	6.7 X10-2 D	HC	SPECFC CONDCTNCE GRAPH	GINN HARR	58008	T L
2.1 E O Q	C12 DIETHANOLAMIDE	25	2.1 X10-2 D	HC	SURFACE TENSION UNSPEC	GINN HARR	58008	T L
1. E 2 I	NA CL	25	4.50 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
4. E 2 I	NA CL	25	3.63 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
1. E 2 I	NA2 CO3	25	4.00 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
4. E 2 I	NA2 CO3	25	2.15 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
1. E 2 I	NA OH	25	4.10 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
4. E 2 I	NA OH	25	3.00 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
1. E 2 I	NA4 P207 PYRO	25	3.46 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
4. E 2 I	NA4 P207 PYRO	25	2.14 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
1. E 2 I	NA PO4	25	3.65 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
4. E 2 I	NA PO4	25	2.12 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
1. E 2 I	NA14 P12037 POLY	25	2.88 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
4. E 2 I	NA14 P12037 POLY	25	1.55 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
1. E 2 I	NA53 P500154 POLY	25	1.90 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
1. E 2 I	NA2 SI03 META	25	4.00 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives			Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
4. E 2 I NA2 SI03 META			25	2.16 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	59009	T L	
2.1 E 0 Q NA5 P3010 TRIPOLY			25	4. X10-2 D	HG	FOTOMTR SPCTR CHNGE PNCC	GINN HARR	58008	T L	
2.1 E 0 Q NA5 P3010 TRIPOLY			25	1.10 X10-1 D	HC	SURFACE TENSION UNSPEC	GINN HARR	58008	T L	
1. E 2 I NA5 P3010 TRIPOLY			25	3.33 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	59009	T L	
4. E 2 I NA5 P3010 TRIPOLY			25	2.20 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	59009	T L	
2. E 2 I NA5 P3010 TRIPOLY			50	7.0 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
3.33 E 1 Q 0206			82	1.8 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	MANK	64010	T L	
5.0 E 1 Q 0206			82	7.5 X10-3 D	HC	FOTOMTR SOLUBLZTN OROT	MANK	64010	T L	
7.5 E 1 Q 0206			82	5. X10-3 D	HC	FOTOMTR SOLUBLZTN OROT	MANK	64010	T L	
47 ENTRIES FOR COMPOUND										
COMPOUND NO = 140 MOL WGT - 320.4 SODIUM DECYL BENZENE SULFONATE										
BRANCHED HYDROCARBON CHAIN										
25 4.1 X10-3 M			25	4.1 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
50 4.8 X10-3 M			50	4.8 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
75 6.3 X10-3 M			75	6.3 X10-3 M	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
3 ENTRIES FOR COMPOUND										
COMPOUND NO = 141 MOL WGT - 362.4 SODIUM TRIDECYL BENZENE SULFONATE										
BRANCHED HYDROCARBON CHAIN										
25 1.3 X10-3 M			25	1.3 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
25 5. X10-2 D			25	5. X10-2 D	HE	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
50 2.8 X10-3 M			50	2.8 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
50 1.0 X10-1 D			50	1.0 X10-1 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
75 2.8 X10-3 M			75	2.8 X10-3 M	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
75 1.0 X10-1 D			75	1.0 X10-1 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
2. E 2 I NA5 P3010 TRIPOLY			50	3.1 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
7 ENTRIES FOR COMPOUND										
COMPOUND NO = 142 MOL WGT - 390.5 SODIUM PENTADECYL BENZENE SULFONATE										
BRANCHED HYDROCARBON CHAIN										
25 4.4 X10-4 M			25	4.4 X10-4 M	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
25 1.7 X10-2 D			25	1.7 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
50 6.5 X10-4 M			50	6.5 X10-4 M	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
50 2.1 X10-2 D			50	2.1 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
75 9.4 X10-4 M			75	9.4 X10-4 M	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	G L	
75 3.5 X10-2 D			75	3.5 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
2. E 2 I NA5 P3010 TRIPOLY			50	1. X10-2 D	HE	FOTOMTR SOLUBLZTN OROT	GINN KINN	61015	T L	
7 ENTRIES FOR COMPOUND										
COMPOUND NO = 143 MOL WGT - 378.6 DECYL /OXYETHYLENE/ 5.0 ALCOHOL										
BRANCHED CHAIN, NATURAL OE DISTRIBUTION										
25 6.3 X10-2 D			25	6.3 X10-2 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
50 6.0 X10-2 D			50	6.0 X10-2 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
75 4.0 X10-2 D			75	4.0 X10-2 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
3 ENTRIES FOR COMPOUND										
COMPOUND NO = 144 MOL WGT - 598.9 DECYL /OXYETHYLENE/ 10.0 ALCOHOL										
BRANCHED CHAIN, NATURAL OE DISTRIBUTION										
25 1.6 X10-1 D			25	1.6 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
35 1.0 X10-1 D			35	1.0 X10-1 D	HE	METHOD NOT CITED	GINN HARR	61014	T L	
50 9.6 X10-2 D			50	9.6 X10-2 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
75 8.5 X10-2 D			75	8.5 X10-2 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
75 8.5 X10-2 D			75	8.5 X10-2 D	HE	METHOD NOT CITED	GINN HARR	61014	T L	
5 ENTRIES FOR COMPOUND										
COMPOUND NO = 145 MOL WGT - 814.8 DECYL /OXYETHYLENE/ 14.9 ALCOHOL										
BRANCHED CHAIN, NATURAL OE DISTRIBUTION										
25 2.8 X10-1 D			25	2.8 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
50 1.44 X10-1 D			50	1.44 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
75 1.2 X10-1 D			75	1.2 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	
75 1.2 X10-1 D			75	1.2 X10-1 D	HE	METHOD NOT CITED	GINN HARR	61014	T L	
4 ENTRIES FOR COMPOUND										
COMPOUND NO = 146 MOL WGT - 1,039.5 DECYL /OXYETHYLENE/ 20.0 ALCOHOL										
BRANCHED CHAIN, NATURAL OE DISTRIBUTION										
25 6.0 X10-1 D			25	6.0 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L	

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; F – wt % solvent; G – mol % surfactant; H – normality; I – mol % surfactant mixture; J – wt % surfactant; K – varied; L – mol/kg; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/l or kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
	50	2.9 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	1.9 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 147 MOL WGT - 1,453.6 DECYL /OXYETHYLENE/ 29.4 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	8.3 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	4.5 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	3.4 X10-1 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 148 MOL WGT - 429.5 TRIDECYL /OXYETHYLENE/ 5.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	5.8 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	5.0 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	6.5 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 149 MOL WGT - 645.4 TRIDECYL /OXYETHYLENE/ 10.1 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	9. X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
	25	1.0 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	25	9.2 X10-3 D	HE	METHOD NOT CITED	GINN HARR	61014	T L
	25	9.2 X10-3 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
	50	7.8 X10-3 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	7.8 X10-3 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
	75	7.6 X10-3 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	7.6 X10-3 D	HE	METHOD NOT CITED	GINN HARR	61014	T L
	75	7.57 X10-3 D	HC	FOTOMTR SOLUBLZTN OROT	GINN KENN	61015	T L
4. E 2 I NA4 P207 PYRO	25	9. X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN KENN	59009	T L
10 ENTRIES FOR COMPOUND							
COMPOUND NO = 150 MOL WGT - 870.1 TRIDECYL /OXYETHYLENE/ 15.2 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	2.3 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	1.5 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	1.0 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 151 MOL WGT - 1,081.6 TRIDECYL /OXYETHYLENE/ 20.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	3.0 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	1.9 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	1.8 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 152 MOL WGT - 1,548.6 TRIDECYL /OXYETHYLENE/ 30.6 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	8.0 X10-2 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	4.6 X10-2 D	HB	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	2.5 X10-2 D	HC	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 153 MOL WGT - 440.6 NONYL BENZENE /OXYETHYLENE/ 5.0 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	2.5 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	1.0 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	1.9 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 154 MOL WGT - 652.1 NONYL BENZENE /OXYETHYLENE/ 9.8 ALCOHOL BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	4.0 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	50	4.3 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
	75	4.2 X10-3 D	HD	FOTOMTR SOLUBLZTN OROT	GINN HARR	60010	T L
3 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 155 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 50 75	898.9 X10-3 D X10-3 D X10-3 D	HD FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT	GINN HARR GINN HARR GINN HARR	60010 60010 60010	T L T L T L	
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 156 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 25 50 75 8.6 E-1 M NA CL	1,101.5 X10-2 D X10-4 M X10-2 D X10-2 D 25	HC FOTOMTR SOLUBLZTN OROT SURFACE TENSION LOG PLOT FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT 7.8 X10-5 M	GINN HARR HSIA DUNN GINN HARR GINN HARR GINN HARR HSIA DUNN	60010 56014 60010 60010 60010 -56014	T L T L T L T L T L T L	
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 157 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 50 75	1,551.0 X10-2 D X10-2 D X10-2 D	HC FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT	GINN HARR GINN HARR GINN HARR	60010 60010 60010	T L T L T L	
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 158 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 50 75	482.7 X10-4 D X10-4 D X10-4 D	HD FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT	GINN HARR GINN HARR GINN HARR	60010 60010 60010	T L T L T L	
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 159 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 25 50 75	711.8 X10-3 D X10-3 D X10-4 D X10-4 D	HD FOTOMTR SOLUBLZTN OROT METHOD NOT CITED FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT HE METHOD NOT CITED	GINN HARR GINN HARR GINN HARR GINN HARR GINN HARR	60010 61014 60010 60010 61014	T L T L T L T L T L	
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 160 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 50 75	927.7 X10-3 D X10-3 D X10-3 D	HD FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT	GINN HARR GINN HARR GINN HARR	60010 60010 60010	T L T L T L	
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 161 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 50 75	1,148.0 X10-3 D X10-3 D X10-3 D	HD FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT	GINN HARR GINN HARR GINN HARR	60010 60010 60010	T L T L T L	
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 162 MOL WGT – BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25 50 75	1,610.7 X10-3 D X10-3 D X10-3 D	HD FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT FOTOMTR SOLUBLZTN OROT	GINN HARR GINN HARR GINN HARR	60010 60010 60010	T L T L T L	
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 163 MOL WGT – NATURAL DISTRIBUTION OF HEAD GROUPS	UNK UNK UNK	538.8 X10-3 D X10-3 D X10-3 D	HC TURBIDITY PLT LITE SCATR FOTOMTR SPCTR CHNG I2 TURBIDITY PLT LITE SCATR	BECH BECH BECH	59006 59006 61003	T L G L T L	
E O CA CL2 E O NA CITRATE		25 25	SUMMARIZING EQN ONLY SUMMARIZING EQN ONLY	BECH BECH	62002 62002	R R	

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture;
D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality
counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/
kg; T – wt % surfactant mixture; U – mol/l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
. E O NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA2 SO4	25			SUMMARIZING EQN ONLY	BECH	62002	R	
7 ENTRIES FOR COMPOUND								
COMPOUND NO = 164 MOL WGT - 715.0 DODECYL /OXYETHYLENE/ 12 ALCOHOL								
NATURAL DISTRIBUTION OF HEAD GROUPS								
UNK 6.5 X10-3 D HC				TURBIDITY PLT LITE SCATR	BECH	59006	T L	
UNK 6.5 X10-3 D HC				FOTOMTR SPCTR CHNG I2	BECH	59006	G L	
UNK 6.5 X10-3 D HC				TURBIDITY PLT LITE SCATR	BECH	61003	T L	
. E O CA CL2	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA CITRATE	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA2 SO4	25			SUMMARIZING EQN ONLY	BECH	62002	R	
7 ENTRIES FOR COMPOUND								
COMPOUND NO = 165 MOL WGT - 979.4 DODECYL /OXYETHYLENE/ 18 ALCOHOL								
NATURAL DISTRIBUTION OF HEAD GROUPS								
UNK 8.5 X10-3 D HC				TURBIDITY PLT LITE SCATR	BECH	59006	T L	
UNK 8.0 X10-3 D HC				FOTOMTR SPCTR CHNG I2	BECH	59006	G L	
UNK 8.0 X10-3 D HC				TURBIDITY PLT LITE SCATR	BECH	61003	T L	
. E O CA CL2	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA CITRATE	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA2 SO4	25			SUMMARIZING EQN ONLY	BECH	62002	R	
7 ENTRIES FOR COMPOUND								
COMPOUND NO = 166 MOL WGT - 1,199.7 DODECYL /OXYETHYLENE/ 23 ALCOHOL								
NATURAL DISTRIBUTION OF HEAD GROUPS								
UN 1.1 X10-2 D HD				FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
UNK 1.4 X10-2 D HD				TURBIDITY PLT LITE SCATR	BECH	59006	T L	
UNK 1.1 X10-2 D HD				FOTOMTR SPCTR CHNG I2	BECH	59006	G L	
UNK 5.8 X10-3 D HC				FOTOMTR SPCTR CHNG I2	ROSS OLIV	59020	T L	
UNK 6. X10-3 D HE				SURFACE TENSION LOG PLOT	ROSS OLIV	59020	T I	
UNK 1.1 X10-2 D HD				METHOD NOT CITED	BECH	63020	T L	
UNK 1.1 X10-2 D HD				TURBIDITY PLT LITE SCATR	BECH	61003	T L	
UNK 1.1 X10-2 D HD				FOTOMTR SPCTR CHNG BZP4	BECH	62001	T L	
UNK 1.1 X10-2 D HD				SURFACE TENSION UNSPEC	BECH	62001	T L	
UNK 1.1 X10-2 D HD				FOTOMTR SPCTR CHNG I2	BECH	62001	T L	
UNK 1.4 X10-2 D HD				TURBIDITY PLT LITE SCATR	BECH CLIF	59005	T L	
. E O CA CL2	25			SUMMARIZING EQN ONLY	BECH	62002	R	
1.1 E O A DIOXANE	UN 1.3 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
2.3 E O A DIOXANE	UN 2.0 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
3.6 E O A DIOXANE	UN 2.6 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
5.0 E O A DIOXANE	UN 3.6 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
1.6 E O A ETHANOL	UN 1.2 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
3.3 E O A ETHANOL	UN 1.5 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
5.2 E O A ETHANOL	UN 2.0 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
7.0 E O A ETHANOL	UN 2.9 X10-2 D HD			FOTOMTR SPCTR CHNG BZP4	BECH	65013	T L	
. E O NA CITRATE	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	R	
. E O NA2 SO4	25			SUMMARIZING EQN ONLY	BECH	62002	R	
5. E O P SUCROSE	UNK 1.1 X10-2 D HD			METHOD NOT CITED	BECH	63020	T L	
1.0 E 1 P SUCROSE	UNK 1.1 X10-2 D HD			METHOD NOT CITED	BECH	63020	T L	
1.5 E 1 P SUCROSE	UNK 1.0 X10-2 D HD			METHOD NOT CITED	BECH	63020	T L	
2.0 E 1 P SUCROSE	UNK 9.3 X10-3 D HD			METHOD NOT CITED	BECH	63020	T L	
27 ENTRIES FOR COMPOUND								
COMPOUND NO = 167 MOL WGT - 660.9 NYLON BENZENE /OXYETHYLENE/ 10 ALCOHOL								
BRANCHED CHAIN, NATURAL OE DISTRIBUTION								
75 4. X10-3 D HE				METHOD NOT CITED	GINN HARR	61014	T L	
UNK 6.4 X10-3 D HC				TURBIDITY PLT LITE SCATR	BECH	61003	T L	
E O NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	R	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 168 MOL WGT - 881.2 NYLON BENZENE /OXYETHYLENE/ 15 ALCOHOL								
BRANCHED CHAIN, NATURAL OE DISTRIBUTION								
25 1.2 X10-4 M HD				SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L	
25 8.3 X10-3 D HC				TURBIDITY PLT LITE SCATR	BECH	62002	T L	
25 7.7 X10-3 D HC				FOTOMTR SPCTR CHNG I2	BECH	62002	T L	
25 8.1 X10-3 D HC				SURFACE TENSION UNSPEC	BECH	62002	T L	
UNK 7.7 X10-3 D HC				TURBIDITY PLT LITE SCATR	BECH	61003	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; F—wt % solvent; G—mol % surfactant; H—normality; I—mol % surfactant mixture; K—normality; L—molar; M—normal; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
8.6 E-1 M	H CL	25	7.4 X10-3 D	HC	FOTOMTR SPCTR CHNG BZP4	BECH	62001	T L
3.1 E O M	H CL	25	8.1 X10-3 D	HC	SURFACE TENSION UNSPEC	BECH	62001	T L
8.6 E-1 M	H NO3	25	7.7 X10-3 D	HC	FOTOMTR SPCTR CHNG I2	BECH	62001	T L
3.1 E O M	H NO3	25	1.20 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5. E-1 M	NA BR	25	1.50 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5. E-1 M	NA BRO3	25	1.30 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5. E-1 M	NA CL	25	2.90 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5. E-1 M	NA CL	25	1.00 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
8.6 E-1 M	NA CL	25	7.6 X10-5 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
. E O	NA CL	25	8.4 X10-5 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5. E-1 M	NA I	25	5.1 X10-5 M	HD	SUMMARIZING EQN ONLY	BECH	62002	R
5. E-1 M	NA NO3	25	1.08 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5. E-1 M	NA OH	25	9.7 X10-5 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
	0337				SEE CMPD NMBR IN ADDITV	HSIA DUNN	56014	X
21 ENTRIES FOR COMPOUND								
COMPOUND NO =	169 MOL WGT -	1,542.1	NONYL BENZENE /OXYETHYLENE/ 30 ALCOHOL					
	BRANCHED CHAIN, NATURAL OE DISTRIBUTION							
8.6 E-1 M	NA CL	25	2.75 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
E O	NA CL	25	2.36 X10-2 D	HC	TURBIDITY PLT LITE SCATR	BECH	61003	T L
	0337	25	1.10 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
					SUMMARIZING EQN ONLY	BECH	62002	R
					SEE CMPD NMBR IN ADDITV	HSIA DUNN	56014	X
5 ENTRIES FOR COMPOUND								
COMPOUND NO =	170 MOL WGT -	1,420.0	DODECYL /OXYETHYLENE/ 28 ALCOHOL					
	REDUCED' DISTRIBUTION OF HEAD GROUPS							
		25	1.09 X10-2 D	EC	ULTRAfiltration	SCHO	64004	T L
			7.670X10-5 M				M	
		25	1.1 X10-2 D	EE	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T L
			7.74 X10-5 M				M	
2 ENTRIES FOR COMPOUND								
COMPOUND NO =	171 MOL WGT -	348.4	SODIUM 2-N-DODECYL BENZENE SULFONATE					
		25	1.19 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T P
		25	7.3 X10-2 D	CB	UNSPECIFIED CONDUCTANCE	SCHI FOWK	57014	T L
			2.09 X10-3 M				M	
		25	6.5 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.86 X10-3 M				M	
		30	1.19 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T P
		55	5.9 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.69 X10-3 M				M	
2.22 E 1 Q	CAPRYLAMIDE	55	5.0 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.43 X10-3 M				M	
2.22 E 1 Q	N-C10 GLYCEROL ETHER	55	3.3 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			9.47 X10-4 M				M	
2.22 E 1 Q	DECANOL-1	55	4.1 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.17 X10-3 M				M	
2.22 E 1 Q	N-C10 SULFOLANYL ETH	55	3.5 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.00 X10-3 M				M	
2.22 E 1 Q	TETRADECANOL-2	55	6.0 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.72 X10-3 M				M	
2.22 E 1 Q	ISOC5 GLYCEROL ETHER	55	5.4 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.54 X10-3 M				M	
2.22 E 1 Q	C12 CLORHYDRIN GLET*	55	5.7 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.63 X10-3 M				M	
5.55 E 0 Q	C12 ETHANOL AMIDE	55	4.7 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.34 X10-3 M				M	
1.11 E 1 Q	C12 ETHANOL AMIDE	55	3.7 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.06 X10-3 M				M	
1.67 E 1 Q	C12 ETHANOL AMIDE	55	3.2 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			9.18 X10-4 M				M	
2.22 E 1 Q	C12 ETHANOL AMIDE	55	3.1 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			9.86 X10-4 M				M	
2.22 E 1 Q	C12 GLYCEROL ETHER	55	2.9 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			8.32 X10-4 M				M	
2.22 E 1 Q	C12 SULFOLANYLAMIDE	55	3.5 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.00 X10-3 M				M	
2.22 E 1 Q	C8 GLYCEROL ETHER	55	3.6 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.03 X10-3 M				M	
2.22 E 1 Q	N-3SOA*	55	4.8 X10-2 D	CG	VISUAL SPCTR CHNG PNCN	SCHI FOWK	57014	T L
			1.37 X10-3 M				M	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; F—wt % solvent; G—mol % surfactant; H—normality; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2.22 E 1 Q	TMCHCGLET*	55	5.3 X10-2 D 1.52 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2.0 E 1 Q	NA2 S04	55	3.2 X10-2 D 9.18 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2.0 E 1 Q	NA2 S04	55	4.0 X10-2 D 1.14 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2.0 E 1 Q	NA2 S04	55	3.6 X10-2 D 1.14 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2.0 E 1 Q	NA2 S04	55	2.8 X10-2 D 8.03 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
8.0 E 1 Q	NA2 S04	55	3.4 X10-2 D 9.75 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
8.0 E 1 Q	NA2 S04	55	3.1 X10-2 D 8.89 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
8.0 E 1 Q	NA2 S04	55	2.8 X10-2 D 8.03 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
8.0 E 1 Q	NA2 S04	55	2.7 X10-2 D 7.74 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
29 ENTRIES FOR COMPOUND									
COMPOUND NO = 172 MOL WGT -		292.3	SODIUM 2-N-OCTYL BENZENE SULFONATE						
2.22 E 1 Q	N-C10 GLYCEROL ETHER	55	5.56 X10-1 D 1.902X10-2 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2.22 E 1 Q	C12 ETHANOL AMIDE	55	3.47 X10-1 D 1.187X10-2 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2.22 E 1 Q	C8 GLYCEROL ETHER	55	3.43 X10-1 D 1.173X10-2 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
4 ENTRIES FOR COMPOUND									
COMPOUND NO = 173 MOL WGT -		320.4	SODIUM 2-N-DECYL BENZENE SULFONATE						
2.22 E 1 Q	C12 ETHANOL AMIDE	55	1.79 X10-1 D 5.586X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 174 MOL WGT -		376.5	SODIUM 2-N-TETRADECYL BENZENE SULFONATE						
2.22 E 1 Q	C12 ETHANOL AMIDE	55	1.5 X10-2 D 3.98 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 175 MOL WGT -		272.3	SODIUM DODECANE 2-SULFONATE						
2.22 E 1 Q	N-C10 GLYCEROL ETHER	55	3.83 X10-1 D 1.406X10-2 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 176 MOL WGT -		300.4	SODIUM TETRADECANE 2-SULFONATE						
2.22 E 1 Q	N-C10 GLYCEROL ETHER	55	1.13 X10-1 D 3.761X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 177 MOL WGT -		328.4	SODIUM HEXADECANE 2-SULFONATE						
2.22 E 1 Q	N-C10 GLYCEROL ETHER	55	3.5 X10-2 D 1.06 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 178 MOL WGT -		356.5	SODIUM OCTADECANE 2-SULFONATE						
		55	1.3 X10-2 D 3.64 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	SCHI FOWK	57014	T M	L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
2.22 E 1 Q N-C10 GLYCEROL ETHER	55	9. X10-3 D 2.5 X10-4 M	CG	VISUAL SPCTR CHNCE PNCN	SCHI FOWK	57014	T M	L
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 179 MOL WGT -	272.3	SODIUM DODECANE 1-SULFONATE	47005	VALUES FRM REF IN CMC	KLEV	48005	R	
25 9.8 X10-3 M AB SURFACE TENSION LOG PLOT BUJA GODD 65007 L 3								
31.5 9.8 X10-3 W DB KRAFFT POINT SOLUBILITY TART WRIG 39002 T L								
33.5 9.2 X10-3 M CG VISUAL SPCTR CHNCE PNCN KLEV 50003 T L								
33.5 9.2 X10-3 M CG FOTOMTR SPCTR CHNCE PNCN KLEV 47004 T L								
35 9.0 X10-3 M BC ELECTROMOTIVE FORCE CORK GOOD 64012 K L								
35 1.05 X10-2 M CC REFRACTIVE INDEX KLEV 47004 T L								
35 1.0 X10-2 M CD REFRACTIVE INDEX KLEV 47005 T L								
40 1.10 X10-2 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L								
40 1.0 X10-2 M DD REFRACTIVE INDEX KLEV 48005 T L								
40 9.7 X10-3 M AB SURFACE TENSION LOG PLOT BUJA GODD 65007 L 3								
40 1.1 X10-2 N CC UNSPECIFIED CONDUCTANCE TART LELO 55021 T L								
45 1.1 X10-2 M CD REFRACTIVE INDEX KLEV 47005 T L								
50 1.1 X10-2 M DD REFRACTIVE INDEX KLEV 48005 T L								
50 6.9 X10-3 M BG VISUAL SPCTR CHNCE PNCN WEIL STIR 63013 T L								
55 2.88 X10-1 D CG VISUAL SPCTR CHNCE PNCN SCHI FOWK 57014 T L								
1.057X10-2 M								
55 1.2 X10-2 M CD REFRACTIVE INDEX KLEV 47005 T L								
60 1.20 X10-2 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L								
65 1.4 X10-2 M CD REFRACTIVE INDEX KLEV 47005 T L								
80 1.40 X10-2 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L								
UNK 1.1 X10-2 M XG METHOD NOT CITED KLEV RAIS 54004 T L								
2.2 E-1 P BENZENE	33.5	7.5 X10-3 M CG VISUAL SPCTR CHNCE PNCN KLEV 50003 T L						
2.22 E 1 Q N-C10 GLYCEROL ETHER	55	1.17 X10-1 D CG VISUAL SPCTR CHNCE PNCN SCHI FOWK 57014 T L						
1.00 E 2 I NA CL	40	8.1 X10-3 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L						
1.00 E 2 I NA CL	60	9.2 X10-3 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L						
1.00 E 2 I NA CL	80	1.17 X10-2 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L						
26 ENTRIES FOR COMPOUND								
COMPOUND NO = 181 MOL WGT -	216.2	SODIUM OCTYL 1-SULFONATE	23 1.55 X10-1 N CB UNSPECIFIED CONDUCTANCE TART LELO 55021 T 3					
25 1.4 X10-1 M CG VISUAL SPCTR CHNCE PNCN LELO TART 51003 L L								
25 1.55 X10-1 M CC REFRACTIVE INDEX KLEV 48005 T 3								
25 1.53 X10-1 M CC REFRACTIVE INDEX KLEV 47004 T L								
25 1.45 X10-1 M CG FOTOMTR SPCTR CHNCE PNCN KLEV 47004 T L								
40 1.62 X10-1 M CC REFRACTIVE INDEX KLEV 48005 T 3								
50 1.77 X10-1 M CC REFRACTIVE INDEX KLEV 48005 T L								
7 ENTRIES FOR COMPOUND								
COMPOUND NO = 182 MOL WGT -	244.3	SODIUM DECYL 1-SULFONATE	47005 VALUES FRM REF IN CMC	KLEV	48005	R		
47006 VALUES FRM REF IN CMC								
22.5 4.0 X10-2 W DB KRAFFT POINT SOLUBILITY TART WRIG 39002 T L								
25 3.8 X10-2 M CG VISUAL SPCTR CHNCE PNCN KLEV 50003 T L								
25 4.2 X10-2 M CC REFRACTIVE INDEX KLEV 47004 T L								
25 4.1 X10-2 M CC REFRACTIVE INDEX KLEV 47005 T L								
25 3.8 X10-2 M CG FOTOMTR SPCTR CHNCE PNCN KLEV 47004 T L								
26 4.00 X10-2 M CG VISUAL SPCTR CHNCE PNCN CORR HARK 47006 T L								
26 3.87 X10-2 M CG VISUAL SPCTR CHNCE RHD6 CORR HARK 47006 T L								
26 4.00 X10-2 M CG VISUAL SPCTR CHNCE PNCN CORR HARK 47010 T L								
30 3.8 X10-2 N UNSPECIFIED CONDUCTANCE TART LELO 55021 T L								
30 1.066X10 0 D BA FOTOMTR SOLUBLZTN OROT SCHO 66036 T 2								
4.363X10-2 M								
35 4.2 X10-2 M CC REFRACTIVE INDEX KLEV 47005 T L								
40 4.0 X10-2 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L								
40 4.1 X10-2 M DC REFRACTIVE INDEX KLEV 48005 T L								
45 4.5 X10-2 M CC REFRACTIVE INDEX KLEV 47005 T L								
50 4.5 X10-2 M DC REFRACTIVE INDEX KLEV 48005 T L								
55 4.9 X10-2 M CC REFRACTIVE INDEX KLEV 47005 T L								
60 4.3 X10-2 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L								
65 5.5 X10-2 M CC REFRACTIVE INDEX KLEV 47005 T L								
80 5.8 X10-2 M DB SPECFC CONDCTNCE GRAPH WRIG ABBO 39007 T L								
UNK 4.00 X10-2 M CG METHOD NOT CITED HARK MITT 49006 T L								
2.2 E-1 P BENZENE	25	3.4 X10-2 M CG VISUAL SPCTR CHNCE PNCN HARK MITT 50003 T L						
3.40 E-2 P BENZENE	UNK	3.81 X10-2 M CG METHOD NOT CITED HARK MITT 49006 T L						
5.50 E-2 P BENZENE	UNK	3.86 X10-2 M CG METHOD NOT CITED HARK MITT 49006 T L						
8.9 E-2 P BENZENE	UNK	3.89 X10-2 M CG METHOD NOT CITED HARK MITT 49006 T L						

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.22 E-1 P	BENZENE	UNK	3.82 X10-2 M	CG	METHOD NOT CITED	HARK MITT	49006	T L	
1.47 E-1 P	BENZENE	UNK	3.85 X10-2 M	CG	METHOD NOT CITED	HARK MITT	49006	T L	
4.92 E-3 M	NA CL	26	3.82 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
1.35 E-2 M	NA CL	26	3.50 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
2.86 E-2 M	NA CL	26	3.17 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
5.27 E-2 M	NA CL	26	2.93 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
1.0 E-1 M	NA CL	30	5.364X10-1 D	BA	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T 2	M
			2.195X10-2 M						
4.21 E-3 M	NA2 SO4	26	3.72 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
7.97 E-3 M	NA2 SO4	26	3.52 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
1.98 E-2 M	NA2 SO4	26	3.12 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
3.46 E-2 M	NA2 SO4	26	2.73 X10-2 M	CG	VISUAL SPCTR CHNGE PNCC	CORR HARK	47010	T L	
37 ENTRIES FOR COMPOUND									
COMPOUND NO = 183 MOL WGT -		300.4	SODIUM TETRADECYL 1-SULFONATE						
39.5	2.7 X10-3 W	DB	KRAFFT POINT SOLUBILITY			TART WRIG	39002	T L	
40	2.5 X10-3 M	DB	SPECFC CONDCTNCE GRAPH			WRIG ABBO	39007	T L	
40	2.5 X10-3 M	CC	REFRACTIVE INDEX			KLEV	48005	T L	
42.5	2.5 X10-3 M	CG	VISUAL SPCTR CHNGE PNCC			KLEV	50003	T L	
42.5	2.5 X10-3 M	CG	FOTOMTR SPCTR CHNGE PNCC			KLEV	47004	T L	
45	2.9 X10-3 M	CC	REFRACTIVE INDEX			KLEV	48005	T L	
45	3.15 X10-3 M	CC	REFRACTIVE INDEX			KLEV	47004	T L	
50	2.9 X10-3 M	CC	REFRACTIVE INDEX			KLEV	48005	T L	
50	2.03 X10-3 M	BG	VISUAL SPCTR CHNGE PNCC			WEIL STIR	63013	T L	
60	3.3 X10-3 M	DB	SPECFC CONDCTNCE GRAPH			WRIG ABBO	39007	T L	
60	3.3 X10-3 N	DC	UNSPECIFIED CONDUCTANCE			TART LELO	55021	TA L	
80	4.6 X10-3 M	DB	SPECFC CONDCTNCE GRAPH			WRIG ABBO	39007	T L	
2.2 E-1 P	BENZENE	42.5	1.8 X10-3 M	CG	VISUAL SPCTR CHNGE PNCC	KLEV	50003	T L	
13 ENTRIES FOR COMPOUND									
COMPOUND NO = 184 MOL WGT -		328.4	SODIUM HEXADECYL 1-SULFONATE						
47.5	1.05 X10-3 W	DB	KRAFFT POINT SOLUBILITY			TART WRIG	39002	T L	
50	7. X10-4 M	CD	REFRACTIVE INDEX			KLEV	48005	T L	
50	8.0 X10-4 M	CG	FOTOMTR SPCTR CHNGE PNCC			KLEV	47004	T L	
50	4.5 X10-4 M	BG	VISUAL SPCTR CHNGE PNCC			WEIL STIR	63013	T L	
52	9. X10-4 M	CD	REFRACTIVE INDEX			KLEV	48005	T L	
52	1.15 X10-3 M	CC	REFRACTIVE INDEX			KLEV	47004	T L	
6 ENTRIES FOR COMPOUND									
COMPOUND NO = 185 MOL WGT -		294.5	POTASSIUM HEXADECANOATE						
35	1.8 X10-3 M	DD	REFRACTIVE INDEX			KLEV	48005	T L	
45	1.9 X10-3 M	DD	REFRACTIVE INDEX			KLEV	48005	T L	
50	2.2 X10-3 M	XC	REFRACTIVE INDEX			KLEV	53010	T L	
1. E-1 K	K ION	UNK	9.0 X10-5 M	CC	SURFACE TENSION LOG PLOT	ROE BRAS	54013	T L	
1.23 E 1	PH OF SOLUTION								
4 ENTRIES FOR COMPOUND									
COMPOUND NO = 186 MOL WGT -		277.9	HEXADECYL AMMONIUM CHLORIDE						
40	1.07 X10-3 M	CB	SPECFC CONDCTNCE GRAPH			SHIR TAMA	57016	P L	
50	8. X10-4 M	CD	REFRACTIVE INDEX			KLEV	48005	TA L	
55	8.5 X10-4 M	XE	REFRACTIVE INDEX			KLEV	53010	T L	
60	9.9 X10-4 M	CC	EQUIV CONDCTNCE GRAPH			RALS HOER	42002	P L	
4 ENTRIES FOR COMPOUND									
COMPOUND NO = 187 MOL WGT -		306.0	OCTADECYL AMMONIUM CHLORIDE						
60	3. X10-4 M	CD	EQUIV CONDCTNCE GRAPH			RALS HOER	42002	T L	
60	2.5 X10-4 M	CD	REFRACTIVE INDEX			KLEV	48005	T L	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 188 MOL WGT -		154.2	POTASSIUM HEXANOATE						
25	1.55 X10 0 M	DC	REFRACTIVE INDEX			KLEV	48005	T L	
25	1.68 X10 0 M	BG	UNSPEC SPCTR CHNG PNCC			KLEV	58011	T L	
UNK	1.5 X10 0 M	XG	METHOD NOT CITED			KLEV RAIS	54004	T L	
2. E 0 I	K OH	25	1.60 X10 0 M	BG	VISUAL SPCTR CHNGE PNCC	SHIN	54005	G L	
5. E-5 N	PINACYANOL CL (DYE)	25	1.22 X10 0 M	DG	VISUAL SPCTR CHNGE PNCC	HERZ	52015	T L	
1. E-4 N	PINACYANOL CL (DYE)	25	1.49 X10 0 M	DG	VISUAL SPCTR CHNGE PNCC	HERZ	52015	T L	
2. E-4 N	PINACYANOL CL (DYE)	25	1.0 X10 0 M	DG	FOTOMTR SPCTR CHNGE PNCC	HERZ	52015	T L	
5.1 E 0 C	0044	25	1.48 X10 0 M	DG	VISUAL SPCTR CHNGE PNCC	HERZ	52015	T L	
2. E 0 I	K OH	25	1.28 X10 0 M	BG	VISUAL SPCTR CHNGE PNCC	SHIN	54005	G L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions: M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.27 E 1 C 0044 2. E O I K OH	25	1.05 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.48 E 1 C 0044 2. E O I K OH	25	8.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
4.58 E 1 C 0044 2. E O I K OH	25	6.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
6.92 E 1 C 0044 2. E O I K OH	25	4.9 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.8 E O C 0090 2. E O I K OH	25	1.07 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.8 E O C 0090 2. E O I K OH	25	7.3 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
8.9 E O C 0090 2. E O I K OH	25	4.7 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.16 E 1 C 0090 2. E O I K OH	25	2.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.53 E 1 C 0090 2. E O I K OH	25	2.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
8. E-1 C 0092 2. E O I K OH	25	2.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.1 E O C 0092 2. E O I K OH	25	2.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.5 E O C 0092 2. E O I K OH	25	1.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
4.8 E O C 0092 2. E O I K OH	25	5.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
5.3 E O C 0296 2. E O I K OH	25	1.52 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.00 E 1 C 0296 2. E O I K OH	25	1.44 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.60 E 1 C 0296 2. E O I K OH	25	1.36 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.34 E 1 C 0296 2. E O I K OH	25	1.28 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.11 E 1 C 0296 2. E O I K OH	25	1.19 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.33 E 1 C 0296 2. E O I K OH	25	1.16 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
3.85 E 1 C 0296 2. E O I K OH	25	1.13 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
4.92 E 1 C 0296 2. E O I K OH	25	1.04 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
6.44 E 1 C 0296 2. E O I K OH	25	9.6 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
7.78 E 1 C 0296 2. E O I K OH	25	9.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
8. E-1 C 0297 2. E O I K OH	25	8.7 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.1 E O C 0297 2. E O I K OH	25	5.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
3.9 E O C 0297 2. E O I K OH	25	3.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
1.00 E 1 C 0297 2. E O I K OH	5	2.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
2.63 E 1 C 0297 2. E O I K OH	25	1.4 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L	
8.8 E O C 0296 7.36 E 1 C 0044 2. E O I K OH	25	4.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.49 E 1 C 0296 5.54 E 1 C 0044 2. E O I K OH	25	5.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
1.93 E 1 C 0296 4.20 E 1 C 0044 2. E O I K OH	25	5.9 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.28 E 1 C 0296 3.17 E 1 C 0044 2. E O I K OH	25	6.6 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.55 E 1 C 0296 2.36 E 1 C 0044 2. E O I K OH	25	7.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
2.76 E 1 C 0296 1.72 E 1 C 0044 2. E O I K OH	25	8.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2.94 E 1 C 0296 1.17 E 1 C 0044 2. E O I K OH	25	9.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
3.10 E 1 C 0296 7.2 E O C 0044 2. E O I K OH	25	9.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
3.22 E 1 C 0296 3.3 E O C 0044 2. E O I K OH	25	1.07 X10 0 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54003	L L	
46 ENTRIES FOR COMPOUND								
COMPOUND NO = 189 MOL WGT -	308.4	ALPHA SULFOMYRISTIC ACID						
	25	1.3 X10-1 D	CD	EQUIV CONDCTNCE GRAPH	WEIL STIR	56008	T L	
		4.21 X10-3 M					M	
	25	5.2 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	WEIL STIR	56008	K 3	
	28	7. X10-2 D	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		2.2 X10-3 M					M	
	UNK	2.27 X10-3 M	CB	SURFACE TENSION UNSPEC	MAUR STIR	64002	L L	
	RM	7.5 X10-2 D	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		2.43 X10-3 M					M	
	RM	8. X10-2 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		2.5 X10-3 M					M	
6 ENTRIES FOR COMPOUND								
COMPOUND NO = 190 MOL WGT -	336.4	ALPHA SULFOPALMITIC ACID						
	28	2. X10-2 D	CE	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		5.9 X10-4 M					M	
	UNK	6.0 X10-4 M	CB	SURFACE TENSION UNSPEC	MAUR STIR	64002	L L	
	RM	2.3 X10-2 D	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		6.83 X10-4 M					M	
	RM	1.7 X10-2 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		5.05 X10-4 M					M	
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 191 MOL WGT -	364.5	ALPHA SULFOSTEARIC ACID						
	28	5. X10-3 D	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		1.3 X10-4 M					M	
	UNK	1.4 X10-4 M	CD	SURFACE TENSION UNSPEC	MAUR STIR	64002	L L	
	RM	4. X10-3 D	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		1.0 X10-4 M					M	
	RM	5. X10-3 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		1.3 X10-4 M					M	
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 192 MOL WGT -	386.5	SODIUM ETHYL ALPHA SULFOPALMITATE						
	28	9. X10-3 D	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		2.3 X10-4 M					M	
	RM	1.3 X10-2 D	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		3.36 X10-4 M					M	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 193 MOL WGT -	400.5	SODIUM PROPYL ALPHA SULFOPALMITATE						
	28	4. X10-3 D	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		9.9 X10-5 M					M	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 194 MOL WGT -	400.5	SODIUM METHYL ALPHA SULFOSTEARATE						
	28	4. X10-3 D	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		9.9 X10-5 M					M	
	UNK	8. X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	MAUR STIR	64002	T L	
	RM	4. X10-3 D	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		9.9 X10-5 M					M	
	RM	3. X10-3 D	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	56008	T L	
		7.4 X10-5 M					M	
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 195 MOL WGT -	414.5	SODIUM ETHYL ALPHA SULFOSTEARATE						
	28	5. X10-3 D	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L	
		1.2 X10-4 M					M	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Meth.	Method	Authors	Reference	Source	Evaluation
		RM	2. X10-3 D 4.8 X10-5 M	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L M	
		RM	2. X10-3 D 4.8 X10-5 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	56008	T T L M	
3 ENTRIES FOR COMPOUND									
COMPOUND NO =	196 MOL WGT -	428.6	SODIUM PROPYL ALPHA SULFOSTEARATE						
		28	5. X10-4 D 1.1 X10-5 M	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	197 MOL WGT -	488.5	DISODIUM 2 SULFOETHYL ALPHA SULFOPALMITATE						
		25	5.3 X10-1 D 1.08 X10-2 M	CC	EQUIV CONDCTNCE GRAPH	WEIL STIR	56008	T L M	
		RM	3. X10-1 D 6.1 X10-3 M	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L M	
		RM	4. X10-1 D 8.1 X10-3 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	56008	T L M	
3 ENTRIES FOR COMPOUND									
COMPOUND NO =	198 MOL WGT -	516.5	DISODIUM 2 SULFOETHYL ALPHA SULFOSTEARATE						
		25	1.6 X10-1 D 3.09 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	WEIL STIR	56008	T L M	
		28	1.3 X10-1 D 2.51 X10-3 M	CD	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L M	
		RM	1.0 X10-1 D 1.93 X10-3 M	CG	FOTOMTR SPCTR CHNGE PNCN	WEIL STIR	56008	T L M	
3 ENTRIES FOR COMPOUND									
COMPOUND NO =	199 MOL WGT -	428.6	SODIUM ISOPROPYL ALPHA SULFOSTEARATE						
		28	1. X10-3 D 2.3 X10-5 M	CE	SURFACE TENSION UNSPEC	WEIL STIR	56008	T L M	
1 ENTRIES FOR COMPOUND									
COMPOUND NO =	200 MOL WGT -	250.3	DODECYL SULFONIC ACID						
		0	1.00 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L	
		0	9.5 X10-3 W	CB	FREEZING POINT	MCBA DYE	39011	K L	
		25	8.5 X10-3 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L	
		25	9.2 X10-3 W	CB	EQUIV CONDCTNCE GRAPH	MCBA DYE	39011	K L	
		25	8. X10-2 W	CF	EQUIV CONDCTNCE GRAPH	MCBA DYE	39011	T L	
		29	7.5 X10-3 M	BB	SURFACE TENSION LOG PLOT	BRAD	48022	T L	
		40	9.2 X10-3 W	CC	ELECTROMOTIVE FORCE	TART LING	48007	P L	
		50	9.5 X10-3 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L	
		50	5.55 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	63013	T L	
		70	1.20 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L	
		90	1.55 X10-2 M	CC	SPECFC CONDCTNCE GRAPH	BRAD HUFF	48004	T L	
		UNK	9. X10-3 M	XG	METHOD NOT CITED	KLEV RAIS	54004	T L	
		UNK	4.39 X10-3 M	BB	SURFACE TENSION LOG PLOT	WEIL STIR	63013	T L	
13 ENTRIES FOR COMPOUND									
COMPOUND NO =	201 MOL WGT -	524.8	DECYL /OXYETHYLENE/8 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS						
		30	6.0 X10-2 D 1.14 X10-3 M	EC	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	
1.50 E-2 Q	N-DECANE	30	4.0 X10-2 D 7.62 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	
1.32 E 0 Q	N-DECANE	30	5.7 X10-2 D 1.08 X10-3 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	
2.33 E 0 Q	N-DECANE	30	5.4 X10-2 D 1.02 X10-3 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	
3.16 E 0 Q	N-DECANE	30	5.2 X10-2 D 9.90 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	
3.78 E 0 Q	N-DECANE	30	5.1 X10-2 D 9.71 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	
4.93 E 0 Q	N-DECANE	30	4.8 X10-2 D 9.14 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	
1.20 E 1 Q	N-DECANE	30	4.0 X10-2 D 7.62 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T L M	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
3.40 E 0 Q	DECANOL-1	30	5.2 X10-2 D 9.90 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
6.19 E 0 Q	DECANOL-1	30	4.5 X10-2 D 8.57 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
8.50 E 0 Q	DECANOL-1	30	4.0 X10-2 D 7.62 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
1.142E 1 Q	DECANOL-1	30	3.3 X10-2 D 6.28 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
1.661E 1 Q	DECANOL-1	30	2.0 X10-2 D 3.81 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
13 ENTRIES FOR COMPOUND									
COMPOUND NO = 202 MOL WGT - 657.0 DECYL /OXYETHYLENE/11 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS									
		30	9.5 X10-2 D 1.44 X10-3 M	EC	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
1.40 E 0 Q	N-DECANE	30	9.2 X10-2 D 1.40 X10-3 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
2.58 E 0 Q	N-DECANE	30	9.0 X10-2 D 1.36 X10-3 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
3.31 E 0 Q	N-DECANE	30	8.8 X10-2 D 1.33 X10-3 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
5.67 E 0 Q	DECANOL-1	30	8.4 X10-2 D 1.27 X10-3 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
1.018E 1 Q	DECANOL-1	30	7.5 X10-2 D 1.14 X10-3 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
1.712E 1 Q	DECANOL-1	30	6.3 X10-2 D 9.58 X10-4 M	ED	TURBIDITY PLT LITE SCATR	NAKA KURI	60006	T M	L
7 ENTRIES FOR COMPOUND									
COMPOUND NO = 203 MOL WGT - 235.8 DECYL TRIMETHYL AMMONIUM CHLORIDE									
		25	6.5 X10-2 M	XG	VISUAL SPCTR CHNGE	KLEV	53010	T L	
3.33 E 1 C 0041		25	6.11 X10-2 M	BB	UNSPECIFIED CONDUCTANCE	HOYE MARM	61002	T 3	
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 204 MOL WGT - 701.0 DECYL /OXYETHYLENE/12 OXYMETHYL REDUCED POLYDISPERSION OF HEAD GROUPS									
		09.7	1.70 X10-1 D 2.425X10-3 M	EC	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		29.0	1.10 X10-1 D 1.569X10-3 M	EC	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		30	1.05 X10-1 D 1.497X10-3 M	EC	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		45	8.2 X10-2 D 1.16 X10-3 M	EC	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		50.7	7.8 X10-2 D 1.11 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		50	7.8 X10-2 D 1.11 X10-3 M	EC	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		58.5	7.0 X10-2 D 9.98 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		69.7	6.2 X10-2 D 8.84 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		73.4	6.0 X10-2 D 8.55 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
		75.0	6.0 X10-2 D 8.55 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.86 E 0 Q	N-DECANE	09.6	1.65 X10-1 D 2.353X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
2.6 E 0 Q	N-DECANE	10.0	1.63 X10-1 D 2.325X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.86 E 0 Q	N-DECANE	30.0	1.05 X10-1 D 1.497X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
4.4 E 0 Q	N-DECANE	30.0	1.02 X10-1 D 1.455X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.86 E 0 Q	N-DECANE	50.0	7.8 X10-2 D 1.11 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
8.7 E 0 Q	N-DECANE	50.0	7.0 X10-2 D 9.98 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.86 E 0 Q	N-DECANE	60.0	7.0 X10-2 D 9.98 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.35 E 1 Q	N-DECANE	60.0	6.8 X10-2 D 9.70 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.86 E 0 Q	N-DECANE	66.6	6.2 X10-2 D 8.84 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.86 E 0 Q	N-DECANE	69.0	6.2 X10-2 D 8.84 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 E 0 Q	DECANOL-1	10.0	1.45 X10-1 D 2.068X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 0	DECANOL-1	29.9	8.8 X10-2 D 1.25 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.87 E 0 Q	DECANOL-1	30	1.05 X10-1 D 1.497X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
4.56 E 0 Q	DECANOL-1	30	9.9 X10-2 D 1.41 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
7.75 E 0 Q	DECANOL-1	30	9.2 X10-2 D 1.31 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 E 0 Q	DECANOL-1	30	8.8 X10-2 D 1.25 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.11 E 1 Q	DECANOL-1	30	8.5 X10-2 D 1.21 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.91 E 1 Q	DECANOL-1	30	7.0 X10-2 D 9.98 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 E 0 Q	DECANOL-1	43.4	7.6 X10-2 D 1.08 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
5.37 E 0 Q	DECANOL-1	45	7.8 X10-2 D 1.11 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 E 0 Q	DECANOL-1	45	7.5 X10-2 D 1.06 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.07 E 1 Q	DECANOL-1	45	7.4 X10-2 D 1.05 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.28 E 1 Q	DECANOL-1	45	7.3 X10-2 D 1.04 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.73 E 1 Q	DECANOL-1	45	6.9 X10-2 D 9.84 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 E 0 Q	DECANOL-1	49.7	7.0 X10-2 D 9.98 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
1.86 E 0 Q	DECANOL-1	50	7.8 X10-2 D 1.11 X10-3 R	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
2.23 E 0 Q	DECANOL-1	50	7.6 X10-2 D 1.08 X10 3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
3.53 ¹ E 0 Q	DECANOL-1	50	7.5 X10-2 D 1.06 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
5.58 E 0 Q	DECANOL-1	50	7.3 X10-2 D 1.04 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
7.52 E 0 Q	DECANOL-1	50	7.1 X10-2 D 1.01 X10-3 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
8.7 E 0 Q	DECANOL-1	50	7.0 X10-2 D 9.98 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 E 0 Q	DECANOL-1	55.4	6.6 X10-2 D 9.41 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L
9.17 E 0 Q	DECANOL-1	61.4	6.2 X10-2 D 8.84 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62009	T M	L

43 ENTRIES FOR COMPOUND

COMPOUND NO = 205 MOL WGT = 729.1 DODECYL /OXYETHYLENE/12 OXYMETHYL
REDUCED POLYDISPERSION OF HEAD GROUPS

5.	E-1 M	CA CL2	30	4. X10-2 D 5.4 X10-4 M	EE	TURBIDITY PLT LITE SCATR	KURI	62011	T M
1.	E 0 M	CA CL2	30	2. X10-2 D 1.37 X10-4 M	EE	TURBIDITY PLT LITE SCATR	KURI	62011	T M
5.	E-1 M	CA CL2	50	1.5 X10-2 D 2.05 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T M
1.	E 0 M	CA CL2	50	8. X10-3 D 1.0 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T M

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/
kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.	E O M	CA CL2	60	5. X10-3 D 6.8 X10-5 M	EE	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
5.	E-1 M	CA CL2	64	1.0 X10-2 D 1.37 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
2.	E-1 M	NA CL	30	2.3 X10-2 D 3.15 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	30	2.3 X10-2 D 3.15 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
1.	E O M	NA CL	30	2.0 X10-2 D 2.74 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
2.	E-1 M	NA CL	50	1.6 X10-2 D 2.19 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	50	1.6 X10-2 D 2.19 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
1.	E O M	NA CL	50	1.5 X10-2 D 2.05 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
1.	E O M	NA CL	60	1.2 X10-2 D 1.64 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
2.	E-1 M	NA CL	70	1.2 X10-2 D 1.64 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	70	1.2 X10-2 D 1.64 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI	62011	T L M	
2.	E-1 M	NA CL	30	2.3 X10-2 D 3.15 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	30	2.5 X10-2 D 3.42 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	30	2.8 X10-2 D 3.84 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	30	2.8 X10-2 D 3.84 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	30	2.9 X10-2 D 3.97 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	50	2.2 X10-2 D 3.01 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E O T	0001	50	2.5 X10-2 D 3.42 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	50	3.0 X10-2 D 4.11 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	50	3.3 X10-2 D 4.52 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	50	3.6 X10-2 D 4.93 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	70	2.0 X10-2 D 2.74 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	70	2.6 X10-2 D 3.56 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	70	3.4 X10-2 D 4.66 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	70	4.0 X10-2 D 5.48 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	
2.	E-1 M	NA CL	70	5.0 X10-2 D 6.85 X10-4 M	ED	TURBIDITY PLT LITE SCATR	KURI NAKA	62010	T L M	

35 ENTRIES FOR COMPOUND

COMPOUND NO = 206 MOL WGT - 624 9 TRITON X-100(P-T-OCTYL BENZENE/OXYETHYLENE/9
NATURAL DISTRIBUTION OF HEAD GROUPS

5.	E-1 M	CA CL2	0	9. X10-4 W	HD	FREEZING POINT	GONI MCBA	47007	T L
1.0	E O M	H NO3	25	1.5 X10-2 D	HD	SURFACE TENSION LOG PLOT	MANK	64010	T L
1.0	E O M	H NO3	25	2.2 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
7.	E-1 M	H CL04	25	1.6 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
7.	E-1 M	H CL04	25	3.7 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
7.	E-1 M	H CL04	25	2.0 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
1.0	E O M	K CL	25	2.2 X10-3 D	HE	SURFACE TNSN LINEAR PLOT	COLI	50012	T L
1.0	E O M	K CL	25	1.5 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
1.0	E O M	K NO3	25	2.5 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
1.0	E O M	K NO3	25	2.2 X10-3 D	HE	SURFACE TNSN LINEAR PLOT	COLI	50012	T L
1.0	E O M	K NO3	25	2.0 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L
1.0	E O M	K OH	25	2.0 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/
kg; T—wt % surfactant mixture; U—mol/(l orgk); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

		Additives	Temp.	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
1.0	E 0 M	K CH3	25	1.8 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L	
5	E-1 M	HA CL 0139	30	4 X10-2 D	HE	TURBIDITY PLT LITE SCATR SEE CMPL NMBR IN ADDITV	KURI MANK	62011 64010	T L X	
2	E-1	IONIC STRENGTH	25	9 X10-3 D	HG	REACTN RATE SOLUBILIZATE	TOHG REEV	65030	T L	
1.01 E-1		PH OF SOLUTION	UNK	8 X10-3 D	HG	FOTOMTR SPCTR CHNGE RDG	TOHG REEV	65030	T L	
2	E-1	IONIC STRENGTH	UNK	1.01 X10-1 PH OF SOLUTION	HG					
5.0	E 0 M	NH3	25	2.0 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L	
5.0	E 0 M	NH3	25	1.8 X10-3 D	HG	POLAROGRAPHIC MAXIMUM	COLI	50012	T L	
1	E-1 M	NH4 CL								
1	E-1 M	NH4 CL								
23	ENTRIES FOR COMPOUND									
COMPOUND NO =	207	MOL WGT =	250.4	P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			15	4.3 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			25	4.86 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L	
			25	4.95 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			35	5.6 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			45	6.3 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			55	7.65 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			65	9.65 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			75	1.29 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			85	1.95 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
9	ENTRIES FOR COMPOUND									
COMPOUND NO =	208	MOL WGT =	294.4	P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			15	7.30 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			25	1.32 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL 3	
			25			QUESTIONABLE CRITERION	CROO FORD	63017	R	
			25	7.65 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			35	7.90 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			45	8.65 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			55	9.65 X10-6 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			65	1.11 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			75	1.32 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			85	1.62 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
10	ENTRIES FOR COMPOUND									
COMPOUND NO =	209	MOL WGT =	338.5	P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			15	1.02 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			25	9.7 X10-6 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL 3	
			25			QUESTIONABLE CRITERION	CROO FORD	63017	R	
			25	1.03 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			35	1.07 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			45	1.13 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			55	1.23 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			65	1.39 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			75	1.68 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			85	2.12 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
10	ENTRIES FOR COMPOUND									
COMPOUND NO =	210	MOL WGT =	382.5	P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL						
		HOMOGENEOUS HEAD GROUP								
			15	1.34 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			25	1.26 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL 3	
			25			QUESTIONABLE CRITERION	CROO FORD	63017	R	
			25	1.29 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			35	1.30 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			45	1.41 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			55	1.59 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			65	1.72 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			75	1.91 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
			85	2.13 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L	
	E 0	0216				QUESTIONABLE CRITERION	CROO FORD	63017	R	
3.33 E-1 C	0216		25	1.47 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L	
5.00 E-1 C	0216		25	1.75 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L	
6.67 E-1 C	0216		25	2.17 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L	
14	ENTRIES FOR COMPOUND									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/d(mg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 211 MOL WGT - HOMOGENEOUS HEAD GROUP		426.6	P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL				
	15	1.81 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	25	1.54 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	L 3
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
	25	1.72 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	35	1.64 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	45	1.64 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	55	1.72 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	65	1.90 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	75	2.2 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	85	2.35 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
10 ENTRIES FOR COMPOUND							
COMPOUND NO = 212 MOL WGT - HOMOGENEOUS HEAD GROUP		470.7	P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL				
	15	2.70 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	25	2.05 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	L 3
	5			QUESTIONABLE CRITERION	CROO FORD	63017	R
	5	2.5 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	35	2.37 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	45	2.28 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	640	L L
	55	2.30 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	65	2.55 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	75	2.87 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	85	3.5 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
10 ENTRIES FOR COMPOUND							
COMPOUND NO = 213 MOL WGT - HOMOGENEOUS HEAD GROUP		514.7	P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL				
	15	2.91 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	25	2.46 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	L 3
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
	25	2.68 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	35	2.44 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	45	2.43 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	55	2.41 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	65	2.50 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	75	2.68 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	85	2.90 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
10 ENTRIES FOR COMPOUND							
COMPOUND NO = 214 MOL WGT - HOMOGENEOUS HEAD GROUP		558.8	P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL				
	15	2.94 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	25	2.80 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL 3
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
	25	2.83 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	35	2.58 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	45	2.50 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	55	2.46 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	65	2.55 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	75	2.74 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	85	2.98 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
10 ENTRIES FOR COMPOUND							
COMPOUND NO = 215 MOL WGT - HOMOGENEOUS HEAD GROUP		602.8	P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL				
	15	3.22 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	25	3.35 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL 3
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
	25	3.04 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	35	2.75 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	45	2.62 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	55	2.70 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	65	2.80 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	75	2.91 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	85	3.18 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
10 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; F—wt % solvent; G—mol % surfactant; H—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 216 MOL WGT - HOMOGENEOUS HEAD GROUP	646.9	P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL					
	15	3.30 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	25	3.23 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	25	3.35 X10-4 M	BC	SURFACE TENSION LOG PLOT	CROO FORD	63017	L 3
	5			QUESTIONABLE CRITERION	CROO FORD	63017	R
	35	3.03 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	45	2.80 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	55	2.90 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	65	3.00 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	75	3.12 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
	85	3.38 X10-4 M	BD	SURFACE TENSION LOG PLOT	CROO FORD	64014	GL L
E O 0210	25			SEE CMPD NMBR IN ADDITV	CROO FORD	63017	X
E O 0210	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
12 ENTRIES FOR COMPOUND							
COMPOUND NO = 217 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	250.4	P-T-OCTYL BENZENE /OXYETHYLENE/1 ALCOHOL					
	25	4.85 X10-5 M	EE	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 218 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	294.4	P-T-OCTYL BENZENE /OXYETHYLENE/2 ALCOHOL					
	25	6.8 X10-5 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 219 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	338.5	P-T-OCTYL BENZENE /OXYETHYLENE/3 ALCOHOL					
	25	1.14 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 220 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	382.5	P-T-OCTYL BENZENE /OXYETHYLENE/4 ALCOHOL					
	25	1.05 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
E O 0226	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
3.33 E 1 C 0226	25	1.28 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	L L
5100 E 1 C 0226	25	1.53 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
6.67 E 1 C 0226	25	2.04 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
6 ENTRIES FOR COMPOUND							
COMPOUND NO = 221 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	426.6	P-T-OCTYL BENZENE /OXYETHYLENE/5 ALCOHOL					
	25	1.17 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 222 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	470.7	P-T-OCTYL BENZENE /OXYETHYLENE/6 ALCOHOL					
	25	1.80 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	L L
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 223 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	514.7	P-T-OCTYL BENZENE /OXYETHYLENE/7 ALCOHOL					
	25	1.84 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 224 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	558.8	P-T-OCTYL BENZENE /OXYETHYLENE/8 ALCOHOL					
	25	2.47 X10-4 M	EC	SURFACE TENSION LOG PLOT	CROO FORD	63017	GL L
	25			QUESTIONABLE CRITERION	CROO FORD	63017	R
2 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation	
COMPOUND NO = 225 MOL WGT -	602.8	P-T-OCTYL BENZENE /OXYETHYLENE/9 ALCOHOL						
NATURAL DISTRIBUTION OF HEAD GROUPS								
25	2.90 X10-4 M	EC	SURFACE TENSION MINIMUM	CROO FORD	63017	L L		
5			QUESTIONABLE CRITERION	CROO FORD	63017	R		
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 226 MOL WGT -	646.9	P-T-OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL						
NATURAL DISTRIBUTION OF HEAD GROUPS								
25	3.20 X10-4 M	EC	SURFACE TENSION MINIMUM	CROO FORD	63017	GL L		
25			QUESTIONABLE CRITERION	CROO FORD	63017	R		
0220			SEE CMPD NMBR IN ADDITV	CROO FORD	63017	X		
E 0			QUESTIONABLE CRITERION	CROO FORD	63017	R		
0220								
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 227 MOL WGT -	911.3	P-T-OCTYL BENZENE /OXYETHYLENE/16 ALCOHOL						
NATURAL DISTRIBUTION OF HEAD GROUPS								
25	4.3 X10-4 M	EC	SURFACE TENSION MINIMUM	CROO FORD	63017	GL L		
25			QUESTIONABLE CRITERION	CROO FORD	63017	R		
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 228 MOL WGT -	1,968.7	P-T-OCTYL BENZENE /OXYETHYLENE/40 ALCOHOL						
NATURAL DISTRIBUTION OF HEAD GROUPS								
25	8.1 X10-4 M	EC	SURFACE TENSION MINIMUM	CROO FORD	63017	GL L		
25			QUESTIONABLE CRITERION	CROO FORD	63017	R		
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 229 MOL WGT -	286.3	SODIUM TRIDECANE 1-SULFONATE						
	50	3.52 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	63013	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 230 MOL WGT -	314.4	SODIUM PENTADECANE 1-SULFONATE						
	50	6.6 X10-4 M	BG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	63013	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 231 MOL WGT -	342.5	SODIUM HEPTADECANE 1-SULFONATE						
	50	2.1 X10-4 M	BG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	63013	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 232 MOL WGT -	334.5	OCTADECANE 1-SULFONIC ACID						
	50	1.0 X10-4 M	BG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	63013	T L	
	UNK	1.0 X10-4 M	BD	SURFACE TENSION LOG PLOT	WEIL STIR	63013	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 233 MOL WGT -	352.3	DISODIUM ALPHA SULFO MYRISTATE						
	25		QUESTIONABLE CRITERION		WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 234 MOL WGT -	380.4	DISODIUM ALPHA SULFO PALMITATE						
	25		QUESTIONABLE CRITERION		WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 235 MOL WGT -	302.3	SODIUM ALPHA SULFO LAURIC ACID						
	25		QUESTIONABLE CRITERION		WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 236 MOL WGT -	330.4	SODIUM ALPHA SULFO MYRISTIC ACID						
	25		QUESTIONABLE CRITERION		WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 237 MOL WGT -	358.4	SODIUM ALPHA SULFO PALMITIC ACID						
	25		QUESTIONABLE CRITERION		WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 238 MOL WGT –	288.3 25	SODIUM DODECANE 1-HYDROXY 2-SULFONATE QUESTIONABLE CRITERION			WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 239 MOL WGT –	316.4 25	SODIUM TETRADECANE 1-HYDROXY 2-SULFONATE QUESTIONABLE CRITERION			WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 240 MOL WGT –	344.4 25	SODIUM HEXADECANE 1-HYDROXY 2-SULFONATE QUESTIONABLE CRITERION			WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 241 MOL WGT –	372.5 25	SODIUM OCTADECANE 1-HYDROXY 2-SULFONATE QUESTIONABLE CRITERION			WEIL STIR	63013	R	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 242 MOL WGT –	264.4 50 UNK	TRIDECANE 1-SULFONIC ACID 2.6 X10-3 M 2.53 X10-3 M	BG BB	VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	WEIL STIR WEIL STIR	63013 63013	T L T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 243 MOL WGT –	278.4 25 25 50 UNK	TETRADECANE 1-SULFONIC ACID 2.6 X10-3 W 2.10 X10-3 W 1.32 X10-3 M 1.36 X10-3 M	CF CA BG BB	EQUIV CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	MCBA DYE MCBA DYE WEIL STIR WEIL STIR	39011 39011 63013 63013	T L P L T L T L	
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 244 MOL WGT –	292.4 50 UNK	PENTADECANE 1-SULFONIC ACID 4.8 X10-4 M 7.9 X10-4 M	BG BC	VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	WEIL STIR WEIL STIR	63013 63013	T L T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 245 MOL WGT –	306.5 40 50 60 80 UNK	HEXADECANE 1-SULFONIC ACID 5.9 X10-4 M 2.8 X10-4 M 8.25 X10-4 M 1.24 X10-3 M 4.2 X10-4 M	DB BG DA DA BC	SPECFC CONDCTNCE GRAPH VISUAL SPCTR CHNGE PNCN SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH SURFACE TENSION LOG PLOT	HART WEIL STIR HART HART WEIL STIR	36002 63013 36002 36002 63013	P L T L P L P L T L	
8.4 E 2 I PENTANOL-1 5.63 E 1 H GLYCEROL 9.58 E 1 I H CL	60 60 60	7.5 X10-4 M 2.80 X10-3 M 6.1 X10-4 M	DB DB DB	SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH	HART HART HART	36002 36002 36002	P L P L P L	
8 ENTRIES FOR COMPOUND								
COMPOUND NO = 246 MOL WGT –	320.5 50 UNK	HEPTADECANE 1-SULFONIC ACID 1.4 X10-4 M 1.8 X10-4 M	BG BD	VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	WEIL STIR WEIL STIR	63013 63013	T L T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 247 MOL WGT –	266.3 50 UNK	DODECANE 1-HYDROXY 2-SULFONIC ACID 1.61 X10-2 M 1.31 X10-2 M	BG BB	VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	WEIL STIR WEIL STIR	63013 63013	T L T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 248 MOL WGT –	294.4 50 UNK	TETRADECANE 1-HYDROXY 2-SULFONIC ACID 2.61 X10-3 M 1.94 X10-3 M	BG BB	VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	WEIL STIR WEIL STIR	63013 63013	T L T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 249 MOL WGT –	322.5 50 UNK	HEXADECANE 1-HYDROXY 2-SULFONIC ACID 5.8 X10-4 M 6.5 X10-4 M	VISUAL SPCTR CHNGE PNCN BB	SURFACE TENSION LOG PLOT	WEIL STIR WEIL STIR	63013 63013	T L T L	
2 ENTRIES FOR COMPOUND								

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC		Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 250 MOL WGT -		350.5	OCTADECANE	1-HYDROXY 2-SULFONIC ACID						
		50	2.1 X10-4 M	CG	VISUAL SPCTR CHNGE PNCN	WEIL STIR	63013	T L		
		UNK	2.2 X10-4 M	CC	SURFACE TENSION LOG PLOT	WEIL STIR	63013	T L		
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 251 MOL WGT -		173.3	OCTYL DIMETHYL AMINE OXIDE							
		25	1.6 X10-2 M	BD	HEAT OF DILUTION	BENJ	64016	L L		
		27	1.5 X10-1 M	BC	TURBIDITY PLT LITE SCATR	HERR	62005	G L		
		30	2.4 X10 0 P	BC	DENSITY	BENJ	66040	T L		
			1.38 X10-1 S					M		
3 ENTRIES FOR COMPOUND										
COMPOUND NO = 252 MOL WGT -		201.3	DECYL DIMETHYL AMINE OXIDE							
		25	1.9 X10-2 M	BD	HEAT OF DILUTION	BENJ	64016	L L		
		27	1.5 X10-2 M	BC	TURBIDITY PLT LITE SCATR	HERR	62005	G L		
		30	3.3 X10-1 P	BC	DENSITY	BENJ	66040	T L		
			1.63 X10-2 S					M		
3 ENTRIES FOR COMPOUND										
COMPOUND NO = 253 MOL WGT -		257.5	TETRADECYL DIMETHYL AMINE OXIDE							
		27	2.7 X10-4 M	BC	TURBIDITY PLT LITE SCATR	HERR	62005	G L		
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 254 MOL WGT -		187.3	NONYL DIMETHYL AMINE OXIDE							
		25	5.4 X10-2 M	BC	HEAT OF DILUTION	BENJ	64016	L 3		
		30	1.1 X10 0 P	BD	DENSITY	BENJ	66040	T L		
			5.87 X10-2 S					M		
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 255 MOL WGT -		354.6	POTASSIUM 9,10 DIHYDROXY STEARATE							
		55	1.1 X10-2 M	XC	REFRACTIVE INDEX	KLEV	53010	T L		
		55	8.0 X10-3 M	XB	REFRACTIVE INDEX	KLEV	53010	T L		
1. E-3 M K OH		60	7.5 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	GREG TART	48012	T 3		
3 ENTRIES FOR COMPOUND										
COMPOUND NO = 256 MOL WGT -		322.6	POTASSIUM STEARATE							
		55	4.5 X10-4 M	XE	REFRACTIVE INDEX	KLEV	53010	T L		
1. E-3 M K OH		60	5. X10-4 M	BE	EQUIV CONDCTNCE GRAPH	GREG TART	48012	T L		
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 257 MOL WGT -		332.3	SODIUM DI-N-BUTYL SULFOSUCCINATE							
		25	2.0 X10-1 M	BC	SURFACE TENSION LOG PLOT	WILL DIXO	57009	TL L		
		29.9	2.1 X10-1 M	BE	SPECFC CONDCTNCE GRAPH	MILL DIXO	58001	T L		
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 258 MOL WGT -		360.3	SODIUM DI-N-AMYL SULFOSUCCINATE							
		48016			VALUES FRM REF IN CMC	KOLT STRI	49005	R		
		25	5.3 X10-2 M	BC	SURFACE TENSION LOG PLOT	WILL DIXO	57009	AL 3		
		29.9	7.3 X10-2 M	BE	SPECFC CONDCTNCE GRAPH	MILL DIXO	58001	T L		
		30	9.5 X10-2 M	HD	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	48016	T L		
5. E-1 N NA CL		50	9.5 X10-2 M	HD	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	48016	T L		
1. E O N NA CL		50	4.5 X10-2 M	HE	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	49005	T L		
7 ENTRIES FOR COMPOUND										
COMPOUND NO = 259 MOL WGT -		388.4	SODIUM DI-N-HEXYL SULFOSUCCINATE							
		25.0	1.0 X10 0 P	HD	DENSITY	VETT	47011	T L		
		25.0	1.1 X10 0 P	HD	VISCOSITY	VETT	47011	T L		
		25	1.24 X10-2 M	BC	SURFACE TENSION LOG PLOT	WILL DIXO	57009	TL 3		
		25	2.7 X10-2 M	XG	VISCOSITY MINIMUM	SATA TYUZ	53006	T L		
		29.9	1.28 X10-2 M	BE	SPECFC CONDCTNCE GRAPH	MILL DIXO	58001	T L		
		29.9	1.19 X10-2 M	BE	EQUIV CONDCTNCE GRAPH	MILL DIXO	58001	T L		
		UNK	3.8 X10-2 M	BG	METHOD NOT CITED	KLEV CARR	56001	T L		
7 ENTRIES FOR COMPOUND										
COMPOUND NO = 260 MOL WGT -		444.5	SODIUM DI-N-OCTYL SULFOSUCCINATE							
		20	4.5 X10-3 M	HG	FOTOMTR SPCTR CHNGE PNCN	TAUB KONS	60033	T L		

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l (or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation	
	20	2.7 X10-3 M	HG	FOTOMTR SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	5.4 X10-3 M	HG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	2.6 X10-3 M	HG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	2.6 X10-3 M	HG	VISUAL SPCTR CHNGE RHD6	TAUB KONS	60033	T L	
	25	6.8 X10-4 M	BC	SURFACE TENSION LOG PLOT	WILL DIXO	57009	TL 3	
	25	9. X10-4 N	HE	UNSPECIFIED CONDUCTANCE	HAFF PICC	42003	T L	
	29.9	6.4 X10-4 M	BE	EQUIV CONDCTNCE GRAPH	MILL DIXO	58001	T L	
	40	5.1 X10-3 M	HC	UNSPECIFIED CONDUCTANCE	TAUB KONS	60033	T L	
	UNK	4. X10-2 M	HE	ELECTROMOTIVE FORCE	STAN RADL	60021	T L	
10 ENTRIES FOR COMPOUND								
COMPOUND NO = 261 MOL WGT -	332.3	SODIUM DI-ISOBUTYL SULFOSUCCINATE						
	25	2.0 X10-1 M	BC	SURFACE TENSION LOG PLOT	WILL DIXO	57009	TL L	
	25			QUESTIONABLE CRITERION	HAFF PICC	42003	R	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 262 MOL WGT -	444.5	SODIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE						
	25	2.5 X10-3 M	BC	SURFACE TENSION LOG PLOT	WILL DIXO	57009	TL 3	
	25	5.5 X10-3 N	HE	UNSPECIFIED CONDUCTANCE	HAFF PICC	42003	T L	
	29.9	5.6 X10-3 M	BE	EQUIV CONDCTNCE GRAPH	MILL DIXO	58001	T L	
	29.9	6.1 X10-3 M	BE	SPECFC CONDCTNCE GRAPH	MILL DIXO	58001	T L	
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 263 MOL WGT -	304.4	SODIUM OLEATE /CIS-9-OCTADECENOATE/						
	20			QUESTIONABLE CRITERION	HESS PHIL	39009		
	24.7	7. X10-2 P	HD	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L	
	25	7.2 X10-4 M	HG	FOTOMTR SPCTR CHNGE PNCN	GINN HARR	58008	T L	
	25	2.64 X10-3 M	HC	SPECFC CONDCTNCE GRAPH	GINN HARR	58008	T L	
	25	2.9 X10-2 D	HE	METHOD NOT CITED	GINN HARR	61014	T L	
	25	2.10 X10-3 M	DC	SPECFC CONDCTNCE GRAPH	FLOC GRAH	53004	P L	
	40	2.15 X10-3 M	DC	SPECFC CONDCTNCE GRAPH	FLOC GRAH	53004	P L	
	40	3.0 X10-3 M	DD	SPECFC CONDCTNCE GRAPH	TAMA SHIR	58007	P L	
	50	3.5 X10-3 M	HC	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L	
	60	3.2 X10-3 M	DD	SPECFC CONDCTNCE GRAPH	TAMA SHIR	58007	P L	
	60			QUESTIONABLE CRITERION	TAMA NAKA	53001	R	
	75	3.5 X10-2 D	HE	METHOD NOT CITED	GINN HARR	61014	T L	
	UNK	1.4 X10-3 M	DD	WIEN EFFECT	EXNE	48018	T L	
	UNK	6.1 X10-2 D	HB	SPECFC CONDCTNCE GRAPH	HARR	58004	K L	
	UNK	1. X10-3 M	DG	FOTOMTR SPCTR CHNGE PNCN	TAMA SHIR	58007	T L	
	UNK	2.7 X10-3 M	DD	EQUIV CONDCTNCE GRAPH	TAMA SHIR	58007	T L	
5. E-2 P	DECANOL-1	24.7	3. X10-2 P	HE	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L
2. E 0 M	ETHANOL	40	1.7 X10-3 M	DD	AVER SP EQUIV COND	TAMA SHIR	58007	P L
7. E-2 P	LAURYL ALCOHOL	24.7	4. X10-2 P	HE	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L
4. E 0 I	NA OH	25	1.5 X10-3 W	DG	VISUAL SPCTR CHNGE RHD6	FINE MCBA	48011	T L
6. E 0 I	NA OH	25	1.5 X10-3 M	CC	EQUIV CONDCTNCE GRAPH	FLOC GRAH	53004	T L
1.67 E 1 I	NA OH	25	1.2 X10-3 M	CD	EQUIV CONDCTNCE GRAPH	FLOC GRAH	53004	T L
3.4 E 0 I	OLEIC ACID	25	2.0 X10-3 M	DD	EQUIV CONDCTNCE GRAPH	FLOC GRAH	53004	K L
1.1 E 1	PH OF SOLUTION	5	8. X10-4 M	BD	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L
1.1 E 1	PH OF SOLUTION	25	1.0 X10-3 M	BD	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L
1.1 E 1	PH OF SOLUTION	40	1.6 X10-3 M	BD	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L
1.1 E 1	PH OF SOLUTION	60	2.3 X10-3 M	BD	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L
1.1 E 1	PH OF SOLUTION	90	3.3 X10-3 M	BD	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L
1.0 E-1 P	TRIBUTYL PHOSPHATE	24.7	4. X10-2 P	HE	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L
2.0 E-1 P	TRIBUTYL PHOSPHATE	24.7	5. X10-2 P	HD	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T L
30 ENTRIES FOR COMPOUND								
* COMPOUND NO = 264 MOL WGT -	304.4	SODIUM ELAIDATE /TRANS-9-OCTADECENOATE/						
	40	1.4 X10-3 M	DD	AVER SP EQUIV COND	TAMA SHIR	58007	P L	
	40	2.5 X10-3 M	DD	SPECFC CONDCTNCE GRAPH	TAMA SHIR	58007	P L	
	60	2.6 X10-3 M	DD	SPECFC CONDCTNCE GRAPH	TAMA SHIR	58007	P L	
	60			QUESTIONABLE CRITERION	TAMA NAKA	53001	R	
	UNK	1. X10-3 M	DG	FOTOMTR SPCTR CHNGE PNCN	TAMA SHIR	58007	T L	
	UNK	2.5 X10-3 M	DD	EQUIV CONDCTNCE GRAPH	TAMA SHIR	58007	T L	
6 ENTRIES FOR COMPOUND								
COMPOUND NO = 265 MOL WGT -	320.0	HEXADECYL TRIMETHYL AMMONIUM CHLORIDE						
	30	1.3 X10-3 M	BD	EQUIV CONDCTNCE GRAPH	RALS EGGE	47003	T L	
	UNK	1.5 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T L	
2 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
Compound No.	Mol. Wgt.							
COMPOUND NO = 266	MOL WGT -	350.0	HEXADECYL DIMETHYL 2-HYDROXYETHYL AMMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 267	MOL WGT -	380.1	HEXADECYL DI-/2-HYDROXYETHYL/METHYL AMMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 268	MOL WGT -	410.1	HEXADECYL TRI-/2-HYDROXYETHYL/ AMMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 269	MOL WGT -	380.1	HEXADECYL DIMETHYL 2,3-DIHYDROXYPROPYLAMMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 270	MOL WGT -	348.1	OCTADECYL TRIMETHYLAMMONIUM CHLORIDE					
4.74 E 0 H METHANOL 5 ENTRIES FOR COMPOUND								
25	3.4	X10-4 M	BD	EQUIV CONDCTNCE GRAPH	GRIE KRAU	48010	T L	
30	4.	X10-4 M	CE	EQUIV CONDCTNCE GRAPH	RALS EGGE	47003	T L	
30	3.	X10-4 M	CD	EQUIV CONDCTNCE GRAPH	RALS EGGE	47003	K L	
UNK	3.46	X10-4 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T L	
25	4.00	X10-4 M	BB	EQUIV CONDCTNCE GRAPH	GRIE KRAU	48010	P 3	
COMPOUND NO = 271	MOL WGT -	306.3	SODIUM TRI-ISO-PROPYL BENZENE SULFONATE					
2 ENTRIES FOR COMPOUND								
31	6.5	X10-2 M	FC	KARFT POINT SOLUBILITY	SHUC LING	49004	K L	
50	6.1	X10-2 M	FG	UNSPEC SPCTR CHNG PNCH	SHUC LING	49004	T L	
COMPOUND NO = 272	MOL WGT -	284.3	TRI-ISOPROPYL BENZENE SULFONIC ACID					
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 273	MOL WGT -	222.3	SODIUM DODECANOATE					
59016 42004								
17	2.7	X10-2 M	DC	EQUIV CONDCTNCE GRAPH	EKWA	27001	L L	
20	3.1	X10-2 M	XG	VISCOSITY MINIMUM	SATA TYUZ	53006	T L	
20				QUESTIONABLE CRITERION	HESS PHIL	39009	R	
20	2.84	X10-2 M	DB	EQUIV CONDCTNCE GRAPH	EKWA	42004	P L	
25	2.60	X10-2 M	HC	FOTOMTR SOLUBLZTN PDMA	GINN KINN	59009	T L	
25	2.64	X10-2 M	HC	FOTOMTR SOLUBLZTN OROT	GINN KINN	59009	T L	
25	2.6	X10-2 M	XG	VISUAL SPCTR CHNGE	KLEV	53010	T L	
25	2.77	X10-2 M	DB	EQUIV CONDCTNCE GRAPH	EKWA	42004	P L	
25	2.37	X10-2 M	DG	VISUAL SPCTR CHNGE PNCH	MERR GETT	48024	T L	
25	2.44	X10-2 M	DB	SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T 3	
25	2.30	X10-2 M	DG	SURFACE TENSION MINIMUM	CAMP LAKS	65024	T L	
30	2.72	X10-2 M	DB	EQUIV CONDCTNCE GRAPH	EKWA	42004	P L	
30	2.55	X10-2 M	DB	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	48016	T L	
30	2.53	X10-2 M	DB	FOTOMTR SOLUBLZTN OROT	KOLT STRI	48016	T L	
35	2.66	X10-2 M	DB	EQUIV CONDCTNCE GRAPH	EKWA	42004	P L	
35	2.50	X10-2 M	DB	SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T L	
40	2.64	X10-2 M	DB	EQUIV CONDCTNCE GRAPH	EKWA	42004	P L	
40	2.5	X10-2 M	DC	FOTOMTR SOLUBLZTN PDMA	KOLT JOHN	46006	T L	
45	2.69	X10-2 M	DB	EQUIV CONDCTNCE GRAPH	EKWA	42004	P L	
50	2.72	X10-2 M	DB	EQUIV CONDCTNCE GRAPH	EKWA	42004	P L	
50	2.55	X10-2 M	DB	FOTOMTR SOLUBLZTN PDMA	KOLT STRI	48016	T L	
50	2.3	X10-2 M	DC	FOTOMTR SOLUBLZTN PDMA	KOLT JOHN	46006	T L	
50	2.80	X10-2 M	DG	SURFACE TENSION MINIMUM	CAMP LAKS	65024	T L	
50	1.91	X10-2 M	DG	VISUAL SPCTR CHNGE PNCH	RAIS	52016	T L	
50	48016			VALUES FRM REF IN CMC	KOLT STRI	49005	R	
50	2.15	X10-2 M	HB	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L	
60	2.6	X10-2 M	DC	PH OR HYDROLYSIS	STAU	39006	G L	
60	2.6	X10-2 M	DG	SOLUBLZTN TOLUENE	DEMC DUMA	60032	T L	
60	5.6	X10-1 D	DG	SOLUBLZTN TOLUENE	DEMC	60034	T L	
	2.51	X10-2 M					M	
60	2.4	X10-2 M	DC	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	T L	
RM	2.7	X10-2 M	DC	ULTRAFILTRATION	EKWA	27001	L L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Concentration units: A = mol %; B = vol % solvent; C = mol % surfactant mixture; D = wt/vol %; E = % saturation; H = wt % solvent; I = mol % surfactant; K = normality

concentrations; M = molar; N = normal; P = wt %; Q = wt % surfactant; R = varied; S = mol/kg; T = wt % surfactant mixture; U = mol/(dm³ kg); W = molar; Y = atm. Details in page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
8.9 E-3 M SI02/NA20 = 1.60	25	1.72 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
1.52 E-2 M SI02/NA20 = 1.60	25	1.47 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
1.57 E-2 M SI02/NA20 = 1.60	25	1.51 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
2.05 E-2 M SI02/NA20 = 1.60	25	1.32 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
2.47 E-2 M SI02/NA20 = 1.60	25	1.19 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
3.34 E-2 M SI02/NA20 = 1.60	25	1.08 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
1.25 E-2 M SI02/NA20 = 1.60	60	1.8 X10-2 M	HD	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	T L	
2.5 E-2 M SI02/NA20 = 1.60	60	1.4 X10-2 M	HD	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	T L	
9.8 E-3 M SI02/NA20 = 2.46	25	1.96 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
2.09 E-2 M SI02/NA20 = 2.46	25	1.56 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
2.84 E-2 M SI02/NA20 = 2.46	25	1.42 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
3.46 E-2 M SI02/NA20 = 2.46	25	1.30 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
4.35 E-2 M SI02/NA20 = 2.46	25	1.09 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
5.82 E-2 M SI02/NA20 = 2.46	25	9.75 X10-3 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
2.5 E-2 M SI02/NA20 = 2.46	60	1.4 X10-2 M	HD	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	K L	
4.3 E-3 M SI02/NA20 = 3.93	25	2.18 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
8.3 E-3 M SI02/NA20 = 3.93	25	2.08 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
1.20 E-2 M SI02/NA20 = 3.93	25	2.00 X10-2 M	HG	VISUAL SPCTR CHNCE PNCN	MERR GETT	48024	T L	
2.5 E-2 M SI02/NA20 = 3.93	60	1.4 X10-2 M	HD	FOTOMTR SOLUBLZTN OROT	MERR GETT	48024	K L	
1.1 E 1 PH OF SOLUTION	24	2.3 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	T L	
1.1 E 1 PH OF SOLUTION	24	2.3 X10-2 M	BC	SURFACE TENSION LOG PLOT	HARV	56018	T 3	
1.1 E 1 PH OF SOLUTION	30	2.25 X10-2 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T 3	
1.1 E 1 PH OF SOLUTION	40	2.25 X10-2 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1 PH OF SOLUTION	50	2.25 X10-2 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1 PH OF SOLUTION	60	2.25 X10-2 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1 PH OF SOLUTION	90	2.25 X10-2 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
3.2 E-4 M 1,2 DECANE DIOL	24	2.11 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	1.86 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
7.0 E-4 M 1,2 DECANE DIOL	24	1.71 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	1.80 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
8.9 E-4 M 1,2 DECANE DIOL	24	1.85 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	1.92 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.20 E-3 M 1,2 DECANE DIOL	24	2.22 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.08 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.02 E-3 M 1,10 DECANE DIOL	24	1.85 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	1.85 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
2.02 E-3 M 1,10 DECANE DIOL	24	1.39 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.20 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
4.10 E-3 M 1,10 DECANE DIOL	24	2.17 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
4.6 E-5 M DECANOL-1	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
8.5 E-5 M DECANOL-1	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.05 E-4 M DECANOL-1	24	2.12 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.46 E-4 M DECANOL-1	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.61 E-4 M DECANOL-1	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.15 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.99 E-4 M DECANOL-1	24	2.12 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	2.00 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
7.7 E-3 M HEXANOL-1	24	1.88 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	1.78 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.12 E-2 M HEXANOL-1	24	1.24 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	1.55 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.58 E-2 M HEXANOL-1	24	1.02 X10-2 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	9.5 X10-3 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
4.02 E-2 M HEXANOL-1	24	8.8 X10-3 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
4.11 E-2 M HEXANOL-1	24	8.8 X10-3 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
4.97 E-2 M HEXANOL-1	24	8.8 X10-3 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	
1.1 E 1 PH OF SOLUTION	24	8.8 X10-3 M	BG	FOTOMTR SPCTR CHNCE PNCN	HARV	56018	G L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

	Additives	Temp. °C	CMC	Qual. Meth.	Method	Authors	Reference	Source	Evaluation
5.70 E-2 M	HEXANOL-1	24	7.1 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
5.85 E-2 M	HEXANOL-1	24	7.5 X10-3 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
1.13 E-3 M	HEPTANOL-1	24	2.13 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
2.02 E-3 M	HEPTANOL-1	24	2.06 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
3.01 E-3 M	HEPTANOL-1	24	2.00 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
6.14 E-3 M	HEPTANOL-1	24	1.68 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
7.98 E-3 M	HEPTANOL-1	24	1.50 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
9.98 E-3 M	HEPTANOL-1	24	1.40 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
1.13 E-2 M	HEPTANOL-1	24	1.45 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
1.24 E-2 M	HEPTANOL-1	24	1.55 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
6.4 E-4 M	OCTANOL-1	24	2.10 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
1.27 E-3 M	OCTANOL-1	24	1.90 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
1.60 E-3 M	OCTANOL-1	24	1.93 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
1.92 E-3 M	OCTANOL-1	24	1.90 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
2.30 E-3 M	OCTANOL-1	24	1.95 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
2.69 E-3 M	OCTANOL-1	24	1.95 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	HARV	56018	G L	
1.1 E 1	PH OF SOLUTION								
174 ENTRIES FOR COMPOUND									
COMPOUND NO = 274 MOL WGT -		340.0	HEXADECYL PYRIDINIUM CHLORIDE						
		13.0	8.4 X10-4 M	BC	KRAFFT POINT SOLUBILITY	ADDI FURM	56019	T L	
		18.5	8.5 X10-4 M	BC	INTERFACIAL TNSN UNSPEC	ADDI FURM	56019	T L	
		23	5.0 X10-4 M	HG	STREAMING CURRENT	CARD	66011	T L	
		25	9. X10-4 M	BD	EQUIV CONDCTNCE GRAPH	MALS HART	34001	K L	
		25	9.0 X10-4 M	BB	FOTOMTR SOLUBLZTN AZBZ	HART	38001	P 3	
		25	6.7 X10-4 M	BG	EQUIV COND MAX BEGINING	GRIE KRAU	49018	T L	
		25	3. X10-5 W	HG	VISUAL SPCTR CHNGE	FINE MCBA	48011	T L	
		80	2.2 X10-3 M	BG	EQUIV COND 1ST DEVIATION	HART	36002	T L	
		80	2.36 X10-3 M	BC	SPECFC CONDCTNCE GRAPH	HART	36002	P 3	
3.2 E-3 N	NA CL	25	4.0 X10-4 M	BD	FOTOMTR SOLUBLZTN AZBZ	HART	38001	P L	
1.0 E-2 N	NA CL	25	1.8 X10-4 M	BD	FOTOMTR SOLUBLZTN AZBZ	HART	38001	P L	
3.2 E-2 N	NA CL	25	8. X10-5 M	BE	FOTOMTR SOLUBLZTN AZBZ	HART	38001	P L	
7.43 E-3 M	NA CL	31	1.6 X10-4 M	BG	TURBIDITY PLT LITE SCATR	ANAC	58009	G L	
1.754E-2 M	NA CL	31	1.2 X10-4 M	BG	TURBIDITY PLT LITE SCATR	ANAC	58009	G L	
4.05 E-2 M	NA CL	31	8.3 X10-5 M	BG	TURBIDITY PLT LITE SCATR	ANAC	58009	G L	
3.08 E-1 M	NA CL	31	3.2 X10-5 M	BG	TURBIDITY PLT LITE SCATR	ANAC	58009	G L	
4.382E-1 M	NA CL	31	6.9 X10-5 M	BG	TURBIDITY PLT LITE SCATR	ANAC	58009	G L	
6.17 E-1 M	NA CL	31	3.3 X10-5 M	BG	TURBIDITY PLT LITE SCATR	ANAC	58009	G L	
7.30 E-1 M	NA CL	31	9.6 X10-6 M	BG	TURBIDITY PLT LITE SCATR	ANAC	58009	G L	
1.00 E 2 E	NITROBENZENE	25	5.8 X10-4 M	BA	EQUIV CONDCTNCE GRAPH	GRIE KRAU	49018	T 3	
20 ENTRIES FOR COMPOUND									
COMPOUND NO = 275 MOL WGT -		396.1	HEXADECYL DIMETHYLBENZYLAMMONIUM CHLORIDE						
		23	4.2 X10-5 M	HG	STREAMING CURRENT	CARD	66011	T L	
		25	2. X10-4 W	HG	VISUAL SPCTR CHNGE	FINE MCBA	48011	T L	
		UNK	4.4 X10-5 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T L	
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 276 MOL WGT -		596.8	NONAETHYLENE GLYCOL MONODODECANOATE						
			NATURAL DISTRIBUTION OF HEAD GROUPS						
1 ENTRIES FOR COMPOUND		0	6.3 X10-4 W	HD	FREEZING POINT	GONI MCBA	47007	T L	
COMPOUND NO = 277 MOL WGT -		349.6	BENZYL TRIMETHYL AMMONIUM DODECANOATE						
4. E O I C6H5 (CH3)3 N OH		25	1 X10-2 M	HE	EQUIV CONDCTNCE GRAPH	BRAD MCBA	48009	T L	
1 ENTRIES FOR COMPOUND									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives			Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 278 MOL WGT - 283.9 DODECYL PYRIDINIUM CHLORIDE									
			25	1.5 X10-2 M	XG	QUESTIONABLE CRITERION VISUAL SPCTR CHNGE	BROW ROBI	52013	R
			25	2.8 X10-3 M	HE	EQUIV CONDCTNCE GRAPH	KLEV	53010	T L
			25	1.47 X10-2 M	CC	TURBIDITY PLT LITE SCATR	BRAD MCBA	48009	T L
			25	1.46 X10-2 M	CB	SURFACE TENSION UNSPEC	FORD OTTE	66028	T 3
			25	1.4 X10-2 M	BC	SURFACE TENSION LOG PLOT	WEIN ZOGR	65026	T L
			30	2.0 X10-2 W	CC	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
			30	1.74 X10-2 M	CB	SPECFC CONDCTNCE GRAPH	MEGU KOND	59024	T L
			30	1.71 X10-2 M	CG	FOTOMTR SPCTR CHNGE EOSN	MEGU KOND	59024	T L
			30	1.5 X10-2 M	XC	REFRACTIVE INDEX	KLEV	53010	T L
			50	2.0 X10-2 W	CC	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
			50	1.51 X10-2 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.	E-2 M	K CL	25	1.13 X10-2 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T L
2.	E-2 M	K CL	25	1.22 X10-2 M	CB	SURFACE TENSION UNSPEC	FORD OTTE	66028	T L
5.	E-2 M	K CL	25	8.46 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T L
5.	E-2 M	K CL	25	8.60 X10-3 M	CB	SURFACE TENSION UNSPEC	FORD OTTE	66028	T L
8.	E-2 M	K CL	25	6.88 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T L
1.	E-1 M	K CL	25	2.3 X10-3 M	BC	SURFACE TENSION LOG PLOT	WEIN ZOGR	65026	T L
2.	E-3 N	K CNS	50	7.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-2 N	K CNS	50	3.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
3.	E-2 N	K CNS	50	2.4 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-2 N	NA BR	50	9.6 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
3.	E-2 N	NA BR	50	6.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-1 N	NA BR	50	2.8 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-1 N	NA BR	50	1.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-3 M	NA CL	50	1.33 X10-2 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-2 M	NA CL	50	1.16 X10-2 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.	E-2 M	NA CL	50	9.3 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-2 M	NA CL	50	6.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.0	E-1 M	NA CL	50	4.9 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
3.	E-1 M	NA CL	50	2.9 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.0	E 0 M	NA CL	50	1.4 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-2 N	NA I	50	6.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.5	E-2 N	NA I	50	4.0 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
9.	E-2 N	NA I	50	2.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-2 N	NA N03	50	9.6 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
3.	E-2 N	NA N03	50	5.6 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-1 N	NA N03	50	3.4 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-1 N	NA N03	50	1.9 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-3 N	NA4 P207 PYRO	50	1.01 X10-2 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.	E-3 N	NA4 P207 PYRO	50	7.5 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-3 N	NA4 P207 PYRO	50	6.0 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-2 N	NA4 P207 PYRO	50	4.7 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.	E-2 N	NA4 P207 PYRO	50	4.6 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-2 N	NA4 P207 PYRO	50	4.8 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.0	E-1 N	NA4 P207 PYRO	50	4.6 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.	E-1 N	NA4 P207 PYRO	50	4.0 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
4.	E-1 N	NA4 P207 PYRO	50	3.3 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.0	E 0 N	NA4 P207 PYRO	50	2.2 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.	E-3 N	NA2 S04	50	1.11 X10-2 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-3 N	NA2 S04	50	8.6 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.	E-2 N	NA2 S04	50	7.8 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
2.	E-2 N	NA2 S04	50	7.1 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
3.	E-2 N	NA2 S04	50	6.7 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
5.	E-2 N	NA2 S04	50	5.9 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.0	E-1 N	NA2 S04	50	5.4 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
3.0	E-1 N	NA2 S04	50	3.8 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
1.0	E 0 N	NA2 S04	50	1.8 X10-3 M	CG	VISUAL SPCTR CHNGE EOSN	LANG	51005	G L
58 ENTRIES FOR COMPOUND									
COMPOUND NO = 279 MOL WGT - 340.0 DODECYL BENZYL DIMETHYLLAMMONIUM CHLORIDE									
			25	2.3 X10-3 M	HC	QUESTIONABLE CRITERION	YANG FOST	53015	R
			25.0	7.8 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	BRAD MCBA	48009	T L
			UNK	2.8 X10-3 M	PG	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3
			UNK	8.1 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	ROSS KWAN	53007	T L
5.	E 2 Y	PRESSURE	25.0	8.25 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	CELL EGGE	52001	T L
1.	E 3 Y	PRESSURE	25.0	8.5 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3
2.	E 3 Y	PRESSURE	25.0	8.2 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3
3.	E 3 Y	PRESSURE	25.0	7.5 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3
4.	E 3 Y	PRESSURE	25.0	7.1 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3
5.	E 3 Y	PRESSURE	25.0	6.8 X10-3 M	CB	SPECFC CONDCTNCE GRAPH	OSUG SATO	65036	G 3
11 ENTRIES FOR COMPOUND									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 280 MOL WGT -		465.8	DECYL TRIMETHYL AMMONIUM DODECYL SULFATE					
	25	1.9 X10-4 M	CE UNSPEC SOLUBLZTN SDN 4		HOYE MARM	61001	TL L	
	25	2.1 X10-4 M	DC SPECFC CONDCTNCE GRAPH		PACK DONB	63030	T L	
	UNK	1.85 X10-3 D	BD DEBYE PLT LIGHT SCATTER		HOYE DOER	64007	T L	
		3.971X10-5 M					M	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 281 MOL WGT -		493.8	DODECYL TRIMETHYLAMMONIUM DODECYL SULFATE					
	25	3. X10-5 M	CE SURFACE TNSN LINEAR PLOT		HOYE MARM	61001	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 282 MOL WGT -		506.8	HEXADECYL /OXYETHYLENE/ 6 ALCOHOL					
HOMOGENEOUS HEAD GROUP								
	20	9.9 X10-5 M	BC SURFACE TENSION LOG PLOT		CARL CHAL	64009	T L	
	25	1.0 X10-6 M	BD SURFACE TENSION LOG PLOT		GOOD OTTE	61004	T P	
	25	1.66 X10-6 M	BC SURFACE TENSION LOG PLOT		ELWO MACF	62027	T P	
	27	9.4 X10-5 M	BG FOTOMTR SPCTR CHNG I2		CARL CHAL	64009	T L	
	28	5. X10-5 D	BE SURFACE TENSION LOG PLOT		CORK GOOD	64023	T L	
		9.8 X10-7 M					M	
5 ENTRIES FOR COMPOUND								
COMPOUND NO = 283 MOL WGT -		399.7	HEXANOLAMINE-CH3CH/OH/CH2C/CH3/2NH2-ELAIDATE					
	0	6. X10-4 W	DD FREEZING POINT		GONI MCBA	46016	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 284 MOL WGT -		261.4	HEXANOLAMINE-CH3CH/OH/CH2C/CH3/2NH2-OCTANOATE					
	0	3.1 X10-1 M	DC FREEZING POINT		GONI	46019	K L	
	20	2.6 X10-1 M	XG VISCOSITY MINIMUM		SATA TYUZ	53006	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 285 MOL WGT -		399.7	HEXANOLAMINE-CH3CH/OH/CH2C/CH3/2NH2-OLEATE					
	0	6.0 X10-4 W	DD FREEZING POINT		GONI MCBA	46008	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 286 MOL WGT -		439.6	AMMONIUM DI-2-ETHYL-HEXYL SULFOSUCCINATE					
	0	4. X10-3 W	CE FREEZING POINT		MCBA BRAD	43006	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 287 MOL WGT -		365.6	OCTYL TRIMETHYLAMMONIUM OCTANE SULFONATE					
	25	2.1 X10-2 M	BC METHOD NOT CITED		CORK GOOD	66014	T L	
	25	2.016X10-2 M	CA SPECFC CONDCTNCE GRAPH		TART LING	43004	T 3	
	RM	1.99 X10-2 M	BB DEBYE PLT LIGHT SCATTER		ANAC	53002	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 288 MOL WGT -		421.7	DECYL TRIMETHYLAMMONIUM DECANESULFONATE					
	25	1.36 X10-3 M	BC METHOD NOT CITED		CORK GOOD	66014	T 3	
	40	1.40 X10-3 M	CA SPECFC CONDCTNCE GRAPH		TART LING	43004	P 3	
	RM	1.3 X10-3 M	BD TURBIDITY PLT LITE SCATR		ANAC	53002	KC L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 289 MOL WGT -		478.7	TETRADECYL/OXYETHYLENE/6 ALCOHOL					
HOMOGENEOUS HEAD GROUP								
	25	5. X10-4 D	BE SURFACE TENSION LOG PLOT		CORK GOOD	64023	T L	
		1.0 X10-5 M					M	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 290 MOL WGT -		328.3	DODECYL PYRIDINIUM BROMIDE					
	5	1.15 X10-2 W	BA SPECFC CONDCTNCE GRAPH		ADDE TAYL	64050	T 3	
	10	1.12 X10-2 W	BA SPECFC CONDCTNCE GRAPH		ADDE TAYL	64050	T 2	
	15	1.10 X10-2 W	BA SPECFC CONDCTNCE GRAPH		ADDE TAYL	64050	T 2	
	20	1.12 X10-2 W	BA SPECFC CONDCTNCE GRAPH		ADDE TAYL	64050	T 2	
	25	1.14 X10-2 W	BA SPECFC CONDCTNCE GRAPH		ADDE TAYL	64050	T 1	
	25	1.6 X10-2 M	XG VISUAL SPCTR CHNG		KLEV	53010	T L	
	25	1.16 X10-2 M	CC TURBIDITY PLT LITE SCATR		FORD OTTE	66028	T L	
	25	1.20 X10-2 M	CB SURFACE TENSION UNSPEC		FORD OTTE	66028	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2. E-2 M K BR	25	1.13 X10-2 M	CA	SPECFC CONDCTNCE GRAPH	BENT SPAR	66038	T	1
2. E-2 M K BR	25	1.21 X10-2 M	CB	MICELLAR SPECTRAL CHANGE	MUKE RAY	66006	T	L
4. E-2 M K BR	30	1.21 X10-2 M	BB	SURFACE TENSION LOG PLOT	VENA NAUM	64001	T	L
5. E-2 M K BR	30	1.18 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	2
6. E-2 M K BR	30	1.25 X10-2 M	CA	SPECFC CONDCTNCE GRAPH	MEGU KOND	59024	T	L
8. E-2 M K BR	30	1.25 X10-2 M	CG	FOTOMTR SPCTR CHNGE EOSN	MEGU KOND	59024	T	L
1.0 E-1 M K BR	35	1.22 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	2
3.0 E-2 M K BR	40	1.28 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	3
3.0 E-2 M K BR	45	1.35 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	3
3.0 E-2 M K BR	50	1.40 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	3
3.0 E-2 M K BR	55	1.48 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	3
3.0 E-2 M K BR	60	1.54 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	3
3.0 E-2 M K BR	65	1.63 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	3
3.0 E-2 M K BR	70	1.72 X10-2 W	BA	SPECFC CONDCTNCE GRAPH	ADDE TAYL	64050	T	3
2. E-2 M K BR	25	7.20 X10-3 M	CB	SURFACE TENSION UNSPEC	FORD OTTE	66028	T	L
2. E-2 M K BR	25	7.32 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T	3
4. E-2 M K BR	25	4.88 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T	3
5. E-2 M K BR	25	4.70 X10-3 M	CB	SURFACE TENSION UNSPEC	FORD OTTE	66028	T	L
6. E-2 M K BR	25	3.96 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T	3
8. E-2 M K BR	25	3.36 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T	3
1.0 E-1 M K BR	25	2.74 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T	L
3.0 E-2 M K BR	30	6.5 X10-3 M	CG	FOTOMTR SPCTR CHNGE EOSN	MEGU KOND	59024	T	L
3.0 E-2 M K I	30	3.5 X10-3 M	CG	FOTOMTR SPCTR CHNGE EOSN	MEGU KOND	59024	T	L
3.0 E-2 M K IO3	30	1.04 X10-2 M	CG	FOTOMTR SPCTR CHNGE EOSN	MEGU KOND	59024	T	L
3.0 E-2 M K2 SO4	30	7.9 X10-3 M	CG	FOTOMTR SPCTR CHNGE EOSN	MEGU KOND	59024	T	L
6. E-2 M LI BR	25	3.96 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T	3
5. E-2 M NA BR	30	3.66 X10-3 M	BB	SURFACE TENSION LOG PLOT	VENA NAUM	64001	TA	L
3.0 E-2 M NA CL	30	9.1 X10-3 M	CG	FOTOMTR SPCTR CHNGE EOSN	MEGU KOND	59024	T	L
6. E-2 M RB BR	25	3.35 X10-3 M	CC	TURBIDITY PLT LITE SCATR	FORD OTTE	66028	T	3
E 0 0427				GRAPH DATA NOT RETRIEVED	LANG	53005	R	
38 ENTRIES FOR COMPOUND								
COMPOUND NO = 291 MOL WGT -	420.6	TETRADECYL TRIPROPYLAMMONIUM BROMIDE						
5. E-2 M NA BR	30	2.05 X10-3 M	BB	SURFACE TENSION LOG PLOT	VENA NAUM	64001	T	L
2 ENTRIES FOR COMPOUND	30	2.76 X10-4 M	BB	SURFACE TENSION LOG PLOT	VENA NAUM	64001	TA	L
COMPOUND NO = 292 MOL WGT -	448.7	HEXADECYL TRIPROPYLAMMONIUM BROMIDE						
1 ENTRIES FOR COMPOUND	30	5.7 X10-4 M	BC	SURFACE TENSION LOG PLOT	VENA NAUM	64001	T	L
COMPOUND NO = 293 MOL WGT -	310.3	DODECYL DIMETHYL AMINE OXIDE HYDROBROMIDE						
1. E-3 M H BR	27	4. X10-2 D	BD	TURBIDITY PLT LITE SCATR	HERR	64006	T	L
2. E-1 M NA BR		1.2 X10-3 M					M	
1. E-3 M H BR	50	7. X10-2 D	BD	TURBIDITY PLT LITE SCATR	HERR	64006	T	L
2. E-1 M NA BR		2.2 X10-3 M					M	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 294 MOL WGT -	366.5	HEXYL/OXYETHYLENE/6 ALCOHOL HOMOGENEOUS HEAD GROUP						
	20	7.4 X10-2 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T	L
	20	7.5 X10-2 M	BE	SURFACE TENSION LOG PLOT	MULL METC	62015	T	L
	25	1.18 X10-1 M	BC	VAPR PRESURE LOWERING	CORK GOOD	64027	L	L
	30	6.5 X10-2 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T	L
	40	5.2 X10-2 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T	L
6 ENTRIES FOR COMPOUND	45	7.8 X10-2 M	BC	VAPR PRESURE LOWERING	CORK GOOD	64027	L	L
COMPOUND NO = 295 MOL WGT -	246.2	SODIUM NONYL 1-SULFATE						
	20	6.5 X10-2 M	BB	INTERFACIAL TENSION LOGM	V VO	60025	T	L
	21	1.66 X10 0 D	BD	REFRACTIVE INDEX	HUIS	64047	T	L
		6.742X10-2 M					M	
	21	1.59 X10 0 D	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T	3
		6.458X10-2 M					M	
	25	1.35 X10 0 D	DB	REFRACTIVE INDEX	PRIN HERM	56011	T	L
		5.483X10-2 M					M	
	25	1.45 X10 0 D	DB	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T	L
		5.889X10-2 M					M	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

		Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.	E-1 K	NA CL	20	4.3 X10-2 M	BB	INTERFACIAL TENSION LOGM	V VO	60025	T L	
3.	E-2 M	NA CL	21	1.30 X10 0 D 5.280X10-2 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3 M	
1.	E-1 M	NA CL	21	1.00 X10 0 D 4.061X10-2 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3 M	
1.	E-1 M	NA CL	21	1.05 X10 0 D 4.264X10-2 M	BD	REFRACTIVE INDEX	HUIS	64047	T L M	
3.	E-1 M	NA CL	21	6.3 X10-1 D 2.55 X10-2 M	BB	TURBIDITY PLT LITE SCATR	HUIS	64047	T 3 M	
7.5	E-2 M	NA CL	25	1.2 X10 0 D 4.87 X10-2 M	DC	TURBIDITY PLT LITE SCATR	PRIN HERM	56011	T L M	
11 ENTRIES FOR COMPOUND										
COMPOUND NO = 296 MOL WGT - 168.2 POTASSIUM HEPTANOATE										
2.	E O I	K OH	25	7.5 X10-1 M	BG	UNSPEC SPCTR CHNG PNCN	KLEV	58011	T L	
1.	E-5 N	PINACYANOL CL (DYE)	25	8.0 X10-1 M	BG	VISUAL SPCTR CHNG PNCN	SHIN	54005	G L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	4.5 X10-1 M	DG	FOTOMTR SPCTR CHNG PNCN	HERZ	52015	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	6.70 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52015	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	6.70 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	7.80 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52015	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	5.5 X10-1 M	DG	FOTOMTR SPCTR CHNG PNCN	HERZ	52015	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	7.80 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
1.	E-4 N	PINACYANOL CL (DYE)	25	7.80 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52015	T L	
	0188		25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
	0188		25			SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
2.5	E-5 N	PINACYANOL CL (DYE)	25	6.62 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
6.6	E-2 M	K CL	25	6.17 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	5.77 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL FL (DYE)	25	4.07 E-1 M	K CL	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	5.43 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	5.22 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	4.85 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	4.30 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
1.08	E 0 M	K CL	25	3.74 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	1.50 E 0 M	K CL	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	3.35 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
1.79	E 0 M	K CL	25	3.07 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	2.85 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.28	E 0 M	K CL	25	2.63 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	2.47 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.64	E 0 M	K CL	25	2.30 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.5	E-5 N	PINACYANOL CL (DYE)	25	2.18 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
2.91	E 0 M	K CL	25	2.18 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	7.28 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
1.46	E-1 M	K CL	25	6.91 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	6.33 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
6.33	E-1 M	K CL	25	5.42 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	1.08 E 0 M	K CL	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	4.30 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
1.72	E 0 M	K CL	25	3.40 X10-1 M	DG	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	2.38 E 0 M	K CL	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
5.	E-5 N	PINACYANOL CL (DYE)	25	2.80 E 0 M	K CL	VISUAL SPCTR CHNG PNCN	HERZ	52017	T L	
1.07	E 1 C	0044	25	7.0 X10-1 M	BG	VISUAL SPCTR CHNG PNCN	SHIN	54005	G L	
2.	E O I	K OH	25	6.6 X10-1 M	BG	VISUAL SPCTR CHNG PNCN	SHIN	54005	G L	
1.57	E 1 C	0044	25	6.1 X10-1 M	BG	VISUAL SPCTR CHNG PNCN	SHIN	54005	G L	
2.	E O I	K OH	25	6.1 X10-1 M	BG	VISUAL SPCTR CHNG PNCN	SHIN	54005	G L	
2.	E O I	K OH	25							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
3.36 E 1 C 0044	25	5.6 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
4.36 E 1 C 0044	25	5.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
5.53 E 1 C 0044	25	4.9 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
6.66 E 1 C 0044	25	4.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
8.18 E 1 C 0044	25	4.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
2.1 E O C 0090	25	5.9 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
6.0 E O C 0090	25	4.4 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
1.40 E 1 C 0090	25	3.1 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
2.88 E 1 C 0090	25	2.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
5.41 E 1 C 0090	25	1.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
1.2 E O C 0091	25	3.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
7.3 E O C 0091	25	1.4 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
1.94 E 1 C 0091	25	8.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
3.97 E 1 C 0091	25	5.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
7.02 E 1 C 0091	25	3.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
E O 0092	25			GRAPH DATA NOT RETRIEVED	SHIN	54005	R
2. E O I K OH							
1.5 E O C 0092	25	8.0 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
5.4 E O C 0092	25	5.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
1.42 E 1 C 0092	25	2.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
3.34 E 1 C 0092	25	1.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
8. E-1 C 0297	25	5.3 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
3.1 E O C 0297	25	3.6 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
8.1 E O C 0297	25	2.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
1.75 E 1 C 0297	25	1.4 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
4.05 E 1 C 0297	25	9.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	54005	G L
2. E O I K OH							
0188	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X

62 ENTRIES FOR COMPOUND

COMPOUND NO = 297 MOL WGT -

224.4 POTASSIUM UNDECANOATE

4.8 E-3 M K CL	25	4.92 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52015	T L
1.44 E-2 M K CL	25	5.0 X10-2 M	BG	UNSPEC SPCTR CHNG PNCN	KLEV	58011	T L
1.87 E-2 M K CL	25	4.92 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
2.73 E-2 M K CL	25	4.56 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
3.86 E-2 M K CL	25	4.37 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
5.34 E-2 M K CL	25	4.09 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
7.64 E-2 M K CL	25	3.86 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
1.03 E-1 M K CL	25	3.56 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
1.33 E-1 M K CL	25	3.27 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
1.69 E-1 M K CL	25	2.94 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
2.16 E-1 M K CL	25	2.66 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
2.87 E-1 M K CL	25	2.42 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
4.03 E-1 M K CL	25	2.16 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
5.06 E-1 M K CL	25	1.91 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
6.42 E-1 M K CL	25	1.61 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
8.30 E-1 M K CL	25	1.45 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
1.01 E 0 M K CL	25	1.28 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
1.18 E 0 M K CL	25	1.11 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
	25	1.01 X10-2 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L
	25	9.4 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality
counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/
kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.35 E 0 M K CL	25	9.0 X10-3 M	DG	VISUAL SPCTR CHNCE PNCN	HERZ	52017	T L	
1.52 E 0 M K CL	25	8.7 X10-3 M	DG	VISUAL SPCTR CHNCE PNCN	HERZ	52017	T L	
1.70 E 0 M K CL	25	8.5 X10-3 M	DG	VISUAL SPCTR CHNCE PNCN	HERZ	52017	T L	
2.01 E 0 M K CL	25	8.1 X10-3 M	DG	VISUAL SPCTR CHNCE PNCN	HERZ	52017	T L	
2.22 E 0 M K CL	25	7.4 X10-3 M	DG	VISUAL SPCTR CHNCE PNCN	HERZ	52017	T L	
2. E 0 I K OH	25	4.9 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54005	G L	
0044	25			SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
0090	25			SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
0188	25			SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
4.9 E 0 C 0091	25	4.5 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	G L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
1.93 E 1 C 0091	25	4.0 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	G L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
2.76 E 1 C 0091	25	3.8 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	G L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
3.38 E 1 C 0091	25	3.6 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	GL L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
4.63 E 1 C 0091	25	3.4 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	G L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
5.76 E 1 C 0091	25	3.2 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	G L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
6.99 E 1 C 0091	25	3.0 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	G L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
8.38 E 1 C 0091	25	2.9 X10-2 M	BG	VISUAL SPCTR CHNCE PNCN	SHIN	54003	G L	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54005	X	
0296	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
2. E 0 I K OH	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
0090	25			SEE CMPD NMBR IN ADDITV	SHIN	54003	X	
40 ENTRIES FOR COMPOUND								
COMPOUND NO = 298 MOL WGT -		250.3 SODIUM TETRADECANOATE						
32001				VALUES FRM REF IN CMC	EKWA	40003	R	
21 6. X10-3 M DD				EQUIV CONDCTNCE GRAPH	EKWA	27001	L L	
21 6. X10-3 M DE				PH OR HYDROLYSIS	EKWA	27001	L L	
25 6.9 X10-3 M DB				SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T 3	
25 6.9 X10-3 M DG				SURFACE TENSION MINIMUM	CAMP LAKS	65024	T L	
35 7. X10-3 M DD				EQUIV CONDCTNCE GRAPH	EKWA	32001	K L	
35 6.95 X10-3 M DB				SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T 3	
48 7. X10-3 M DD				EQUIV CONDCTNCE GRAPH	EKWA	32001	K L	
50 7.5 X10-3 M HB				SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L	
50 7.1 X10-3 M DG				SURFACE TENSION MINIMUM	CAMP LAKS	65024	T L	
58 7. X10-3 M DD				EQUIV CONDCTNCE GRAPH	EKWA	32001	K L	
60 1.00 X10-2 M DC				PH OR HYDROLYSIS	STAU	39006	G L	
65 7. X10-3 M DD				EQUIV CONDCTNCE GRAPH	EKWA	32001	K L	
80 9. X10-3 M DD				EQUIV CONDCTNCE GRAPH	EKWA	32001	K L	
UNK 2.1 X10-1 D XG				VISUAL SPCTR CHNCE PNCN	DEMC	61031	T L	
8.38 X10-3 M							M	
1. E-1 H NA OH	70	2.9 X10-3 M CC		SURFACE TENSION LOG PLOT	POWN ADDI	38006	T L	
1. E-1 H NA OH	70	2.9 X10-3 M CC		INTERFACIAL TENSION LOGM	POWN ADDI	38006	T L	
1.1 E 1 PH OF SOLUTION	40	4.4 X10-3 M BD		SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1 PH OF SOLUTION	50	4.4 X10-3 M BD		SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1 PH OF SOLUTION	60	4.4 X10-3 M BD		SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1 PH OF SOLUTION	90	4.4 X10-3 M BD		SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
21 ENTRIES FOR COMPOUND								
COMPOUND NO = 299 MOL WGT -		194.2 SODIUM DECANOATE						
41003				VALUES FRM REF IN CMC	EKWA	40003	R	
20 1.0 X10-1 M XG				VISCOSITY MINIMUM	SATA TYUZ	53006	T L	
20 1.24 X10-1 M DE				VISCOSITY	HESS PHIL	39009	T L	
20 9.5 X10-2 M DE				PH OR HYDROLYSIS	EKWA	41003	T L	
20 9.7 X10-2 M DC				EQUIV CONDCTNCE GRAPH	EKWA	41003	T L	
25 9.40 X10-2 M DB				SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T 3	
25 9.55 X10-2 M DB				SURFACE TENSION LOG PLOT	CAMP LAKS	65024	T L	
30 1.06 X10-1 M DB				FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L	
30 1.1 X10-1 M DD				FOTOMTR SOLUBLZTN PDMAB	KOLT JOHN	46006	T L	
35 9.80 X10-2 M DB				SPECFC CONDCTNCE GRAPH	CAMP LAKS	65024	T 3	
50 1.05 X10-1 M DB				FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L	
50 1.05 X10-1 M DB				FOTOMTR SOLUBLZTN PDMAB	KOLT JOHN	46006	T L	
50 1.060 X10-1 M DB				SURFACE TENSION LOG PLOT	CAMP LAKS	65024	T L	
UNK 1.0 X10-1 M DE				ELECTROMOTIVE FORCE	CARR JOHN	47013	T L	
UNK 1.05 X10-1 M DB				FOTOMTR SOLUBLZTN PDMAB	CARR JOHN	47013	T L	
UNK 2.00 X10 0 D XG				VISUAL SPCTR CHNCE PNCN	DEMC	61031	T L	
1.029 X10-1 M							M	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
1.1 E 1	PH OF SOLUTION	5	1.03 X10-1 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1	PH OF SOLUTION	20	1.03 X10-1 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1	PH OF SOLUTION	60	1.03 X10-1 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
1.1 E 1	PH OF SOLUTION	90	1.03 X10-1 M	BC	SPECFC CONDCTNCE GRAPH	MARK TSIK	64051	T L	
20 ENTRIES FOR COMPOUND									
COMPOUND NO = 300 MOL WGT -		278.4	SODIUM HEXADECANOATE						
		42004		VALUES FRM REF IN CMC		EKWA	40003	R	
		50	2.1 X10-3 M	HC	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L	
		52	3.2 X10-3 M	DD	EQUIV CONDCTNCE GRAPH	EKWA	42004	K L	
		58	3.3 X10-3 M	DD	EQUIV CONDCTNCE GRAPH	EKWA	42004	K L	
		60	3.0 X10-3 M	DC	PH OR HYDROLYSIS	STAU	39006	G L	
		60	2.5 X10-3 M	DC	PH OR HYDROLYSIS	STAU	39006	K L	
		60	4. X10-4 M	DE	FOTOMTR SOLUBLZTN OROT		48024	T L	
		67	3.3 X10-3 M	DD	EQUIV CONDCTNCE GRAPH	EKWA	42004	K L	
		UNK	9. X10-2 D	XG	VISUAL SPCTR CHNGE PNCC	DEMCO	61031	T L	
			3.2 X10-3 M					M	
1. E-1 H	NA OH	70	3.9 X10-4 M	CC	SURFACE TENSION LOG PLOT	POWN ADDI	38006	K L	
1. E-1 H	NA OH	70	6.5 X10-4 M	CC	INTERFACIAL TENSION LOGM	POWN ADDI	38006	K L	
11 ENTRIES FOR COMPOUND									
COMPOUND NO = 301 MOL WGT -		348.4	SODIUM 3-N-DODECYL BENZENE SULFONATE						
		25	1.46 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T 3	
		30	1.46 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T L	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 302 MOL WGT -		348.4	SODIUM 4-N-DODECYL BENZENE SULFONATE						
		25	1.59 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T 3	
		30	1.59 X10-3 M	FA	SPECFC CONDCTNCE GRAPH	LUDL	56005	T L	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 303 MOL WGT -		245.4	DI-ISOPROPYLAMMONIUM CAPRYLATE						
		0	4.3 X10-1 M	DC	FREEZING POINT	GONI	46019	K L	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 304 MOL WGT -		352.6	CATOL 607 (N-/LAURYL-COLAMINO-FORMYL-METHYL/-PYRIDINIUM CHLORIDE)						
		0	4.8 X10-3 W	DC	FREEZING POINT				
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 305 MOL WGT -		320.6	POTASSIUM OLEATE/ CIS-9-OCTADECENOATE/						
		25	8. X10-4 M	XE	REFRACTIVE INDEX	KLEV	53010	T L	
		25	9.5 X10-4 M	XG	VISUAL SPCTR CHNGE	KLEV	53010	T L	
		25.8	1.0 X10-3 M	DG	FOTOMTR SPCTR CHNGE PNCC	CORR KLEV	46010	T L	
		30	6. X10-4 M	DE	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L	
		30	48016		VALUES FRM REF IN CMC	KOLT STRI	49005	R	
		50	1.2 X10-3 M	XE	REFRACTIVE INDEX	KLEV	53010	T L	
		50	1.3 X10-3 M	XG	VISUAL SPCTR CHNGE	KLEV	53010	T L	
		50	1.1 X10-3 M	DD	FOTOMTR SOLUBLZTN PDMAB	KOLT STRI	48016	T L	
8 ENTRIES FOR COMPOUND									
COMPOUND NO = 306 MOL WGT -		496.9	DECYL TRIMETHYLAMMONIUM SULFATE						
		UNK	5.03 X10-2 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 307 MOL WGT -		553.0	DODECYL TRIMETHYLAMMONIUM SULFATE						
		25	1.64 X10-2 N	BA	EQUIV CONDCTNCE GRAPH	VOEK TART	55006	T L	
		UNK	9.3 X10-3 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L	
		UNK			QUESTIONABLE CRITERION	WASI HUBB	64043	R	
E 0 NA2 SO4 3 ENTRIES FOR COMPOUND									
COMPOUND NO = 308 MOL WGT -		609.1	TETRADECYL TRIMETHYLAMMONIUM SULFATE						
		UNK	1.5 X10-3 M	BD	METHOD NOT CITED	WASI HUBB	64043	T L	
6.7 E-3 M	LA2 (SO4)3	UNK	8.7 X10-4 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L	
2. E-2 M	MG SO4	UNK	8.9 X10-4 M	BC	METHOD NOT CITED	WASI HUBB	64043	T L	
E 0	NA2 SO4	UNK			QUESTIONABLE CRITERION	WASI HUBB	64043	R	
4 ENTRIES FOR COMPOUND									

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; L – molar; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 317 MOL WGT - 647.0 T-OCTYL BENZENE/OXYETHYLENE/10 ALCOHOL								
REDUCED POLYDISPERSITY OF HEAD GROUPS								
25.0 1.00 X10-4 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
55.0 6.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 318 MOL WGT - 1,087.6 T-OCTYL BENZENE/OXYETHYLENE/20 ALCOHOL								
REDUCED POLYDISPERSITY OF HEAD GROUPS								
25.0 1.30 X10-4 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
55.0 7.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 319 MOL WGT - 1,528.2 T-OCTYL BENZENE/OXYETHYLENE/30 ALCOHOL								
REDUCED POLYDISPERSITY OF HEAD GROUPS								
25.0 1.60 X10-4 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
55.0 9.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 320 MOL WGT - 661.0 NYLON BENZENE/OXYETHYLENE/10 ALCOHOL								
BRANCHED CHAIN, REDUCED OE DISTRIBUTION								
62019 VALUES FRM REF IN CMC SCHI GILB 65011 R								
62019 VALUES FRM REF IN CMC SCHI ATLA 62020 R								
25.0 7.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
55.0 4.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
1.5 E O M DIOXANE 25.0 1.00 X10-4 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
3.0 E O M DIOXANE 25.0 1.80 X10-4 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
3. E O M UREA 25.0 1.00 X10-4 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
6. E O M UREA 25.0 2.40 X10-4 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
3. E O M GUANIDINIUM CL 25.0 1.40 X10-4 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
4.6 E O PH OF SOLUTION 25.0 2.50 X10-4 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
5. E O M GUANIDINIUM CL 25.0 7.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
4.6 F O PH OF SOLUTION 25.0 7.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
E O H CL 25.0 7.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI GILB 65011 T L								
4.6 E O PH OF SOLUTION 11 ENTRIES FOR COMPOUND								
COMPOUND NO = 321 MOL WGT - 881.3 NYLON BENZENE/OXYETHYLENE/15 ALCOHOL								
BRANCHED CHAIN, REDUCED OE DISTRIBUTION								
62019 VALUES FRM REF IN CMC SCHI ATLA 62020 R								
25.0 1.10 X10-4 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
55.0 4.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N BA CL2 25.0 5.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N CA CL2 25.0 5.7 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N CA NO3 25.0 7.6 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M H CL 25.0 1.50 X10-4 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M K CL 25.0 5.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M K NO3 25.0 5.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M LI CL 25.0 6.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M LI NO3 25.0 8.7 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N MG CL2 25.0 5.3 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N MG (NO3)2 25.0 7.6 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M NA BR 25.0 8.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M NA BRO3 25.0 5.2 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N NA CITRATE 25.0 2.5 X10-5 M ED SURFACE TENSION LOG PLOT SCHI 62019 T L								
4.3 E-1 M NA CL 25.0 6.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 L L								
8.6 E-1 M NA CL 25.0 5.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
1.29 E O M NA CL 25.0 4.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 L L								
8.6 E-1 M NA F 25.0 3.2 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M NA I 25.0 8.3 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M NA NO3 25.0 8.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M NA OH 25.0 8.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N NA2 SO4 25.0 2.3 X10-5 M ED SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M NA CNS 25.0 8.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 N SR CL2 25.0 5.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
8.6 E-1 M (CH3)4 N CL 25.0 4.5 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
5. E-1 M UREA 25.0 9.0 X10-5 M EC SURFACE TENSION LOG PLOT SCHI ATLA 62020 T L								
8.6 E-1 N NA2 SO4 25.0 4.2 X10-5 M EC SURFACE TENSION LOG PLOT SCHI 62019 T L								
4.3 E-1 M NA CNS 29 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 322 MOL WGT - 1.101.6 NONYL BENZENE/OXYETHYLENE/20 ALCOHOL									
BRANCHED CHAIN, REDUCED OE DISTRIBUTION									
1.5 E 0 M DIOXANE									
25.0 1.40 X10-4 M EC SURFACE TENSION LOG PLOT									
55.0 6.0 X10-5 M EC SURFACE TENSION LOG PLOT									
3.0 E 0 M DIOXANE									
25.0 3.90 X10-4 M EC SURFACE TENSION LOG PLOT									
3. E 0 M UREA									
25.0 1.80 X10-4 M EC SURFACE TENSION LOG PLOT									
6. E 0 M UREA									
25.0 4.75 X10-4 M EC SURFACE TENSION LOG PLOT									
3. E 0 M GUANIDINIUM CL									
25.0 1.80 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
5. E 0 M GUANIDINIUM CL									
25.0 5.60 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
E 0 H CL									
25.0 1.40 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
11 ENTRIES FOR COMPOUND									
COMPOUND NO = 323 MOL WGT - 1.542.2 NONYL BENZENE/OXYETHYLENE/30 ALCOHOL									
BRANCHED CHAIN, REDUCED OE DISTRIBUTION									
1.5 E 0 M DIOXANE									
25.0 1.85 X10-4 M EC SURFACE TENSION LOG PLOT									
55.0 8.0 X10-5 M EC SURFACE TENSION LOG PLOT									
3.0 E 0 M DIOXANE									
25.0 5.70 X10-4 M EC SURFACE TENSION LOG PLOT									
3. E 0 M UREA									
25.0 3.50 X10-4 M EC SURFACE TENSION LOG PLOT									
6. E 0 M UREA									
25.0 7.40 X10-4 M EC SURFACE TENSION LOG PLOT									
3. E 0 M GUANIDINIUM CL									
25.0 4.25 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
5. E 0 M GUANIDINIUM CL									
25.0 9.60 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
E 0 H CL									
25.0 1.85 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
11 ENTRIES FOR COMPOUND									
COMPOUND NO = 324 MOL WGT - 2.423.4 NONYL BENZENE/OXYETHYLENE/50 ALCOHOL									
BRANCHED CHAIN, REDUCED OE DISTRIBUTION									
62019 VALUES FRM REF IN CMC									
62019 VALUES FRM REF IN CMC									
25.0 1.85 X10-4 M EC SURFACE TENSION LOG PLOT									
55.0 8.0 X10-5 M EC SURFACE TENSION LOG PLOT									
1.5 E 0 M DIOXANE									
25.0 2.60 X10-4 M EC SURFACE TENSION LOG PLOT									
3.0 E 0 M DIOXANE									
25.0 5.70 X10-4 M EC SURFACE TENSION LOG PLOT									
3. E 0 M UREA									
25.0 3.50 X10-4 M EC SURFACE TENSION LOG PLOT									
6. E 0 M UREA									
25.0 7.40 X10-4 M EC SURFACE TENSION LOG PLOT									
3. E 0 M GUANIDINIUM CL									
25.0 4.25 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
5. E 0 M GUANIDINIUM CL									
25.0 9.60 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
E 0 H CL									
25.0 1.85 X10-4 M EC SURFACE TENSION LOG PLOT									
4.6 E 0 PH OF SOLUTION									
10 ENTRIES FOR COMPOUND									
COMPOUND NO = 325 MOL WGT - 362.6 DODECYL/OXYETHYLENE/4 ALCOHOL									
REDUCED POLYDISPERSITY OF HEAD GROUPS									
05.0 7.8 X10-5 M EC SURFACE TENSION LOG PLOT									
25 4.0 X10-5 M EC SURFACE TENSION LOG PLOT									
25.0 4.0 X10-5 M EC SURFACE TENSION LOG PLOT									
25.0 4.0 X10-5 M ED SURFACE TENSION LOG PLOT									
45.0 2.2 X10-5 M ED SURFACE TENSION LOG PLOT									
55.0 1.7 X10-5 M ED SURFACE TENSION LOG PLOT									
2.5 E 1 I 0001									
05.0 1.1 X10-4 M ED SURFACE TENSION LOG PLOT									
5.0 E 1 I 0001									
05.0 1.8 X10-4 M ED SURFACE TENSION LOG PLOT									
7.5 E 1 I 0001									
05.0 3.5 X10-4 M EC SURFACE TENSION LOG PLOT									
9.0 E 1 I 0001									
05.0 5.5 X10-4 M EC SURFACE TENSION LOG PLOT									
2.5 E 1 I 0001									
25.0 8.5 X10-5 M EC SURFACE TENSION LOG PLOT									
5.0 E 1 I 0001									
25.0 1.4 X10-4 M ED SURFACE TENSION LOG PLOT									
7.5 E 1 I 0001									
25.0 2.8 X10-4 M EC SURFACE TENSION LOG PLOT									
9.0 E 1 I 0001									
25.0 5.0 X10-4 M EC SURFACE TENSION LOG PLOT									
2.5 E 1 I 0001									
45.0 5.0 X10-5 M EC SURFACE TENSION LOG PLOT									
5.0 E 1 I 0001									
45.0 9.0 X10-5 M EC SURFACE TENSION LOG PLOT									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; F—wt % solvent; G—mol % surfactant; H—varied; I—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/l; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
7.5 E 1 I	0001		45.0	1.4 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L
9.0 E 1 I	0001		45.0	3.0 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L
	0097		25			SEE CMPD NMBR IN ADDITV	SCHI	66025	X
19 ENTRIES FOR COMPOUND									
COMPOUND NO = 326 MOL WGT - 803.2 DODECYL/OXYETHYLENE/14 ALCOHOL									
REDUCED POLYDISPERSITY OF HEAD GROUPS									
		62019				VALUES FRM REF IN CMC	SCHI ATLA	62020	R
		25.0	5.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
		55.0	2.5 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 327 MOL WGT - 1,199.8 DODECYL/OXYETHYLENE/23 ALCOHOL									
REDUCED POLYDISPERSITY OF HEAD GROUPS									
		62019				VALUES FRM REF IN CMC	SCHI ATLA	62020	R
		05.0	1.0 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
		25	6.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	66025	L L	
		25.0	6.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
		25.0	6.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
		45.0	3.6 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
		55.0	3.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
9.0 E 0 C	0001	05.0	8.0 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
2.5 E 1 I	0001	05.0	1.0 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
5.0 E 1 I	0001	05.0	1.2 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
7.5 E 1 I	0001	05.0	3.2 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
7.5 E 0 C	0001	25.0	1.6 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
9.0 E 0 C	0001	25.0	4.8 X10-4 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
2.5 E 1 I	0001	25.0	6.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
5.0 E 1 I	0001	25.0	7.6 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
7.5 E 0 C	0001	45.0	7.1 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
9.0 E 0 C	0001	45.0	2.2 X10-4 M	ED	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
2.5 E 1 I	0001	45.0	3.6 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
5.0 E 1 I	0001	45.0	5.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI MANN	66001	L L	
	0097	25			SEE CMPD NMBR IN ADDITV	SCHI	66025	X	
20 ENTRIES FOR COMPOUND									
COMPOUND NO = 328 MOL WGT - 887.4 OCTADECYL/OXYETHYLENE/14 ALCOHOL									
REDUCED POLYDISPERSITY OF HEAD GROUPS									
		62019				VALUES FRM REF IN CMC	SCHI ATLA	62020	R
		25.0	6.0 X10-5 M	EC	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
		55.0	2.0 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 329 MOL WGT - 4,676.6 OCTADECYL/OXYETHYLENE/100 ALCOHOL									
REDUCED POLYDISPERSITY OF HEAD GROUPS									
		62019				VALUES FRM REF IN CMC	SCHI ATLA	62020	R
		25.0	2.0 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI	62019	T L	
8.6 E-1 M	NA CL	25.0	2.0 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI ATLA	62020	T L	
1.29 E 0 M	NA CL	25.0	2.0 X10-5 M	ED	SURFACE TENSION LOG PLOT	SCHI ATLA	62020	T L	
4 ENTRIES FOR COMPOUND									
COMPOUND NO = 330 MOL WGT - 550.9 HEXADECYL/OXYETHYLENE/7 ALCOHOL									
HOMOGENEOUS HEAD GROUP									
		25	1.74 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T 3	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 331 MOL WGT - 639.0 HEXADECYL/OXYETHYLENE/9 ALCOHOL									
HOMOGENEOUS HEAD GROUP									
		18	8.0 X10-6 M	CG	FLOCCULATION RATE	MATH OTTE	66037	T L	
		20	3.6 X10-5 M	BE	SURFACE TENSION LOG PLOT	CARL CHAL	64009	T L	
		20	3.5 X10-5 M	BG	FOTOMTR SPCTR CHNG I2	CARL CHAL	64009	T L	
		25	2.09 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T 3	
4 ENTRIES FOR COMPOUND									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/g; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 332 MOL WGT - HOMOGENEOUS HEAD GROUP	25	771.2 2.34 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T 3
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 333 MOL WGT - HOMOGENEOUS HEAD GROUP	25	903.4 3.09 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T 3
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 334 MOL WGT - HOMOGENEOUS HEAD GROUP	25	1,167.8 3.89 X10-6 M	BC	SURFACE TENSION LOG PLOT	ELWO MACF	62027	T 3
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 335 MOL WGT - BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	580.9 2.05 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 336 MOL WGT - BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	639.0 9.3 X10-3 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
	25	8.5 X10-5 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 337 MOL WGT - BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	683.0 8.3 X10-5 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
8.6 E-1 M NA CL	25	4.6 X10-5 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5.0 E-1 C 0168	25	9.5 X10-5 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
5.0 E-1 C 0169	25	1.30 X10-4 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
4 ENTRIES FOR COMPOUND							
COMPOUND NO = 338 MOL WGT - BRANCHED CHAIN, NATURAL OE DISTRIBUTION	25	4,626.4 1.00 X10-3 M	HD	SURFACE TENSION LOG PLOT	HSIA DUNN	56014	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 339 MOL WGT -	25	354.7 6.3 X10-1 N	CG	VISUAL SPCTR CHNGE PNCN	LELO TART	51003	L L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 340 MOL WGT -	23	410.8 1.10 X10-1 N	CB	UNSPECIFIED CONDUCTANCE	TART LELO	55021	T 3
	25	1.1 X10-1 N	CD	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L L
	60	1.1 X10-1 N	CD	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 341 MOL WGT -	60	466.9 -2.0 X10-2 N	CC	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L 3
	60	2.0 X10-2 N	CC	UNSPECIFIED CONDUCTANCE	TART LELO	55021	T L
	80	2.5 X10-2 N	CC	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 342 MOL WGT -	60	523.1 3.6 X10-3 N	CB	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L 3
	60	3.3 X10-3 N	CC	UNSPECIFIED CONDUCTANCE	TART LELO	55021	T L
	80	4.9 X10-3 N	CB	SPECFC CONDCTNCE GRAPH	LELO TART	51003	L L
3 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 343 MOL WGT -	174.2 25	SODIUM PENTANE SULFONATE 9.9 X10-1 M CD	METHOD NOT CITED		LELO TART	51003	L L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 344 MOL WGT -	188.2 25	SODIUM HEXANE SULFONATE 4.6 X10-1 M CG	VISUAL SPCTR CHNGE PNCN		LELO TART	51003	L L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 345 MOL WGT -	418.3 1.	DIDODECYL DIMETHYLAMMONIUM CHLORIDE X10-4 N CE	UNSPECIFIED CONDUCTANCE	TART LELO	55021	T L	
3. E-3 N NA CL 1 ENTRIES FOR COMPOUND							
COMPOUND NO = 346 MOL WGT -	437.8 63014 25	DECYL TRIMETHYLAMMONIUM DECYL SULFATE 4.6 X10-4 M BB	VALUES FRM REF IN CMC SURFACE TENSION LOG PLOT	CORK GOOD	65005	R	
1.00 E 2 I NA BR 4 ENTRIES FOR COMPOUND	25	4.5 X10-4 M BC	METHOD NOT CITED	CORK GOOD	66014	T L	
	25	4.6 X10-4 M BB	SURFACE TENSION LOG PLOT	CORK GOOD	63014	L 3	
COMPOUND NO = 347 MOL WGT -	381.6 25	OCTYL TRIMETHYLAMMONIUM OCTYL SULFATE 7.5 X10-3 M BC	METHOD NOT CITED	CORK GOOD	66014	T L	
1.00 E 2 I NA BR 2 ENTRIES FOR COMPOUND	25	7.5 X10-3 M BB	SURFACE TENSION LOG PLOT	CORK GOOD	65005	T 3	
COMPOUND NO = 348 MOL WGT -	443.2 25	NN-DIMETHYL 1-1-DIHYDROPENTADECALUORO OCTYL AMINE N-OXIDE /C7F15CH2N/CH3/20/ 4.7 X10-4 M BB	SURFACE TENSION LOG PLOT	CORK GOOD	65005	T 3	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 349 MOL WGT -	356.5 57.0	SODIUM OCTADECANONE 1-SULFONATE 7.5 X10-4 W DB	KRAFFT POINT SOLUBILITY	TART WRIG	39002	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 350 MOL WGT -	196.4 25	POTASSIUM NONANOATE 2.1 X10-1 M DG	FOTOMTR SPCTR CHNGE PNCN	HERZ	52015	T L	
3.7 E-2 M K CL 8.7 E-2 M K CL 1.59 E-1 M K CL 2.78 E-1 M K CL 4.61 E-1 M K CL 6.77 E-1 M K CL 8.55 E-1 M K CL 1.29 E 0 M K CL 1.63 E 0 M K CL 1.92 E 0 M K CL 2.11 E 0 M K CL 2.39 E 0 M K CL 2.64 E 0 M K CL 17 ENTRIES FOR COMPOUND	25	2.01 X10-1 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52015	T L	
	25	2.00 X10-1 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	2.0 X10-1 M BG	UNSPEC SPCTR CHNGE PNCN	KLEV	58011	T L	
	25	1.87 X10-1 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.75 X10-1 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.59 X10-1 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.39 X10-1 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.15 X10-1 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	9.7 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	8.6 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	6.4 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	5.4 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	4.8 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	TA L	
	25	4.2 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	4.0 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	3.8 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
COMPOUND NO = 351 MOL WGT -	252.5 25	POTASSIUM TRIDECANOATE 1.26 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52015	T L	
1.34 E-3 M K CL 2.91 E-3 M K CL 4.78 E-3 M K CL 7.06 E-3 M K CL 9.29 E-3 M K CL 1.34 E-2 M K CL 1.56 E-2 M K CL 1.87 E-2 M K CL 2.51 E-2 M K CL 2.76 E-2 M K CL 3.27 E-2 M K CL	25	1.2 X10-2 M BG	UNSPEC SPCTR CHNGE PNCN	KLEV	58011	T L	
	25	1.26 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.21 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.16 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.12 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	1.06 X10-2 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	9.29 X10-3 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	8.92 X10-3 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	8.42 X10-3 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	8.00 X10-3 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	7.17 X10-3 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	6.90 X10-3 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	
	25	6.54 X10-3 M DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality
counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

		Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
4.11 E-2 M	K CL		25	5.87 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
5.29 E-2 M	K CL		25	5.29 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
6.16 E-2 M	K CL		25	4.92 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
9.13 E-2 M	K CL		25	4.06 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
1.16 E-1 M	K CL		25	3.57 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
1.44 E-1 M	K CL		25	3.19 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
1.69 E-1 M	K CL		25	2.94 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
1.94 E-1 M	K CL		25	2.77 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
2.16 E-1 M	K CL		25	2.62 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
2.41 E-1 M	K CL		25	2.53 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
2.68 E-1 M	K CL		25	2.38 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
3.12 E-1 M	K CL		25	2.27 X10-3 M	DG	VISUAL SPCTR CHNGE PNCN	HERZ	52017	T	L
26 ENTRIES FOR COMPOUND										
COMPOUND NO = 352	MOL WGT -		388.4	SODIUM DI-1-METHYLISOAMYL SULFOSUCCINATE						
			25	3. X10-2 N	HE	UNSPECIFIED CONDUCTANCE	HAFF PICC	42003	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 353	MOL WGT -		393.7	OCTYL TRIMETHYLLAMMONIUM DECANE SULFONATE						
1.78 E-1 M	K CL		RM	5.75 X10-3 M	BB	DEBYE PLT LIGHT SCATTER	ANAC	53002	T	3
2.65 E-1 M	K CL		RM	5.67 X10-3 M	BB	DEBYE PLT LIGHT SCATTER	ANAC	53002	T	3
3 ENTRIES FOR COMPOUND										
COMPOUND NO = 354	MOL WGT -		255.9	HEXYL BENZYL DIMETHYLLAMMONIUM CHLORIDE						
			UNK	4.34 X10-2 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 355	MOL WGT -		363.1	AMMONIUM DODECAFLUOROHEPTANOATE	H/CF2/6COONH4					
			UNK	2.5 X10-1 M	CG	METHOD NOT CITED	ARRI PATT	53003	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 356	MOL WGT -		312.0	DECYL BENZYL DIMETHYLLAMMONIUM CHLORIDE						
			UNK	6.1 X10-3 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 357	MOL WGT -		368.1	TETRADECYL BENZYL DIMETHYLLAMMONIUM CHLORIDE						
			UNK	3.7 X10-4 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 358	MOL WGT -		424.2	OCTADECYL BENZYL DIMETHYLLAMMONIUM CHLORIDE						
			23	8.5 X10-6 M	HG	STREAMING CURRENT	CARD	66011	T	L
			UNK	7.1 X10-6 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
2 ENTRIES FOR COMPOUND										
COMPOUND NO = 359	MOL WGT -		328.9	OCTYL 4-NITROBENZYL DIMETHYLLAMMONIUM CHLORIDE						
			UNK	5.7 X10-2 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 360	MOL WGT -		357.0	DECYL 4-NITROBENZYL DIMETHYLLAMMONIUM CHLORIDE						
			UNK	2.3 X10-2 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 361	MOL WGT -		385.0	DODECYL 4-NITROBENZYL DIMETHYL AMMONIUM CHLORIDE						
			UNK	3.6 X10-3 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 362	MOL WGT -		413.1	TETRADECYL 4-NITROBENZYL DIMETHYLLAMMONIUM CHLORIDE						
			UNK	5.1 X10-4 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 363	MOL WGT -		441.2	HEXADECYL 4-NITROBENZYL DIMETHYLLAMMONIUM CHLORIDE						
			UNK	1.3 X10-4 M	PG	VISUAL SPCTR CHNGE INPX	ROSS KWAR	53007	T	L
1 ENTRIES FOR COMPOUND										

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 364 MOL WGT –	469.2 UNK 2.9	OCTADECYL 4-NITROBENZYL DIMETHYLMONIUM CHLORIDE X10-5 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 365 MOL WGT –	374.5 UNK 2.8	DODECYL 2-CHLOROBENZYL DIMETHYLMONIUM CHLORIDE X10-4 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 366 MOL WGT –	408.9 UNK 3.7	DODECYL 2-4-DICHLOROBENZYL DIMETHYLMONIUMCHLORIDE X10-4 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 367 MOL WGT –	374.5 UNK 4.2	DODECYL 4-CHLOROBENZYL DIMETHYLMONIUM CHLORIDE X10-4 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 368 MOL WGT –	401.0 UNK 6.9	DODECYL 2-HYDROXY-5-NITROBENZYL DIMETHYLMONIUM CHLORIDE X10-4 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 369 MOL WGT –	408.9 UNK 1.1	DODECYL 3-4-DICHLOROBENZYL DIMETHYLMONIUMCHLORIDE X10-3 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 370 MOL WGT –	384.1 UNK 3.8	DODECYL 3-4-METHYLENEDIOXYBENZYL DIMETHYLMONIUM CHLORIDE X10-3 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 371 MOL WGT –	400.1 UNK 3.9	DODECYL 3-4-DIMETHOXYBENZYL DIMETHYLMONIUMCHLORIDE X10-3 M PG VISUAL SPCTR CHNGE INPX			ROSS KWAR	53007	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 372 MOL WGT –	463.1 0092 UNK 3.8	AMMONIUM HEXADECAFLUORONONANOATE H/CF2/8 COO NH4 X10-2 M CG METHOD NOT CITED SEE CMPD NMBR IN ADDITV			ARRI PATT	53003	T L
2 ENTRIES FOR COMPOUND					ARRI PATT	53003	X
COMPOUND NO = 373 MOL WGT –	563.2 UNK 9.	AMMONIUM EICOSAFLUOROUNDECANOATE H/CF2/10 COO NH4 X10-3 M CG METHOD NOT CITED			ARRI PATT	53003	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 374 MOL WGT –	346.1 25 1.5 UNK 1.5	DODECAFLUOROHEPTANOIC ACID H/CF2/6COOH X10-1 M HG UNSPEC SPCTR CHNG PNCN X10-1 M CG METHOD NOT CITED			KLEV	58011	T L
2 ENTRIES FOR COMPOUND					ARRI PATT	53003	T L
COMPOUND NO = 375 MOL WGT –	446.1 25 3. UNK 3.	HEXADECAFLUORONONANOIC ACID H/CF2/8COOH X10-2 M HG UNSPEC SPCTR CHNG PNCN X10-2 M CG METHOD NOT CITED			KLEV	58011	T L
2 ENTRIES FOR COMPOUND					ARRI PATT	53003	T L
COMPOUND NO = 376 MOL WGT –	375.4 60015 18.0 4.1 18.5 4.3 20.3 5.10 24.9 5.26 25 5.60 25 5.70 25 5.26 30 5.0 30 4.5 30.1 5.60 34.9 5.85 40.0 6.30	DODECYL PYRIDINIUM IODIDE VALUES FRM REF IN CMC X10-3 M BC KRAFT POINT SOLUBILITY X10-3 M BC INTERFACIAL TNSN UNSPEC X10-3 M CB MICELLAR SPECTRAL CHANGE X10-3 M CB MICELLAR SPECTRAL CHANGE X10-3 M DC TURBIDITY PLT LITE SCATR X10-3 M CB SURFACE TENSION UNSPEC X10-3 M DB MICELLAR SPECTRAL CHANGE X10-3 M CG FOTOMTR SPCTR CHNG EOSN X10-3 M CA SPECFC CONDCTNCE GRAPH X10-3 M CB MICELLAR SPECTRAL CHANGE X10-3 M CB MICELLAR SPECTRAL CHANGE X10-3 M CB MICELLAR SPECTRAL CHANGE			FORD OTTE	66028	R
					ADDI FURM	56019	T L
					ADDI FURM	56019	T L
					MUKE RAY	66006	T L
					MUKE RAY	66006	T 3
					PARR	60015	T 3
					FORD OTTE	66028	T 3
					MUKE RAY	63032	T L
					MEGU KOND	59024	T L
					MEGU KOND	59024	T L
					MUKE RAY	66006	T L
					MUKE RAY	66006	T L
					MUKE RAY	66006	T L
					MUKE RAY	66006	T L

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; F – wt % solvent; G – mol % surfactant; H – normality; I – mol % surfactant mixture; J – mol/l (or kg); K – molal; L – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation	
2.5 E-3 M K I	44.9	6.70 X10-3 M	CB	MICELLAR SPECTRAL CHANGE	MUKE RAY	66006	T L	
5.0 E-3 M K I	45	6.70 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
1.00 E-2 M K I	RM	5.0 X10-3 M	CC	MICELLAR SPECTRAL CHANGE	HARK DRIZ	51010	T L	
2. E-2 M K I	RM	5.3 X10-3 M	CG	VISUAL SPCTR CHNG SKYB	HARK DRIZ	51010	T L	
2.02 E-2 M K I	RM	5.6 X10-3 M	CC	FOTOMTR SOLUBLZTN OROT	HARK DRIZ	51010	T L	
5.01 E-2 M K I	25	4.53 X10-3 M	DC	TURBIDITY PLT LITE SCATR	PARR	60015	T L	
1.002E-1 M K I	25	3.87 X10-3 M	DC	TURBIDITY PLT LITE SCATR	PARR	60015	T L	
2. E-2 M K I	25	2.94 X10-3 M	DC	TURBIDITY PLT LITE SCATR	PARR	60015	T L	
2.02 E-2 M K I	30	1.80 X10-3 M	CB	SURFACE TENSION UNSPEC	FORD OTTE	66028	T L	
5.01 E-2 M K I	30	1.94 X10-3 M	CB	MICELLAR SPECTRAL CHANGE	MUKE RAY	66006	T L	
1.0002E-1 M K I	30	1.12 X10-3 M	CB	MICELLAR SPECTRAL CHANGE	MUKE RAY	66006	T L	
1. E-4 M NA2 S203 TH10SULF	30	6.5 X10-4 M	CB	MICELLAR SPECTRAL CHANGE	MUKE RAY	66006	T L	
1. E-3 M NA2 S203 TH10SULF	25	5.15 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
1. E-3 M NA2 S203 TH10SULF	25	4.75 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
3.4 E 0 M UREA	45	5.63 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
5.9 E 0 M UREA	25	9.34 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
3.4 E 0 M UREA	25	1.36 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
5.9 E 0 M UREA	45	1.18 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
8.0 E 0 M UREA	45	1.71 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
1. E-4 M NA2 S203 TH10SULF	45	2.13 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
3.4 E 0 M UREA	25	9.30 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
1. E-4 M NA2 S203 TH10SULF	25	1.39 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
5.9 E 0 M UREA	25	5.75 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
1. E-3 M NA2 S203 TH10SULF	9.6 E-1 M UREA	25	9.10 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L
3.4 E 0 M UREA	25	1.33 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L	
1. E-3 M NA2 S203 TH10SULF	5.9 E 0 M UREA	45	7.10 X10-3 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L
9.6 E-1 M UREA	1. E-3 M NA2 S203 TH10SULF	45	1.10 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L
3.4 E 0 M UREA	1. E-3 M NA2 S203 TH10SULF	45	1.57 X10-2 M	DB	MICELLAR SPECTRAL CHANGE	MUKE RAY	63032	T L
5.9 E 0 M UREA	41 ENTRIES FOR COMPOUND							
COMPOUND NO = 377 MOL WGT - HOMOGENEOUS HEAD GROUP	406.7	DODECYL/OXYETHYLENE/ 5 ALCOHOL						
	20	4.0 X10-5 M	BD	SURFACE TENSION LOG PLOT	CARL CHAL	64009	T L	
	20	3.5 X10-5 M	BG	FOTOMTR SPCTR CHNG I2	CARL CHAL	64009	T L	
	20	5.00 X10-5 M	BD	REFRACTIVE INDEX	DONB JAN	63021	T L	
	23	5.7 X10-5 M	PB	SURFACE TENSION LOG PLOT	LANG	60012	T L	
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 378 MOL WGT - HOMOGENEOUS HEAD GROUP	378.6	DECYL/OXYETHYLENE/5 ALCOHOL						
	20	8.6 X10-4 M	BC	SURFACE TENSION LOG PLOT	CARL CHAL	64009	T L	
	20	1.0 X10-3 M	BG	FOTOMTR SPCTR CHNG I2	CARL CHAL	64009	T L	
	20	7.8 X10-4 M	BC	REFRACTIVE INDEX	DONB JAN	63021	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 379 MOL WGT - HOMOGENEOUS HEAD GROUP	334.6	DECYL/OXYETHYLENE/4 ALCOHOL						
	16	4.2 X10-4 M	BG	FOTOMTR SPCTR CHNG I2	CARL CHAL	64009	T L	
	20	6.4 X10-4 M	BC	SURFACE TENSION LOG PLOT	CARL CHAL	64009	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 380 MOL WGT - HOMOGENEOUS HEAD GROUP	278.4	HEXYL/OXYETHYLENE/4 ALCOHOL						
	20	9. X10-2 M	BD	SURFACE TENSION LOG PLOT	CARL CHAL	64009	T L	
	20	9. X10-2 M	BD	FOTOMTR SPCTR CHNG I2	CARL CHAL	64009	T L	
	20	9.0 X10-2 M	BB	REFRACTIVE INDEX	DONB JAN	63021	T 3	
	20	7.5 X10-2 M	BE	SURFACE TENSION LOG PLOT	MULL METC	62015	T L	
4 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO =	381 MOL WGT – HOMOGENEOUS HEAD GROUP	322.5	HEXYL/OXYETHYLENE/5 ALCOHOL					
		20	9.25 X10-2 M	BB	REFRACTIVE INDEX	DONB JAN	63021	T 3
		20	7.5 X10-2 M	BE	SURFACE TENSION LOG PLOT	MULL METC	62015	T L
2 ENTRIES FOR COMPOUND								
COMPOUND NO =	382 MOL WGT –	353.6	ETHYL TRIMETHYLAMMONIUM DODECYL SULFATE					
		25	4.30 X10-3 M	DA	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	383 MOL WGT –	381.6	BUTYL TRIMETHYLAMMONIUM DODECYL SULFATE					
		25	2.38 X10-3 M	DA	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	384 MOL WGT –	409.7	HEXYL TRIMETHYLAMMONIUM DODECYL SULFATE					
		25	1.25 X10-3 M	DB	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	385 MOL WGT –	437.8	OCTYL TRIMETHYLAMMONIUM DODECYL SULFATE					
		25	4.3 X10-4 M	BC	METHOD NOT CITED	CORK GOOD	66014	T 3
		25	4.0 X10-4 M	DB	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
2 ENTRIES FOR COMPOUND								
COMPOUND NO =	386 MOL WGT –	283.4	AMMONIUM DODECYL SULFATE					
		25	6.16 X10-3 M	DA	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
		30	7.2 X10-3 M	CB	EQUIV CONDCTNCE GRAPH	MEGU KOND	59026	T L
		30	6.6 X10-3 M	CG	FOTOMTR SPCTR CHNGE RHD6	MEGU KOND	59026	T L
3 ENTRIES FOR COMPOUND								
COMPOUND NO =	387 MOL WGT –	297.5	METHYLAMMONIUM DODECYL SULFATE					
		25	5.70 X10-3 M	DA	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	388 MOL WGT –	311.5	ETHYLAMMONIUM DODECYL SULFATE					
		25	5.00 X10-3 M	DA	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	389 MOL WGT –	339.5	BUTYLAMMONIUM DODECYL SULFATE					
		25	2.92 X10-3 M	DA	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	390 MOL WGT –	367.6	HEXYLAMMONIUM DODECYL SULFATE					
		25	1.12 X10-3 M	DB	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	391 MOL WGT –	395.7	OCTYLAMMONIUM DODECYL SULFATE					
		25	2.8 X10-4 M	DC	SPECFC CONDCTNCE GRAPH	PACK DONB	63030	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	392 MOL WGT –	165.7	OCTYLAMMONIUM CHLORIDE					
		25	1.75 X10-1 M	XC	REFRACTIVE INDEX	KLEV	53010	T L
1 ENTRIES FOR COMPOUND								
COMPOUND NO =	393 MOL WGT – HOMOGENEOUS HEAD GROUP	338.5	BUTYL/OXYETHYLENE/6 ALCOHOL					
		20	7.96 X10-1 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T 3
		30	7.60 X10-1 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T 3
		40	7.10 X10-1 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T L
3 ENTRIES FOR COMPOUND								
COMPOUND NO =	394 MOL WGT – HOMOGENEOUS HEAD GROUP	338.5	I-METHYL PROPYL/OXYETHYLENE/6 ALCOHOL					
		20	9.1 X10-1 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T 3

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
	30	8.8 X10-1 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T 3
	40	8.5 X10-1 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 395 MOL WGT — HOMOGENEOUS HEAD GROUP	366.6	2-ETHYL BUTYL/OXYETHYLENE/6 ALCOHOL					
	20	1.00 X10-1 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T 3
	30	9.3 X10-2 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T 3
	40	8.7 X10-2 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 396 MOL WGT — HOMOGENEOUS HEAD GROUP	394.6	2-PROPYL PENTYL/OXYETHYLENE/6 ALCOHOL					
	20	2.30 X10-2 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T 3
	30	2.0 X10-2 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, 3
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 397 MOL WGT — HOMOGENEOUS HEAD GROUP	422.7	2-BUTYL HEXYL/OXYETHYLENE/6 ALCOHOL					
	15	3.36 X10-3 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, L
	20	3.10 X10-3 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, 3
	25	2.84 X10-3 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, 3
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 398 MOL WGT — HOMOGENEOUS HEAD GROUP	554.9	2-BUTYL HEXYL/OXYETHYLENE/9 ALCOHOL					
	20	3.20 X10-3 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, 3
	30	2.79 X10-3 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, 3
	40	2.43 X10-3 M	BC	SURFACE TENSION LOG PLOT	ELWO FLOR	64049	T, L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 399 MOL WGT —	278.0	DODECYL DIMETHYL ETHYLMONIUM CHLORIDE					
	25	1.9 X10-2 M	BC	SURFACE TENSION LOG PLOT	WEIN ZOGR	65026	T, L
1. E-1 K CL	UNK	2.13 X10-2 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 400 MOL WGT —	292.0	DODECYL METHYL DIETHYLMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND	UNK	1.99 X10-2 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L
COMPOUND NO = 401 MOL WGT —	306.0	DODECYL TRIETHYLMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND	UNK	1.93 X10-2 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L
COMPOUND NO = 402 MOL WGT —	278.0	TRIDECYL TRIMETHYLMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND	UNK	1.12 X10-2 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L
COMPOUND NO = 403 MOL WGT —	326.0	DODECYL DIMETHYLPHENYLMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND	UNK	7.65 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L
COMPOUND NO = 404 MOL WGT — C6H5CH2CH2/N/CH3/2/C12H25	354.1	DODECYL DIMETHYL 2-PHENYLETHYLMONIUM CHLORIDE					
1 ENTRIES FOR COMPOUND	UNK	4.1 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L
COMPOUND NO = 405 MOL WGT —	354.1	DODECYL METHYLETHYL BENZYLAMMONIUM CHLORIDE					
1 ENTIRE FOR COMPOUND	UNK	7.7 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L
COMPOUND NO = 406 MOL WGT — CHLORIDE CF3C6H4CH2/N/CH3/2/C12H25	408.0	DODECYL DIMETHYL META-TRIFLUOROMETHYL BENZYL AMMONIUM					
1 ENTRIES FOR COMPOUND	UNK	3.2 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T, L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l (or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation	
COMPOUND NO = 407 MOL WGT – C6H5CH2CH2/N/CH3/2/C12H25	368.1	DODECYL DIMETHYL 3-PHENYLPROPYLAMMONIUM CHLORIDE							
	UNK	3.13 X10-3 M	CC	UNSPECIFIED CONDUCTANCE	CELL EGGE	52001	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 408 MOL WGT –	344.6	POTASSIUM HEXADECANE 1-SULFONATE							
	80	1.80 X10-3 W	CD	EQUIV CONDCTNCE GRAPH	MURR HART	35001	K L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 409 MOL WGT –	415.6	TRIETHANOLAMMONIUM DODECYL SULFATE							
	59017	VALUES FRM REF IN CMC			KASH	58021	R		
	33	4. X10-3 M	BE	SURFACE TNSN LINEAR PLOT	KASH EZAK	59017	T L		
	40	4. X10-3 M	BE	FOAMING POWER	KASH EZAK	59017	T L		
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 410 MOL WGT –	353.5	MORPHOLINIUM DODECYL SULFATE							
	59017	VALUES FRM REF IN CMC			KASH	58021	R		
	40	3. X10-3 M	BE	FOAMING POWER	KASH EZAK	59017	T L		
	40	4. X10-3 M	BE	SURFACE TNSN LINEAR PLOT	KASH EZAK	59017	T L		
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 411 MOL WGT –	217.4	DECYLLAMMONIUM ACETATE							
	UNK	4. X10-2 M	CE	METHOD NOT CITED	SOMA HEAL	64035	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 412 MOL WGT –	245.5	DODECYLLAMMONIUM ACETATE							
	UNK	1.3 X10-2 M	CD	METHOD NOT CITED	SOMA HEAL	64035	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 413 MOL WGT –	273.5	TETRADECYLAMMONIUM ACETATE							
	UNK	4. X10-3 M	CE	METHOD NOT CITED	SOMA HEAL	64035	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 414 MOL WGT –	301.6	HEXADECYLAMMONIUM ACETATE							
	UNK	8. X10-4 M	CE	METHOD NOT CITED	SOMA HEAL	64035	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 415 MOL WGT –	329.6	OCTADECYLAMMONIUM ACETATE							
	UNK	3 X10-4 M	CE	METHOD NOT CITED	SOMA HEAL	64035	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 416 MOL WGT –	314.1	PERFLUORO HEXANOIC ACID							
		QUESTIONABLE CRITERION			KLEV RAIS	54010	R		
		QUESTIONABLE CRITERION			KLEV VERG	57017	R		
	57017	VALUES FRM REF IN CMC			KLEV VERG	56010	R		
	0	1.09 X10-1 M	BC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3		
	18.5	8.2 X10-2 M	BC	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T L		
	18	1.06 X10-1 M	BC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3		
	25	5.4 X10-2 M	BG	UNSPEC SPCTR CHNG PNCC	KLEV	58011	T I		
	UNK	5.0 X10-2 M	CG	METHOD NOT CITED	KLEV CARR	56001	T L		
	UNK	5.1 X10-2 M	XG	METHOD NOT CITED	KLEV RAIS	54004	T L		
	UNK			GRAPH DATA NOT RETRIEVED	KLEV RAIS	54004	R		
E O PH OF SOLUTION									
10 ENTRIES FOR COMPOUND									
COMPOUND NO = 417 MOL WGT –	414.1	PERFLUORO OCTANOIC ACID							
		QUESTIONABLE CRITERION			KLEV RAIS	54010	R		
		QUESTIONABLE CRITERION			KLEV VERG	57017	R		
	57017	VALUES FRM REF IN CMC			KLEV VERG	56010	R		
	18.5	9.0 X10-3 M	BC	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T L		
	18	9.8 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3		
	25	5.6 X10-3 M	BG	UNSPEC SPCTR CHNG PNCC	KLEV	58011	T L		
	30	8.7 X10-3 M	BC	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T L		
	35	9.3 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3		
	45	1.02 X10-2 M	BB	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P 3		
	UNK	5.1 X10-3 M	CG	METHOD NOT CITED	KLEV CARR	56001	T L		
	2.5 E-2 M	H CL	30	3.2 X10-3 M	BC	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T L
	5. E-2 M	H CL	30	2. X10-3 M	BE	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T L

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality; counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives			Temp. °C	CMC		Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
5. E-2 M	K CL		30	5.	X10-3 M	BE	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
5. E-3 M	K OH		30	1.12	X10-2 M	BD	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
1. E-2 M	K OH		30	1.52	X10-2 M	BD	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
2.5 E-2 M	K OH		30	3.02	X10-2 M	BD	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
5. E-2 M	K OH		30	2.4	X10-2 M	BD	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
1.0 E-1 M	K OH		30	1.9	X10-2 M	BD	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
. E 0	PH OF SOLUTION		UNK				GRAPH DATA NOT RETRIEVED	KLEV RAIS	54004	R	
2.5 E 1 C	0001		30	1.10	X10-2 M	CD	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
5.0 E 1 C	0001		30	4.5	X10-3 M	CD	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
8.0 E 1 C	0001		30	5.	X10-3 M	CE	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
9.0 E 1 C	0001		30	5.5	X10-3 M	CE	SURFACE TENSION MINIMUM	KLEV RAIS	54010	T	L
23 ENTRIES FOR COMPOUND											
COMPOUND NO =	418	MOL WGT -	258.3	SODIUM UNDECYL SULFONATE							
			20	1.9	X10-2 M	BC	INTERFACIAL TENSION LOGM	V VO	60025	T	L
3. E-2 K	NA CL		20	1.4	X10-2 M	BC	INTERFACIAL TENSION LOGM	V VO	60025	T	L
2 ENTRIES FOR COMPOUND											
COMPOUND NO =	419	MOL WGT -	356.5	SODIUM ALPHA-HEPTYL UNDECYL SULFONATE							
			20	3.6	X10-4 M	BB	INTERFACIAL TENSION LOGM	V VO	60025	T	L
1. E-3 K	NA CL		20	2.9	X10-4 M	BB	INTERFACIAL TENSION LOGM	V VO	60025	T	L
2 ENTRIES FOR COMPOUND											
COMPOUND NO =	420	MOL WGT -	360.6	POTASSIUM 1-1-2-OCTANE TRICARBOXYLATE							
			25	7.9	X10-1 M	DG	VISUAL SPCTR CHNGE PNCRN	SHIN	56003	T	L
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	421	MOL WGT -	416.7	POTASSIUM 1-1-2-DODECANE TRICARBOXYLATE							
			25	9.5	X10-2 M	DG	VISUAL SPCTR CHNGE PNCRN	SHIN	56003	T	L
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	422	MOL WGT -	472.8	POTASSIUM 1-1-2-HEXADECANE TRICARBOXYLATE							
			25	1.2	X10-2 M	DG	VISUAL SPCTR CHNGE PNCRN	SHIN	56003	T	L
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	423	MOL WGT -	174.3	OCTYL/OXYETHYLENE/1 ALCOHOL--OCTYL GLYCOL ETHER							
	HOMOGENEOUS HEAD GROUP										
			25	4.9	X10-3 M	BC	SURFACE TENSION LOG PLOT	SHIN YAMA	59013	T	3
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	424	MOL WGT -	204.3	OCTYL ALPHA-GLYCERYL ETHER							
			25	5.8	X10-3 M	BC	SURFACE TENSION LOG PLOT	SHIN YAMA	59013	T	3
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	425	MOL WGT -	388.6	POTASSIUM 1-1-2-DECANE TRICARBOXYLATE							
			25				THEORETICALLY ESTIMATED	SHIN	56003	T	R
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	426	MOL WGT -	444.7	POTASSIUM 1-1-2-TETRADECANE TRICARBOXYLATE							
			25				THEORETICALLY ESTIMATED	SHIN	56003	T	R
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	427	MOL WGT -	384.5	HEXADECYL PYRIDINIUM BROMIDE							
			25	6.2	X10-4 M	CB	SPECFC CONDCTNCE GRAPH	BENT SPAR	66038	T	L
			25	5.81	X10-4 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P	2
			25	7.00	X10-4 W	BD	SPECFC CONDCTNCE GRAPH	CZER	65037	T	L
			27.0	4.8	X10-4 M	BC	KRAFT POINT SOLUBILITY	ADDI FURM	56019	T	L
			30.0	4.7	X10-4 M	BC	INTERFACIAL TNSN UNSPEC	ADDI FURM	56019	T	L
			35	7.5	X10-4 M	BB	EQUIV CONDCTNCE GRAPH	HART COLL	36001	T	3
			35	7.7	X10-4 M	BB	SPECFC CONDCTNCE GRAPH	HART COLL	36001	P	3
			35	7.80	X10-4 W	BD	SPECFC CONDCTNCE GRAPH	CZER	65037	T	L
			45	8.90	X10-4 W	BD	SPECFC CONDCTNCE GRAPH	CZER	65037	T	L
			55	1.035X10-3 W	BD	SPECFC CONDCTNCE GRAPH	CZER	65037	T	L	
3. E-3 M	K BR		40	3.4	X10-4 M	BD	TURBIDITY PLT LITE SCATR	TART	59010	T	L
6.4 E 0 H	METHANOL		25	7.51	X10-4 M	BC	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P	3
1.47OE 1 H	METHANOL		25	1.18	X10-3 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P	2
1.99IE 1 H	METHANOL		25	1.69	X10-3 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P	2

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2.602E 1 H METHANOL	25	2.81 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P	3
3.520E 1 H METHANOL	25	6.01 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P	2
E 0 0099				GRAPH DATA NOT RETRIEVED	LANG	53005	R	
E 0 0290				GRAPH DATA NOT RETRIEVED	LANG	53005	R	
18 ENTRIES FOR COMPOUND								
COMPOUND NO = 428 MOL WGT -	114.0	PERFLUORO ACETIC ACID						
				QUESTIONABLE CRITERION	KLEV RAIS	54010	R	
	57017			VALUES FRM REF IN CMC	KLEV VERG	56010	R	
	25	2.6 X10 0 M	BE	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
	25			QUESTIONABLE CRITERION	KLEV	58011	R	
	35			QUESTIONABLE CRITERION	KLEV VERG	57017	R	
5 ENTRIES FOR COMPOUND								
COMPOUND NO = 429 MOL WGT -	164.0	PERFLUORO PROPIONIC ACID						
	57017			VALUES FRM REF IN CMC	KLEV VERG	56010	R	
	25	1.11 X10-3 M	BG	UNSPEC SPCTR CHNG PN CN	KLEV	58011	T	L
	35			QUESTIONABLE CRITERION	KLEV VERG	57017	R	
	35	1.21 X10 0 M	AC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P	L
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 430 MOL WGT -	214.0	PERFLUORO BUTYRIC ACID						
				QUESTIONABLE CRITERION	KLEV RAIS	54010	R	
				QUESTIONABLE CRITERION	KLEV VERG	57017	R	
	57017			VALUES FRM REF IN CMC	KLEV VERG	56010	R	
	0	5.5 X10-1 M	AC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P	L
	04.9-	1.49 X10 0 W	AC	FREEZING POINT	HOLL CADY	59023	C	L
	18	5.0 X10-1 M	AC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P	L
	25	7.1 X10-1 M	BC	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L
	25	5.3 X10-1 M	BG	UNSPEC SPCTR CHNG PN CN	KLEV	58011	T	L
	35	4.9 X10-1 M	AC	EQUIV CONDCTNCE GRAPH	KLEV VERG	57017	P	L
9 ENTRIES FOR COMPOUND								
COMPOUND NO = 431 MOL WGT -	574.9	TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/8.5 ALCOHOL						
BRANCHED CHAIN, NATURAL OE DISTRIBUTION								
	UNK	8.8 X10-3 D	HC	FOTOMTR SPCTR CHNG I2	BECH	59006	G	L
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 432 MOL WGT -	641.0	TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL						
BRANCHED CHAIN, NATURAL OE DISTRIBUTION								
	UNK	8.1 X10-3 D	HC	FOTOMTR SPCTR CHNG I2	BECH	59006	G	L
E 0 NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	G	R
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 433 MOL WGT -	861.3	TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/15 ALCOHOL						
BRANCHED CHAIN, NATURAL OE DISTRIBUTION								
	UNK	1.3 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G	L
E 0 NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	G	R
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 434 MOL WGT -	1,169.7	TRIDECYL/OXO-PROCESS/ /OXYETHYLENE/22 ALCOHOL						
BRANCHED CHAIN, NATURAL OE DISTRIBUTION								
	UNK	2.8 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G	L
E 0 NA CL	25			SUMMARIZING EQN ONLY	BECH	62002	G	R
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 435 MOL WGT -	711.2	OCTADECYL/OXYETHYLENE/10 ALCOHOL						
NATURAL DISTRIBUTION OF HEAD GROUPS								
				QUESTIONABLE CRITERION	BECH	62002	R	
				VALUES FRM REF IN CMC	BECH	59006	R	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 436 MOL WGT -	1,151.8	OCTADECYL/OXYETHYLENE/20 ALCOHOL						
NATURAL DISTRIBUTION OF HEAD GROUPS								
				QUESTIONABLE CRITERION	BECH	62002	R	
				VALUES FRM REF IN CMC	BECH	59006	R	
2 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 437 MOL WGT - 797.3 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/12 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	UNK	3.3 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 438 MOL WGT - 973.5 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/16 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	UNK	3.2 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 439 MOL WGT - 1,149.7 OLEYL/CIS-9-OCTADECENOYL/ /OXYETHYLENE/20 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	UNK	2.9 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 440 MOL WGT - 522.8 /OXYETHYLENE/4 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	UNK	1.3 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 441 MOL WGT - 699.0 /OXYETHYLENE/8 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	UNK	1.2 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 442 MOL WGT - 787.1 /OXYETHYLENE/10 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	UNK	1.2 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 443 MOL WGT - 1,227.7 /OXYETHYLENE/20 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	UNK	1.4 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 444 MOL WGT - 1,448.0 /OXYETHYLENE/25 SORBITAN MONOLAURATE ISOMERS AND DISTRIBUTED MULTIPLE OE CHAINS	UNK	1.4 X10-2 D	HD	FOTOMTR SPCTR CHNG I2	BECH	59006	G L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 445 MOL WGT - 292.3 SODIUM PARA-BIS-/N-BUTYL/ BENZENE SULFONATE	60		QUESTIONABLE CRITERION	KOBE KLAM	60017	R	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 446 MOL WGT - 348.5 SODIUM PARA-BIS-/N-HEXYL/ BENZENE SULFONATE	60		QUESTIONABLE CRITERION	KOBE KLAM	60017	R	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 447 MOL WGT - 404.6 SODIUM PARA-BIS-/N-OCTYL/ BENZENE SULFONATE	60		QUESTIONABLE CRITERION	KOBE KLAM	60017	R	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 448 MOL WGT - 306.5 SODIUM OCTADECANOATE /STEARATE/							
50	1.8 X10-3 M	HC	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L	
UNK	4. X10-2 D	XG	VISUAL SPCTR CHNG PNCN	DEMC	61031	T L	
	1.3 X10-3 M					M	
1. E-1 H NA OH	70	7.5 X10-5 M	CC	SURFACE TENSION LOG PLOT	POWN ADDI	38006	K L
4 ENTRIES FOR COMPOUND							
COMPOUND NO = 449 MOL WGT - 235.9 DODECYLMETHYL AMMONIUM CHLORIDE	30	1.46 X10-2 M	BC	EQUIV CONDCTNCE GRAPH	RALS BROO	49013	T 3
1 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 450 MOL WGT –	249.9 DODECYLDIMETHYL AMMONIUM CHLORIDE	30	1.61 X10-2 M	BC EQUIV CONDCTNCE GRAPH		RALS BROO	49013	T	3
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 451 MOL WGT –	448.1 PARA DI-ISOBUTYLPHENOXYETHOXYETHYL DIMETHYLBENZYL AMMONIUM CHLORIDE /HYAMINE 1622/				QUESTIONABLE CRITERION GRAPH DATA NOT RETRIEVED	ROSS HUDS COHE VASS	57010 61027	R	R
5. E O DC ANTI FOAM A (PMS*)	24.7 1.6 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T	L		
5. E-2 P DECANOL-1	24.7 1.7 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T	L		
5. E-2 P LAURYL ALCOHOL	24.7 9. X10-2 P	HD	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T	L		
5. E-2 P TRIBUTYL PHOSPHATE	24.7 1.1 X10-1 P	HD	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T	L		
5.0 E-1 P TRIBUTYL PHOSPHATE	24.7 1.5 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T	L		
5. E-2 P 2-ETHYL HEXANOL	24.7 1.4 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T	L		
	24.7 1.3 X10-1 P	HC	SPECFC CONDCTNCE GRAPH	ROSS BRAM	57031	T	L		
9 ENTRIES FOR COMPOUND									
COMPOUND NO = 452 MOL WGT –	246.9 3-4-DICHLOROPERFLUORO BUTYRIC/KEL-F/ACID	25	7.0 X10-1 M	HG UNSPEC SPCTR CHNG PN CN	KLEV		58011	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 453 MOL WGT –	363.4 3-5-6-TRICHLOROPERFLUORO HEXANOIC/KEL-F/ACID	25	6.2 X10-2 M	HG UNSPEC SPCTR CHNG PN CN	KLEV		58011	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 454 MOL WGT –	479.9 3-5-7-8-TETRACHLOROPERFLUORO OCTANOIC/KEL-F/ACID	25	9.1 X10-3 M	HG UNSPEC SPCTR CHNG PN CN	KLEV		58011	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 455 MOL WGT –	514.1 PERFLUORO DECANOIC ACID	25	4.8 X10-4 M	BC UNSPEC SPCTR CHNG PN CN	KLEV		58011	T	L
	30 8.9 X10-4 M	BC	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L		
	UNK 8. X10-4 M	XG	METHOD NOT CITED	KLEV RAIS	54004	T	L		
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 456 MOL WGT –	452.2 POTASSIUM PERFLUORO OCTANOATE	25	2.88 X10-2 W	BB UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T	3	
	30 2.74 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T	3		
	30 2.63 X10-2 M	BC	SURFACE TENSION LOG PLOT	KLEV RAIS	54010	T	L		
	40 2.65 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T	3		
	55 2.76 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T	3		
	70 3.07 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T	3		
	85 3.54 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	T	3		
	UNK 2.7 X10-2 M	XG	METHOD NOT CITED	KLEV RAIS	54004	T	L		
9.4 E-3 W K N03	30 2.43 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
1.82 E-2 W K N03	30 2.22 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
2.96 E-2 W K N03	30 2.01 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
8.3 E-3 W K N03	40 2.40 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
2.39 E-2 W K N03	40 2.07 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
4.08 E-2 W K N03	40 1.79 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
7.31 E-2 W K N03	40 1.46 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
1.13 E-2 W K N03	55 2.42 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
2.30 E-2 W K N03	55 2.17 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
3.54 E-2 W K N03	55 1.93 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
4.65 E-2 W K N03	55 1.82 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
6.01 E-2 W K N03	55 1.65 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
7.77 E-2 W K N03	55 1.49 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
1.28 E-2 W K N03	70 2.75 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
3.77 E-2 W K N03	70 2.22 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
8.22 E-2 W K N03	70 1.82 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
4.65 E-2 W K N03	85 2.59 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
7.71 E-2 W K N03	85 2.27 X10-2 W	BB	UNSPECIFIED CONDUCTANCE	SHIN KATS	64011	L	3		
26 ENTRIES FOR COMPOUND									
COMPOUND NO = 457 MOL WGT –	118.2 1-4-HEXANEDIOL	24	1.9 X10-1 M	BC SURFACE TENSION LOG PLOT	KATO		63037	T	L
	UNK 2.0 X10-1 M	BC	FOTOMTR SOLUBLZTN SDN 4	KATO		63037	T	L	
	UNK 1.9 X10-1 M	BC	REFRACTIVE INDEX	KATO		63037	T	L	
3 ENTRIES FOR COMPOUND									

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; F – wt % solvent; G – mol % surfactant; H – normality; I – mol % surfactant mixture; J – molar; K – varied; L – normal; M – wt % surfactant; N – varied; O – mol/l or kg; P – mol/kg; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; V – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 458 MOL WGT -	347.3	DECYL PYRIDINIUM IODIDE						
RM 2.4 X10-2 M	CC	FOTOMTR SOLUBLZTN OROT	HARK KRIZ	51010	T L			
RM 2.25 X10-2 M	CB	MICELLAR SPECTRAL CHANGE	HARK KRIZ	51010	T 3			
RM 2.23 X10-2 M	CG	VISUAL SPCTR CHNG SKYB	HARK KRIZ	51010	T L			
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 459 MOL WGT - /CETOMACROGOL 1000/	.0	ALKYL/15-17/ /OXYETHYLENE/ /19-23/ALCHOL						
20 6.3 X10-3 D	HC	SURFACE TENSION LOG PLOT	ELWO	60027	T L			
20 7. X10-3 D	HE	FOTOMTR SOLUBLZTN DMYL	ELWO	60027	T L			
UNK 2.1 X10-3 D	HG	SURFACE TENSION UNSPEC	WAN	66018	T L			
UNK 1.1 X10-3 D	HE	SURFACE TENSION UNSPEC	HUGO NEWT	60026	T L			
UNK 1.2 X10-3 D	HE	FOTOMTR SPCTR CHNG I2	HUGO NEWT	60026	T L			
UNK 7.0 X10-3 D	HD	FOTOMTR SPCTR CHNG I2	ELWO	60027	T L			
2. E-1 M NA CL	UNK 6.7 X10-3 D	HD FOTOMTR SPCTR CHNG I2	ELWO	60027	G L			
7. E-1 M NA CL	UNK 6.2 X10-3 D	HD FOTOMTR SPCTR CHNG I2	ELWO	60027	G L			
1.0 E 0 M NA CL	UNK 5.8 X10-3 D	HD FOTOMTR SPCTR CHNG I2	ELWO	60027	G L			
1.5 E 0 M NA CL	UNK 4.7 X10-3 D	HD FOTOMTR SPCTR CHNG I2	ELWO	60027	G L			
2.0 E 0 M NA CL	UNK 3.5 X10-3 D	HD FOTOMTR SPCTR CHNG I2	ELWO	60027	G L			
2.5 E 0 M NA CL	UNK 3. X10-3 D	HE FOTOMTR SPCTR CHNG I2	ELWO	60027	T L			
12 ENTRIES FOR COMPOUND								
COMPOUND NO = 460 MOL WGT - 8.5 E 1 H H3 PO4	356.5 25 1.0 X10-3 M	DODECYL TROPYLIUM MONOPHOSPHATE	ABUH MYSE	65019	T L			
1 ENTRIES FOR COMPOUND		SURFACE TENSION LOG PLOT						
COMPOUND NO = 461 MOL WGT - HOMOGENEOUS HEAD GROUP	118.2	BUTYL/OXYETHYLENE/1 ALCOHOL--BUTYL GLYCOL ETHER						
25 9.8 X10-1 M	CC	REFRACTIVE INDEX	DONB JACO	66019	T L			
UNK 8.8 X10-1 M	CC	SURFACE TENSION LOG PLOT	DONB JACO	66019	T L			
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 462 MOL WGT -	376.4	SODIUM MONOLAURIN SULFATE						
25 5.2 X10-3 M	CC	SPECFC CONDCTNCE GRAPH	BISW MUKH	60028	T L			
UNK 2.6 X10-3 M	CG	VISUAL SPCTR CHNG RHD6	BISW MUKH	60028	T L			
4. E-3 M NA CL	25 3.4 X10-3 M	CC SPECFC CONDCTNCE GRAPH	BISW MUKH	60028	T L			
1. E-2 M NA CL	25 1.6 X10-3 M	CC SPECFC CONDCTNCE GRAPH	BISW MUKH	60028	T L			
2. E-2 M NA CL	25 1.03 X10-3 M	CD SPECFC CONDCTNCE GRAPH	BISW MUKH	60028	T L			
3. E-2 M NA CL	25 6.025X10-3 M	CE SPECFC CONDCTNCE GRAPH	BISW MUKH	60028	T L			
4. E-3 M NA CL	UNK 1.75 X10-3 M	CG VISUAL SPCTR CHNG RHD6	BISW MUKH	60028	T L			
1. E-2 M NA CL	UNK 1.3 X10-3 M	CG VISUAL SPCTR CHNG RHD6	BISW MUKH	60028	T L			
2. E-2 M NA CL	UNK 9. X10-4 M	CG VISUAL SPCTR CHNG RHD6	BISW MUKH	60028	T L			
3. E-2 M NA CL	UNK 6.3 X10-4 M	CG VISUAL SPCTR CHNG RHD6	BISW MUKH	60028	T L			
4. E-2 M NA CL	UNK 4.5 X10-4 M	CG VISUAL SPCTR CHNG RHD6	BISW MUKH	60028	T L			
11 ENTRIES FOR COMPOUND								
COMPOUND NO = 463 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	705.0 28 2.47 X10-5 M	NONYL BENZENE/OXYETHYLENE/11 ALCOHOL	MANK	66021	T L			
1 ENTRIES FOR COMPOUND		SURFACE TENSION LOG PLOT						
COMPOUND NO = 464 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	881.3	NONYL BENZENE/OXYETHYLENE/15 ALCOHOL						
21.2 3.10 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L			
25 2.85 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L			
28 2.71 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L			
33.5 2.69 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L			
41.5 2.36 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L			
45 2.27 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L			
6 ENTRIES FOR COMPOUND								
COMPOUND NO = 465 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	596.9 28 1.49 X10-5 M	TRIDECYL/OXYETHYLENE/9 ALCOHOL	MANK	66021	T L			
1 ENTRIES FOR COMPOUND		SURFACE TENSION LOG PLOT						

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 466 MOL WGT - 596.9 TRIDECYL/SECONDARY/ /OXYETHYLENE/9 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	28	5.15 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 467 MOL WGT - 729.1 TRIDECYL/SECONDARY/ /OXYETHYLENE/12 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	28	8.67 X10-5 M	HD	SURFACE TENSION LOG PLOT	MANK	66021	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 468 MOL WGT - 459.6 DODECYL/OXYETHYLENE/6.2 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	8 X10-5 M	ED	FOTOMTR SOLUBLZTN YLOB	TOKI	64024	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 469 MOL WGT - 521.2 DODECYL/OXYETHYLENE/7.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	9. X10-5 M	ED	FOTOMTR SOLUBLZTN YLOB	TOKI	64024	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 470 MOL WGT - 653.4 DODECYL/OXYETHYLENE/10.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	1.2 X10-4 M	ED	FOTOMTR SOLUBLZTN SDN 4	TOKI	64024	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 471 MOL WGT - 882.5 DODECYL/OXYETHYLENE/15.8 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	1.8 X10-4 M	ED	FOTOMTR SOLUBLZTN YLOB	TOKI	64024	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 472 MOL WGT - 1,085.2 DODECYL/OXYETHYLENE/20.4 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	2.1 X10-4 M	EC	FOTOMTR SOLUBLZTN SDN 4	TOKI	64024	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 473 MOL WGT - 1,380.4 DODECYL/OXYETHYLENE/27.1 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	2.5 X10-4 M	EC	FOTOMTR SOLUBLZTN SDN 4	TOKI	64024	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 474 MOL WGT - 1,887.1 DODECYL/OXYETHYLENE/38.6 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	3.6 X10-4 M	EC	FOTOMTR SOLUBLZTN SDN 4	TOKI	64024	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 475 MOL WGT - 3,151.6 DODECYL/OXYETHYLENE/67.3 ALCOHOL NATURAL DISTRIBUTION OF HEAD GROUPS	30	5.7 X10-4 M	EC	FOTOMTR SOLUBLZTN SDN 4	TOKI	64024	T L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 476 MOL WGT - 166.2 SODIUM OCTANOATE								
20 7. X10-2 M XG VISCOSITY MINIMUM								
20 3.6 X10-1 M DE VISCOSITY								
25 3.40 X10-1 M DB SPECFC CONDCTNCE GRAPH								
25 3.51 X10-1 M DB SURFACE TENSION LOG PLOT								
35 3.60 X10-1 M DB SPECFC CONDCTNCE GRAPH								
50 3.85 X10-1 M DB SURFACE TENSION LOG PLOT								
6 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 477 MOL WGT -	392.6 23 3.0	OCTADECYL TRIMETHYLLAMMONIUM BROMIDE X10-4 M HG		STREAMING CURRENT	CARD	66011	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 478 MOL WGT -	378.6 23 4.8	HEXADECYL DIMETHYLETHYLAMMONIUM BROMIDE X10-3 M GH		STREAMING CURRENT	CARD	66011	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 479 MOL WGT -	403.4 28.0 7.9 40.0 1.2	TETRADECYL PYRIDINIUM IODIDE X10-4 M BC X10-3 M BD	KRAFT POINT SOLUBILITY INTERFACIAL TNSN UNSPEC		ADDI FURM ADDI FURM	56019 56019	T L T L
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 480 MOL WGT -	431.5 35.0 3.1 50.0 4.4	HEXADECYL PYRIDINIUM IODIDE X10-4 M BC X10-4 M BC	KRAFT POINT SOLUBILITY INTERFACIAL TNSN UNSPEC		ADDI FURM ADDI FURM	56019 56019	T L T L
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 481 MOL WGT -	459.6 45.5 1.3	OCTADECYL PYRIDINIUM IODIDE X10-4 M BD	KRAFT POINT SOLUBILITY		ADDI FURM	56019	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 482 MOL WGT -	248.4	DODECYL AMMONIUM NITRATE 30 9.9 30 9.9 30 7.1 30 6.9 30 6.9	X10-3 M BC X10-3 M BC X10-3 M BC X10-3 M BC X10-3 M BC	EQUIV CONDCTNCE GRAPH EQUIV CONDCTNCE GRAPH EQUIV CONDCTNCE GRAPH EQUIV CONDCTNCE GRAPH EQUIV CONDCTNCE GRAPH	RALS EGGE RALS EGGE RALS EGGE RALS EGGE RALS EGGE	49008 49008 49008 49008 49008	K 3 K L K L K L K L
2. E-2 N ACETIC ACID 2. E-2 N H CL 2. E-2 N NA ACETATE 2. E-2 N NA CL							
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 483 MOL WGT -	306.0 30 2.83	DIOCTYL DIMETHYL AMMONIUM CHLORIDE X10-2 M BB	EQUIV CONDCTNCE GRAPH		RALS EGGE	48014	K 3
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 484 MOL WGT -	124.1 20 2.35	SODIUM PENTANOATE/VALERATE/ X10 0 M DD	X-RAY DIFFRACTION		HESS PHIL	39009	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 485 MOL WGT -	138.2 20 1.57	SODIUM HEXANOATE/CAPROATE/ X10 0 M DE	VISCOSITY		HESS PHIL	39009	T L
1.1 E 1 PH OF SOLUTION 1.1 E 1 PH OF SOLUTION 1.1 E 1 PH OF SOLUTION 1.1 E 1 PH OF SOLUTION	5 6.60 20 7.30 60 8.90 90 1.11	X10-1 M BC X10-1 M BC X10-1 M BC X10 0 M BC	SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH		MARK TSIK MARK TSIK MARK TSIK MARK TSIK	64051 64051 64051 64051	T L T L T L T L
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 486 MOL WGT -	152.2 20 9.5	SODIUM HEPTANOATE X10-1 M DE	VISCOSITY		HESS PHIL	39009	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 487 MOL WGT -	180.2 20 2.4 20 2.2	SODIUM NONANOATE X10-1 M XG X10-1 M DE	VISCOSITY MINIMUM VISCOSITY		SATA TYUZ HESS PHIL	53006 39009	T L T L
1.1 E 1 PH OF SOLUTION 1.1 E 1 PH OF SOLUTION 1.1 E 1 PH OF SOLUTION	20 1.59 60 1.82 90 2.04	X10-1 M BC X10-1 M BC X10-1 M BC	SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH SPECFC CONDCTNCE GRAPH		MARK TSIK MARK TSIK MARK TSIK	64051 64051 64051	T L T L T L
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 488 MOL WGT -	494.8 HOMOGENEOUS HEAD GROUP 23 7.1 23 8.0	DODECYL/OXYETHYLENE/7 ALCOHOL X10-5 M PE X10-5 M PB	FOTOMTR SPCTR CHNG I2 SURFACE TENSION LOG PLOT		LANG LANG	60012 60012	T L T L
2 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—moi % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 489 MOL WGT - HOMOGENEOUS HEAD GROUP		23	582.9 X10-5 M	PE	FOTOMTR SPCTR CHNG I2	LANG	60012	T L
		23	1.0 X10-4 M	PC	SURFACE TENSION LOG PLOT	LANG	60012	T L
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 490 MOL WGT - HOMOGENEOUS HEAD GROUP		23	715.1 X10-4 M	PC	SURFACE TENSION LOG PLOT	LANG	60012	T L
		23	1.4 X10-4 M	PE	FOTOMTR SPCTR CHNG I2	LANG	60012	T L
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 491 MOL WGT - 6.0 E 1 H H2 SO4		25	356.5 X10-3 M	BD	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T L
7.8 E 1 H H2 SO4		25	1.0 X10-3 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3
9.6 E 1 H H2 SO4		25	2.2 X10-3 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3
9.6 E 1 H H2 SO4		25	1.25 X10-2 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3
9. E-1 M NA2 SO4		25	5.4 X10-3 M	BC	SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3
4 ENTRIES FOR COMPOUND								
COMPOUND NO = 492 MOL WGT - BENZENE SULFONATE		75	348.5 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6	GRIE	55028	T L
2. E-1 M NA CL		20	3.14 X10-4 M	CC	SURFACE TENSION LOG PLOT	HARR	59001	K L
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 493 MOL WGT - 1 ENTRIES FOR COMPOUND		20	306.4 X10-3 M	BB	SODIUM TETRAPROPYLENE/1-3-5-7-TETRAMETHYL-OCTYL/SURFACE TENSION LOG PLOT	HARR	59001	L 3
COMPOUND NO = 494 MOL WGT -		25.8	.0 X10-4 M	HG	POTASSIUM DILINOLEATE FOTOMTR SPCTR CHNGE PNCH	CORR KLEV	46010	T L
		25.8	2.5 X10-4 M	HC	SPECFC CONDCTNCE GRAPH	CORR KLEV	46010	T L
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 495 MOL WGT -		20	524.7 X10-4 M	BE	SUCROSE MONOLAURATE INTERFACIAL TENSION LOGM	WACH HAYA	62023	T L
		20	1.85 X10-4 M	BE	SURFACE TENSION LOG PLOT	WACH HAYA	62023	T L
		27.1	1.3 X10-4 M	CD	SURFACE TENSION LOG PLOT	OSIP SNEL	57024	T L
		27.1	6.4 X10-6 M	CD	SURFACE TENSION LOG PLOT	OSIP SNEL	57024	T L
4 ENTRIES FOR COMPOUND		27.1	3.4 X10-4 M	CD	SURFACE TENSION LOG PLOT	OSIP SNEL	57024	T L
COMPOUND NO = 496 MOL WGT - 1 ENTRIES FOR COMPOUND		50	608.9 X10-5 M	BE	SUCROSE MONOSTEARATE SURFACE TENSION LOG PLOT	WACH HAYA	62023	TL L
COMPOUND NO = 497 MOL WGT - 1 ENTRIES FOR COMPOUND		20	552.7 X10-5 M	BE	SUCROSE MONOMYRISTATE INTERFACIAL TENSION LOGM	WACH HAYA	62023	T L
COMPOUND NO = 498 MOL WGT - 1 ENTRIES FOR COMPOUND		50	580.8 X10-5 M	BE	SUCROSE MONOPALMITATE SURFACE TENSION LOG PLOT	WACH HAYA	62023	TL L
COMPOUND NO = 499 MOL WGT - 1 ENTRIES FOR COMPOUND		819.3	SUCROSE DI-PALMITATE QUESTIONABLE CRITERION			WACH HAYA	62023	R
COMPOUND NO = 500 MOL WGT - 1 ENTRIES FOR COMPOUND		30	354.0 X10-2 M	BB	DODECYL TRI/2-HYDROXYETHYL/AMMONIUM CHLORIDE EQUIV CONDCTNCE GRAPH	RALS EGGE	49009	K 3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

	Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 501 MOL WGT -		264.3	SODIUM HEXYL BENZENE SULFONATE						
	75	3.71 X10-2 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	1.61 X10 0 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		6.091X10-2 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 502 MOL WGT -		278.3	SODIUM HEPTYL BENZENE SULFONATE						
	75	2.09 X10-2 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	5.98 X10-1 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		2.148X10-2 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 503 MOL WGT -		292.3	SODIUM OCTYL BENZENE SULFONATE						
	75	1.06 X10-2 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	3.2 X10-1 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		1.09 X10-2 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 504 MOL WGT -		306.4	SODIUM NYONYL BENZENE SULFONATE						
	75	6.50 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 505 MOL WGT -		320.4	SODIUM DECYL BENZENE SULFONATE						
	75	3.70 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	1.2 X10-1 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		3.74 X10-3 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 506 MOL WGT -		348.5	SODIUM DODECYL BENZENE SULFONATE						
	75	1.19 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	4.1 X10-2 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		1.17 X10-3 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 507 MOL WGT -		376.5	SODIUM TETRADECYL BENZENE SULFONATE						
	75	6.6 X10-4 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	2.5 X10-2 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		6.64 X10-4 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 508 MOL WGT -		404.6	SODIUM HEXADECYL BENZENE SULFONATE						
	75	5.35 X10-4 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	1.6 X10-2 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		3.95 X10-4 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 509 MOL WGT -		432.6	SODIUM OCTADECYL BENZENE SULFONATE						
	75	6.38 X10-4 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
	UNK	1.0 X10-2 D	XG	VISUAL SPCTR CHNGE PNCN		DEMC	61031	T	L
		2.31 X10-4 M						M	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 510 MOL WGT -		292.3	SODIUM 2-ETHYL-HEXYL BENZENE SULFONATE						
	75	2.54 X10-2 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 511 MOL WGT -		320.4	SODIUM 2-PROPYL-HEPTYL BENZENE SULFONATE						
	75	8.48 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 512 MOL WGT -		348.5	SODIUM 2-BUTYL-OCTYL BENZENE SULFONATE						
	75	3.20 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6		GRIE	55028	T	L
1 ENTRIES FOR COMPOUND									

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 513 MOL WGT -	376.5 75	SODIUM 2-AMYL-NONYL BENZENE SULFONATE 3.32 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6	GRIE	55028	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 514 MOL WGT -	348.5 75	SODIUM 6-N-DODECYL BENZENE SULFONATE 3.12 X10-3 M	BG	VISUAL SPCTR CHNGE RHD6	GRIE	55028	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 515 MOL WGT -	306.5 BRANCHED CHAIN, NATURAL OE DISTRIBUTION 25	ISO-OCTYL/OXYETHYLENE/4 ALCOHOL 5.00 X10-1 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 516 MOL WGT -	320.5 BRANCHED CHAIN, NATURAL OE DISTRIBUTION 25	NONYL/OXYETHYLENE/4 ALCOHOL 3.45 X10-1 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 517 MOL WGT -	598.9 BRANCHED CHAIN, NATURAL OE DISTRIBUTION 25	DECYL/OXO-PROCESS/ /OXYETHYLENE/10 ALCOHOL 1.30 X10-1 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 518 MOL WGT -	450.7 BRANCHED CHAIN, NATURAL OE DISTRIBUTION 25	TRIMETHYLNONYL/OXYETHYLENE/6 ALCOHOL 8.4 X10-2 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 519 MOL WGT -	334.6 BRANCHED CHAIN, NATURAL OE DISTRIBUTION 25	DECYL/OXO-PROCESS/ /OXYETHYLENE/4 ALCOHOL 6.6 X10-2 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 520 MOL WGT -	524.8 NATURAL DISTRIBUTION OF HEAD GROUPS 25	UNDECYL/OXYETHYLENE/8 ALCOHOL 2.3 X10-2 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 521 MOL WGT -	641.0 BRANCHED CHAIN, NATURAL OE DISTRIBUTION 25	TRIDECYL/OXYETHYLENE/10 ALCOHOL 9.4 X10-3 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 522 MOL WGT -	727.9 NATURAL DISTRIBUTION OF HEAD GROUPS 25	DECYL BENZENE ORTHO/OXYETHYLENE/11.2 ALCOHOL 2.15 X10-3 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 523 MOL WGT -	719.1 NATURAL DISTRIBUTION OF HEAD GROUPS 25	DECYL BENZENE PARA/OXYETHYLENE/11.0 ALCOHOL 1.4 X10-3 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 524 MOL WGT -	859.3 BRANCHED CHAIN, NATURAL OE DISTRIBUTION 25	HEXADECYL/OXYETHYLENE/14 ALCOHOL 1.3 X10-3 P	HD	SURFACE TENSION LOG PLOT	KOMO BEIS	66022	T L
1 ENTRIES FOR COMPOUND							

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture;
D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality
counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg;
T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 525	MOL WGT –	316.4 60 RM	SODIUM TETRADECYL 6-SULFATE 9.80 X10-3 M 1.23 X10-2 M	CB UNSPECIFIED CONDUCTANCE CG VISUAL SPCTR CHNGE PNCN	WINS WINS	48008 48008	L 3 T L		
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 526	MOL WGT –	316.4 RM	SODIUM 2-DI-N-HEXYL ETHYL SULFATE 8.38 X10-3 M	CG VISUAL SPCTR CHNGE PNCN	WINS	48008	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 527	MOL WGT –	222.3 RM	SODIUM UNDECANE-3-CARBOXYLATE 7.8 X10-2 M	CG VISUAL SPCTR CHNGE PNCN	WINS	48008	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 528	MOL WGT – 7.0 E 1 H H ClO4	358.9 25	DODECYL TROPYLIUM PERCHLORATE 8.6 X10-4 M	BC SURFACE TENSION LOG PLOT	ABUH MYSE	65019	T 3		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 529	MOL WGT – 8.5 E 1 H H2 SO4 E 0 PH OF SOLUTION	144.2 UNK 27 UNK	OCTANOIC ACID X10-3 M X10-1 M GRAPH DATA NOT RETRIEVED	XG SURFACE TENSION UNSPEC CE SURFACE TENSION LOG PLOT	KLEV RAIS STEI SHAN KLEV RAIS	54004 65025 54004	T L T L R		
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 530	MOL WGT – 8.5 E 1 H H2 SO4	172.3 27	DECANOIC ACID 2.4 X10-2 M	CE SURFACE TENSION LOG PLOT	STEI SHAN	65025	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 531	MOL WGT – 9.5 E 1 H H2 SO4 9.5 E 1 H H2 SO4	200.4 27 UNK	DODECANOIC ACID X10-2 M X10 0 D 5.98 X10-2 M	CD SURFACE TENSION LOG PLOT CB TURBIDITY PLT LITE SCATR	STEI SHAN STEI SHAN	65025 65025	T L C L		
9.5 E 1 H H2 SO4 5. E 0 M K H SO4									
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 532	MOL WGT – 9.5 E 1 H H2 SO4	228.4 27	TETRADECANOIC ACID 1.3 X10-2 M	CD SURFACE TENSION LOG PLOT	STEI SHAN	65025	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 533	MOL WGT – 9.5 E 1 H H2 SO4	256.5 27	HEXADECANOIC ACID 2.8 X10-3 M	CD SURFACE TENSION LOG PLOT	STEI SHAN	65025	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 534	MOL WGT – 9.5 E 1 H H2 SO4 9.55 E 1 H H2 SO4 9.62 E 1 H H2 SO4 9.73 E 1 H H2 SO4 9.5 E 1 H H2 SO4	284.5 27 27 27 27 UNK	OCTADECANOIC ACID X10-4 M X10-4 M X10-3 M X10-3 M X10-3 D	CD SURFACE TENSION LOG PLOT CD SURFACE TENSION LOG PLOT CD SURFACE TENSION LOG PLOT CD TURBIDITY PLT LITE SCATR	STEI SHAN STEI SHAN STEI SHAN STEI SHAN STEI SHAN	65025 65025 65025 65025 65025	T L T L T L T L C L		
5. E 0 M K H SO4									
6 ENTRIES FOR COMPOUND									
COMPOUND NO = 535	MOL WGT – 0099	1,035.6 25 25	HEXADECYL/OXYETHYLENE/18 ALCOHOL 1.3 X10-5 M	ED SURFACE TENSION LOG PLOT SEE CMPD NMBR IN ADDITV	SCHI SCHI	66025 66025	L L X		
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 536	MOL WGT –	208.3 25	NONYL SULFONIC ACID 4. X10-2 W	CE EQUIV CONDUCTNCE GRAPH	MCBA DYE	39011	T L		
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 537	MOL WGT –	236.4 25	UNDECYL SULFONIC ACID 1.5 X10-2 W	CD EQUIV CONDUCTNCE GRAPH	MCBA DYE	39011	T L		
1 ENTRIES FOR COMPOUND									

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture;
 D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality
 counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l(kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 538 MOL WGT — 1 ENTRIES FOR COMPOUND		PLURONIC L62 UNK 2.40 X10 0 D	HC	FOTOMTR SPCTR CHNG I2	ROSS OLIV	59020	T L	
COMPOUND NO = 539 MOL WGT — 1 ENTRIES FOR COMPOUND		RENEX 698 UNK 4.7 X10-3 D	HC	FOTOMTR SPCTR CHNG I2	ROSS OLIV	59020	T L	
COMPOUND NO = 540 MOL WGT — 2 ENTRIES FOR COMPOUND		SIPONIC BC UNK 1.1 X10-2 D UNK 1.20 X10-2 D	HD HC	FOTOMTR SPCTR CHNG I2 SURFACE TENSION LOG PLOT	ROSS OLIV ROSS OLIV	59020 59020	T L T L	
COMPOUND NO = 541 MOL WGT — 1 ENTRIES FOR COMPOUND	50	332.4 SODIUM DODECYL MONO-OXYETHYLENE SULFATE 4.78 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 542 MOL WGT — 1 ENTRIES FOR COMPOUND	50	376.5 SODIUM DODECYL DIOXYETHYLENE SULFATE 3.00 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 543 MOL WGT — 1 ENTRIES FOR COMPOUND	50	464.6 SODIUM DODECYL TETRA-OXYETHYLENE SULFATE 1.26 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 544 MOL WGT — 1 ENTRIES FOR COMPOUND	50	360.5 SODIUM TETRADECYL MONO-OXYETHYLENE SULFATE 1.39 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 545 MOL WGT — 1 ENTRIES FOR COMPOUND	50	404.5 SODIUM TETRADECYL DI-OXYETHYLENE SULFATE 1.00 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 546 MOL WGT — 1 ENTRIES FOR COMPOUND	50	448.6 SODIUM TETRADECYL TRI-OXYETHYLENE SULFATE 6.92 X10-4 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 547 MOL WGT — 1 ENTRIES FOR COMPOUND	50	414.6 SODIUM OLEYL MONO-OXYETHYLENE SULFATE 2.00 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 548 MOL WGT — 1 ENTRIES FOR COMPOUND	50	458.6 SODIUM OLEYL DI-OXYETHYLENE SULFATE 1.77 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 549 MOL WGT — 1 ENTRIES FOR COMPOUND	50	502.7 SODIUM OLEYL TRI-OXYETHYLENE SULFATE 1.19 X10-3 M	CD	SURFACE TENSION UNSPEC	GOTT	60018	G L	
COMPOUND NO = 550 MOL WGT — . E O K CL . E O K I . E O K NO3 4 ENTRIES FOR COMPOUND		.0 LAURIC ACID DIETHANOLAMINE CONDENSATE UNK 4.0 X10-5 M 25 3.98 X10-5 M 25 2.50 X10-5 M 25 1.00 X10-5 M	CE CG CG CG	FOTOMTR SPCTR CHNG I2 POLAROGRAPHIC MAXIMUM POLAROGRAPHIC MAXIMUM POLAROGRAPHIC MAXIMUM	MALI CHAN MALI CHAN MALI CHAN MALI CHAN	66023 66023 66023 66023	T L T L T L T L	
COMPOUND NO = 551 MOL WGT — . E O K CL . E O K I . E O K NO3 4 ENTRIES FOR COMPOUND		440.6 PHENYL SULFOSTEARIC ACID 8.0 X10-5 M 25 8.20 X10-5 M 25 1.21 X10-4 M 25 9.80 X10-5 M	CE CG CG CG	FOTOMTR SPCTR CHNG I2 POLAROGRAPHIC MAXIMUM POLAROGRAPHIC MAXIMUM POLAROGRAPHIC MAXIMUM	MALI CHAN MALI CHAN MALI CHAN MALI CHAN	66023 66023 66023 66023	T L T L T L T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality
counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 552 MOL WGT -		454.7 TOLYL SULFOSTEARIC ACID					
. E 0 K CL	UNK	1.00 X10-4 M	CE	FOTOMTR SPCTR CHNG I2	MALI CHAN	66023	T L
. E 0 K I	25	4.898X10-4 M	CG	POLAROGRAPHIC MAXIMUM	MALI CHAN	66023	T L
. E 0 K NO3	25	7.94 X10-5 M	CG	POLAROGRAPHIC MAXIMUM	MALI CHAN	66023	T L
4 ENTRIES FOR COMPOUND	25	2.19 X10-5 M	CG	POLAROGRAPHIC MAXIMUM	MALI CHAN	66023	T L
COMPOUND NO = 553 MOL WGT -		468.7 XYLYL SULFOSTEARIC ACID					
. E 0 K CL	UNK	1.20 X10-4 M	CE	FOTOMTR SPCTR CHNG I2	MALI CHAN	66023	T L
. E 0 K I	25	3.631X10-4 M	CG	POLAROGRAPHIC MAXIMUM	MALI CHAN	66023	T L
. E 0 K NO3	25	5.37 X10-5 M	CG	POLAROGRAPHIC MAXIMUM	MALI CHAN	66023	T L
4 ENTRIES FOR COMPOUND	25	1.29 X10-5 M	CG	POLAROGRAPHIC MAXIMUM	MALI CHAN	66023	T L
COMPOUND NO = 554 MOL WGT -		348.5 SANTOMERSE 3 /SODIUM DODECYL BENZENE SULFONATE/					
	30	5. X10-3 W	HE	QUESTIONABLE CRITERION	BROW ROBI	52013	R
	50	6.5 X10-3 W	HD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
3 ENTRIES FOR COMPOUND				VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 555 MOL WGT -		.0 TERGITOL TMN		QUESTIONABLE CRITERION	BROW ROBI	52013	R
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 556 MOL WGT -		328.3 ARESKAP 100 /MONOBUTYL PHENYLPHENOL SODIUM MONOSULFONATE/					
1 ENTRIES FOR COMPOUND	30	1.4 X10-2 W	HD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 557 MOL WGT -		312.3 ARESKET 300 /MONOBUTYL BIPHENYL SODIUM MONOSULFONATE/					
1 ENTRIES FOR COMPOUND	30	2.2 X10-2 W	HD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 558 MOL WGT -		486.4 ARESKLENE 400 /DIBUTYL PHENYLPHENOL DISODIUMDISULFONATE/					
1 ENTRIES FOR COMPOUND	30	1.7 X10-2 W	HD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 559 MOL WGT -		.0 CATOL 605 /(N-(2-KETO-2-(2,-LAUROYL OXYETHYLAMINO))ETHYL) TRIMETHYLAMMONIUM CHLORIDE/					
1 ENTRIES FOR COMPOUND	30	5.5 X10-3 W	HD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 560 MOL WGT -		.0 EMULSOL 607L (N-(2-KETO-2-(2,-LAUROYLOXETHYL AMINO))ETHYL) PYRIDINIUM CHLORIDE					
3 ENTRIES FOR COMPOUND	0	5. X10-3 W	HE	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
	30	6. X10-3 W	HE	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
	50	8. X10-3 W	HE	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 561 MOL WGT -		320.4 SANTOMERSE D /SODIUM DECYLBENZENE SULFONATE/					
2 ENTRIES FOR COMPOUND	30	2.1 X10-2 W	HD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
	50	1.1 X10-2 W	HD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 562 MOL WGT -		286.3 SODIUM DODECENYL SULFATE					
1 ENTRIES FOR COMPOUND	50	1.6 X10-2 W	CD	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
COMPOUND NO = 563 MOL WGT -		.0 ZEPHIRAN /COCONUT DIMETHYLBENZYL AMMONIUM CHLORIDE/					
2 ENTRIES FOR COMPOUND	30	3. X10-3 W	HE	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L
	50	5. X10-3 W	HE	VAPR PRESURE LOWERING	HUFF MCBA	51004	T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 564 MOL WGT -	50	.0 2.3 X10-3 M	HC	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 565 MOL WGT -	50	.0 8.0 X10-2 D	HC	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 566 MOL WGT -	50	.0 3.0 X10-1 D	HC	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 567 MOL WGT -	50	.0 1.9 X10-1 D	HC	SPECFC CONDCTNCE GRAPH	MARO ELDE	54006	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 568 MOL WGT -	25 30 40 40 40 54 30	555.1 8.8 X10-4 M 1.25 X10-3 M 1.1 X10-3 M 1.0 X10-3 M 1.1 X10-3 M 1.1 X10-3 M 3. X10-4 M	BB BB BC BG BG BC BE	SURFACE TENSION LOG PLOT UNSPECIFIED CONDUCTANCE SURFACE TENSION LOG PLOT VISUAL SPCTR CHNGE PNCN VISUAL SPCTR CHNGE RHD6 SURFACE TENSION LOG PLOT SURFACE TENSION LOG PLOT	MIYA SATA IWAM MIYA MIYA MIYA MIYA SATA IWAM	60029 63034 60029 60029 60029 60029 63034	T L T 3 T L T L T L
1. E-1 M MG SO4 7 ENTRIES FOR COMPOUND							
COMPOUND NO = 569 MOL WGT -	67	618.4 1.1 X10-3 M	BC	SURFACE TENSION LOG PLOT	MIYA	60029	T L
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 570 MOL WGT -	54 67	737.9 1.0 X10-3 M 9.8 X10-4 M	BC BB	SURFACE TENSION LOG PLOT SURFACE TENSION LOG PLOT	MIYA MIYA	60029 60029	T L T L
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 571 MOL WGT -	25 40 40 40 54	585.7 1.1 X10-3 M 1.1 X10-3 M 1.1 X10-3 M 1.2 X10-3 M 1.1 X10-3 M	BC BC BG BG BC	SURFACE TENSION LOG PLOT SURFACE TENSION LOG PLOT VISUAL SPCTR CHNGE RHD6 VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	MIYA MIYA MIYA MIYA MIYA	60029 60029 60029 60029 60029	T L T L T L
5 ENTRIES FOR COMPOUND							
COMPOUND NO = 572 MOL WGT -	30 40 40 30	589.7 1.23 X10-3 M 1.3 X10-3 M 1.0 X10-3 M 3. X10-4 M	BB BG BG BE	UNSPECIFIED CONDUCTANCE VISUAL SPCTR CHNGE RHD6 VISUAL SPCTR CHNGE PNCN SURFACE TENSION LOG PLOT	SATA IWAM MIYA MIYA SATA IWAM	63034 60029 60029 63034	T 3 T L T L T L
1. E-1 M CO SO4 4 ENTRIES FOR COMPOUND							
COMPOUND NO = 573 MOL WGT -	30 40 40 30	594.3 1.20 X10-3 M 1.3 X10-3 M 1.2 X10-3 M 3. X10-4 M	BB BG BG BE	UNSPECIFIED CONDUCTANCE VISUAL SPCTR CHNGE RHD6 VISUAL SPCTR CHNGE PNCN GRAPH DATA NOT RETRIEVED SURFACE TENSION LOG PLOT	SATA IWAM MIYA MIYA SATA IWAM	63034 60029 60029 63034	T 3 T L T L T R
1. E 0 CU SO4 1. E-1 M CU SO4 1. E 0 NA2 SO4 6 ENTRIES FOR COMPOUND				GRAPH DATA NOT RETRIEVED	SATA IWAM	63034	T R
COMPOUND NO = 574 MOL WGT -	40 40	596.1 1.1 X10-3 M 1.1 X10-3 M	BG BG	VISUAL SPCTR CHNGE RHD6 VISUAL SPCTR CHNGE PNCN	MIYA MIYA	60029 60029	T L T L
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 575 MOL WGT -	30	589.5 1.24 X10-3 M	BB	UNSPECIFIED CONDUCTANCE	SATA IWAM	63034	T 3

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC		Qual. Mat.	Method	Authors		Reference	Source	Evaluation
1. E-1 M	NI S04	30	1.1	X10-3 M	BE	METHOD NOT CITED	SATA	IWAM	63034	T	L
3 ENTRIES FOR COMPOUND		30	3.	X10-4 M	BE	SURFACE TENSION LOG PLOT	SATA	IWAM	63034	T	L
COMPOUND NO =	576 MOL WGT -	650.4	CUPRIC	TETRADECYL SULFATE							
1 ENTRIES FOR COMPOUND		47	2.5	X10-4 M	BD	SURFACE TENSION LOG PLOT	SATA	IWAM	63034	T	L
COMPOUND NO =	577 MOL WGT -	706.5	CUPRIC	HEXADECYL SULFATE							
1 ENTRIES FOR COMPOUND		47	5.8	X10-5 M	BC	SURFACE TENSION LOG PLOT	SATA	IWAM	63034	T	L
COMPOUND NO =	578 MOL WGT -	710.6	METHYL /OXYETHYLENE/	11.9 DECANOATE							
	REDUCED POLYDISPERSITY OF HEAD GROUPS										
1. E 1 C	0579	27	1.8	X10-3 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA	KURI	57020	T	L
2.2 E 1 C	0579	27	1.40	X10-3 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA	KURI	57020	T	L
3. E 1 C	0579	27	1.45	X10-3 M	EG	FOTOMTR SPCTR CHNG ERTS	NAKA	KURI	57020	T	L
4. E 1 C	0579	27	1.4	X10-3 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
6. E 1 C	0579	27	1.1	X10-3 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
7. E 1 C	0579	27	8.5	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
10 ENTRIES FOR COMPOUND		27	7.0	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
		27	5.4	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
		27	4.4	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
		27	3.8	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
COMPOUND NO =	579 MOL WGT -	765.1	METHYL /OXYETHYLENE/	12.5 DODECANOATE							
	REDUCED POLYDISPERSITY OF HEAD GROUPS										
		27	3.5	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA	KURI	57020	T	L
		27	3.4	X10-4 M	EG	FOTOMTR SPCTR CHNG ERTS	NAKA	KURI	57020	T	L
		27	2.8	X10-4 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA	KURI	57020	T	L
0578		27	3.4	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA		58017	T	L
5 ENTRIES FOR COMPOUND						SEE CMPD NMBR IN ADDITV	NAKA		58017	X	
COMPOUND NO =	580 MOL WGT -	494.7	METHYL /OXYETHYLENE/	7.0 DECANOATE							
	REDUCED POLYDISPERSITY OF HEAD GROUPS										
		27	1.0	X10-3 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA	KURI	57020	T	L
		27	8.0	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA	KURI	57020	T	L
3 ENTRIES FOR COMPOUND		27	9.5	X10-4 M	EG	FOTOMTR SPCTR CHNG ERTS	NAKA	KURI	57020	T	L
COMPOUND NO =	581 MOL WGT -	640.1	METHYL /OXYETHYLENE/	10.3 DECANOATE							
	REDUCED POLYDISPERSITY OF HEAD GROUPS										
		27	1.4	X10-3 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA	KURI	57020	T	L
		27	1.05	X10-3 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA	KURI	57020	T	L
3 ENTRIES FOR COMPOUND		27	1.15	X10-3 M	EG	FOTOMTR SPCTR CHNG ERTS	NAKA	KURI	57020	T	L
COMPOUND NO =	582 MOL WGT -	891.3	METHYL /OXYETHYLENE/	16.0 DECANOATE							
	NATURAL DISTRIBUTION OF HEAD GROUPS										
		27	1.6	X10-3 M	GG	FOTOMTR SPCTR CHNGE PNCN	NAKA	KURI	57020	T	L
1 ENTRIES FOR COMPOUND											
COMPOUND NO =	583 MOL WGT -	478.7	METHYL /OXYETHYLENE/	6.0 DODECANOATE							
	REDUCED POLYDISPERSITY OF HEAD GROUPS										
		27	1.6	X10-4 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA	KURI	57020	T	L
		27	1.5	X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCN	NAKA	KURI	57020	T	L
3 ENTRIES FOR COMPOUND		27	2.0	X10-4 M	EG	FOTOMTR SPCTR CHNG ERTS	NAKA	KURI	57020	T	L
COMPOUND NO =	584 MOL WGT -	584.5	METHYL /OXYETHYLENE/	8.4 DODECANOATE							
	REDUCED POLYDISPERSITY OF HEAD GROUPS										
		27	2.0	X10-4 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA	KURI	57020	T	L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
	27	2.7 X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCH	NAKA KURI	57020	T L
	27	2.7 X10-4 M	EG	FOTOMTR SPCTR CHNG ERTS	NAKA KURI	57020	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 585 MOL WGT -	707.9	METHYL /OXYETHYLENE/ 11.2 DODECANOATE					
REDUCED POLYDISPERSITY OF HEAD GROUPS							
	27	2.5 X10-4 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA KURI	57020	T L
	27	3.2 X10-4 M	EG	FOTOMTR SPCTR CHNGE PNCH	NAKA KURI	57020	T L
	27	3.1 X10-4 M	EG	FOTOMTR SPCTR CHNG ERTS	NAKA KURI	57020	T L
3 ENTRIES FOR COMPOUND							
COMPOUND NO = 586 MOL WGT -	493.1	METHYL /OXYETHYLENE/ 7.6 OCTANOATE					
REDUCED POLYDISPERSITY OF HEAD GROUPS							
	10	2.00 X10-2 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA INOU	58028	T L
	11	1.25 X10-2 M	EG	FOTOMTR SPCTR CHNGE PNCH	NAKA INOU	58028	T L
	25	9.8 X10-3 M	EG	FOTOMTR SPCTR CHNGE PNCH	NAKA INOU	58028	T L
	25	1.48 X10-2 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA INOU	58028	T L
	40	8.2 X10-3 M	EG	FOTOMTR SPCTR CHNGE PNCH	NAKA INOU	58028	T L
	40	1.28 X10-2 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA INOU	58028	T L
	43	7.8 X10-3 M	EG	FOTOMTR SPCTR CHNGE PNCH	NAKA INOU	58028	T L
	43	1.20 X10-2 M	ED	FOTOMTR SOLUBLZTN SDN 4	NAKA INOU	58028	T L
8 ENTRIES FOR COMPOUND							
COMPOUND NO = 587 MOL WGT -	307.5	DECYL DIMETHYLAMMONIOPROPANE SULFONATE					
	30	1.20 X10 0 D	BB	TURBIDITY PLT LITE SCATR	HERR	66013	T 3
		3.902X10-2 M					M
	30	1.1 X10 0 P	BD	DENSITY	BENJ	66040	T L
		3.57 X10-2 S					M
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 588 MOL WGT -	335.6	DODECYL DIMETHYLAMMONIOPROPANE SULFONATE					
	30	1.2 X10-1 D	BC	TURBIDITY PLT LITE SCATR	HERR	66013	T 3
		3.57 X10-3 M					M
	30	1.2 X10-1 P	BD	DENSITY	BENJ	66040	T L
		3.57 X10-3 S					M
2. E-1 M NA CL	30	1.0 X10-1 D	BC	METHOD NOT CITED	HERR	66013	G 3
1. E 0 M NA CL	30	2.97 X10-3 M					M
	30	5.8 X10-2 D	BC	METHOD NOT CITED	HERR	66013	G 3
		1.72 X10-3 M					M
4 ENTRIES FOR COMPOUND							
COMPOUND NO = 589 MOL WGT -	391.7	HEXADECYL DIMETHYLAMMONIOPROPANE SULFONATE					
	30	2.4 X10-3 D	BD	TURBIDITY PLT LITE SCATR	HERR	66013	T L
		6.12 X10-5 M					M
	30	1.2 X10-3 D	BD	SURFACE TENSION LOG PLOT	HERR	66013	T L
		3.06 X10-5 M					M
2 ENTRIES FOR COMPOUND							
COMPOUND NO = 590 MOL WGT -	299.6	DODECYL DIMETHYLAMMONIOPROPANE CARBOXYLATE					
	30	1.6 X10-1 D	BC	TURBIDITY PLT LITE SCATR	HERR	66013	T 3
		5.34 X10-3 M					M
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 591 MOL WGT -	352.5	DODECYL DIMETHYLPHOSPHONIOPROPANE SULFONATE					
	30	8.5 X10-2 D	BC	TURBIDITY PLT LITE SCATR	HERR	66013	T 3
		2.41 X10-3 M					M
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 592 MOL WGT -	299.6	DODECYL N-DIETHYL N-BETAINE					
	30	8.4 X10-2 D	XC	METHOD NOT CITED	HERR	66013	T L
		2.80 X10-3 M					M
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 593 MOL WGT -	351.6	DIMETHYL DODECYLAMMONIOPROPANE HYDROXY SULFONATE					
	30	7.0 X10-2 D	BC	METHOD NOT CITED	HERR	66013	T 3
		1.99 X10-3 M					M
1 ENTRIES FOR COMPOUND							

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 594 MOL WGT -	391.7 30 1.78 X10-3 M	DODECYL DIPROPYL AMMONIOPROPANE SULFONATE X10-2 D BC METHOD NOT CITED		HERR	66013	T 3 M	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 595 MOL WGT - 2. E-1 M NA P-TOLUENE-SO3	351.6 30 5.68 X10-4 M	DODECYL DIMETHYL AMMONIOPROPANE SULFATE X10-2 D BC METHOD NOT CITED		HERR	66013	T L M	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 596 MOL WGT -	353.6 50 2.2 X10-4 M	TETRAHYDROANACARDOL AMMONIUM MONOSULFONATE X10-3 D XG VELOCITY OF SOUND		KUPP SURY	65028	T L M	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 597 MOL WGT -	346.5 UNK 2.69 X10-3 M	SODIUM DODECYL MONO-OXYPROPYL SULFATE CG VISUAL SPCTR CHNGE PNCN		WEIL STIR	66003	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 598 MOL WGT -	374.5 UNK 5.8 X10-4 M	SODIUM TETRADECYL MONO-OXYPROPYL SULFATE CG VISUAL SPCTR CHNGE PNCN		WEIL STIR	66003	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 599 MOL WGT -	432.6 UNK 3.6 X10-4 M	SODIUM TETRADECYL DI-OXYPROPYL SULFATE CG VISUAL SPCTR CHNGE PNCN		WEIL STIR	66003	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 600 MOL WGT -	402.6 UNK 1.6 X10-4 M	SODIUM HEXADECYL MONO-OXYPROPYL SULFATE CG VISUAL SPCTR CHNGE PNCN		WEIL STIR	66003	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 601 MOL WGT -	430.6 UNK 7. X10-5 M	SODIUM OCTADECYL MONO-OXYPROPYL SULFATE CG VISUAL SPCTR CHNGE PNCN		WEIL STIR	66003	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 602 MOL WGT -	280.4 UNK 1.30 X10-2 M	ALPHA SULFO LAURIC ACID CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 603 MOL WGT -	344.4 UNK 5.3 X10-3 M	SODIUM PROPYL ALPHA SULFO LAURATE CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 604 MOL WGT -	344.4 UNK 2.8 X10-3 M	SODIUM METHYL ALPHA SULFO MYRISTATE CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 605 MOL WGT -	330.4 UNK 1.82 X10-2 M	SODIUM AMYL ALPHAPHOSPHONO PELARGONATE CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 606 MOL WGT -	344.4 UNK 9.0 X10-3 M	SODIUM AMYL ALPHAPHOSPHONO CAPRATE CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 607 MOL WGT -	344.4 UNK 6.68 X10-3 M	SODIUM ISOPROPYL ALPHAPHOSPHONO LAURATE CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L	
1 ENTRIES FOR COMPOUND							
COMPOUND NO = 608 MOL WGT -	344.4 UNK 7.26 X10-3 M	SODIUM METHYL ALPHAPHOSPHONO MYRISTATE CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L	
1 ENTRIES FOR COMPOUND							

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 609 MOL WGT - 1 ENTRIES FOR COMPOUND	400.5 UNK	SODIUM ISOPROPYL ALPHAPHOSPHONO PALMITATE X10-4 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L		
COMPOUND NO = 610 MOL WGT - 1 ENTRIES FOR COMPOUND	400.5 UNK	SODIUM METHYL ALPHAPHOSPHONO STEARATE X10-4 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	T L		
COMPOUND NO = 611 MOL WGT - 1 ENTRIES FOR COMPOUND	238.2 UNK	ALPHAPHOSPHONO PELARGONIC ACID X10-2 M CB SURFACE TENSION UNSPEC		MAUR STIR	64002	LT L		
COMPOUND NO = 612 MOL WGT - 1 ENTRIES FOR COMPOUND	252.3 UNK	ALPHAPHOSPHONO DECANOIC ACID X10-2 M CB SURFACE TENSION UNSPEC		MAUR STIR	64002	LT L		
COMPOUND NO = 613 MOL WGT - 1 ENTRIES FOR COMPOUND	280.3 UNK	ALPHAPHOSPHONO DODECANOIC ACID X10-3 M CC SURFACE TENSION UNSPEC		MAUR STIR	64002	LT L		
COMPOUND NO = 614 MOL WGT - 1 ENTRIES FOR COMPOUND	308.4 UNK	ALPHAPHOSPHONO TETRADECANOIC ACID X10-4 M CB SURFACE TENSION UNSPEC		MAUR STIR	64002	LT L		
COMPOUND NO = 615 MOL WGT - 1 ENTRIES FOR COMPOUND	336.5 UNK	ALPHAPHOSPHONO HEXADECANOIC ACID X10-5 M CB SURFACE TENSION UNSPEC		MAUR STIR	64002	LT L		
COMPOUND NO = 616 MOL WGT - 1 ENTRIES FOR COMPOUND	274.3 UNK	MONOSODIUM ALPHAPHOSPHONO DECANOATE X10-2 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 617 MOL WGT - 1 ENTRIES FOR COMPOUND	302.3 UNK	MONOSODIUM ALPHAPHOSPHONO DODECANOATE X10-2 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 618 MOL WGT - 1 ENTRIES FOR COMPOUND	330.4 UNK	MONOSODIUM ALPHAPHOSPHONO TETRADECANOATE X10-3 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 619 MOL WGT - 1 ENTRIES FOR COMPOUND	358.4 UNK	MONOSODIUM ALPHAPHOSPHONO HEXADECANOATE X10-4 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 620 MOL WGT - 1 ENTRIES FOR COMPOUND	324.3 UNK	DISODIUM ALPHAPHOSPHONO DODECANOATE X10-2 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 621 MOL WGT - 1 ENTRIES FOR COMPOUND	352.4 UNK	DISODIUM ALPHAPHOSPHONO TETRADECANOATE X10-2 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 622 MOL WGT - 1 ENTRIES FOR COMPOUND	380.4 UNK	DISODIUM ALPHAPHOSPHONO HEXADECANOATE X10-3 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 623 MOL WGT - 1 ENTRIES FOR COMPOUND	408.5 UNK	DISODIUM ALPHAPHOSPHONO OCTADECANOATE X10-3 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		
COMPOUND NO = 624 MOL WGT - 1 ENTRIES FOR COMPOUND	374.3 UNK	TRISODIUM ALPHAPHOSPHONO TETRADECANOATE X10-2 M CG VISUAL SPCTR CHNGE PNCN		MAUR STIR	64002	L L		

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 625 MOL WGT - 1 ENTRIES FOR COMPOUND	402.4 UNK	TRISODIUM ALPHAPHOSPHONO HEXADECANOATE 1.62 X10-2 M CG VISUAL SPCTR CHNGE PNCN			MAUR STIR	64002	L L
COMPOUND NO = 626 MOL WGT - 1 ENTRIES FOR COMPOUND	430.5 UNK	TRISODIUM ALPHAPHOSPHONO OCTADECANOATE 8.2 X10-3 M CG VISUAL SPCTR CHNGE PNCN			MAUR STIR	64002	L L
COMPOUND NO = 627 MOL WGT - 1 ENTRIES FOR COMPOUND	332.2 25	CESIUM DODECANOATE 2.5 X10-2 M XG VISUAL SPCTR CHNGE			KLEV	53010	T L
COMPOUND NO = 628 MOL WGT - 1 ENTRIES FOR COMPOUND	266.3 30	DODECYLAMMONIUM BROMIDE 1.2 X10-2 M XG VISUAL SPCTR CHNGE			KLEV	53010	T L
COMPOUND NO = 629 MOL WGT - 1 ENTRIES FOR COMPOUND	320.6 50	POTASSIUM ELAIDATE/TRANS-9-OCTADECENOATE/ 1.5 X10-3 M XC REFRACTIVE INDEX			KLEV	53010	T L
COMPOUND NO = 630 MOL WGT - 1 ENTRIES FOR COMPOUND	336.6 55	POTASSIUM RICINOLEATE/12 HYDROXY OLEATE/ 3.6 X10-3 M XC REFRACTIVE INDEX			KLEV	53010	T L
COMPOUND NO = 631 MOL WGT - 1 ENTRIES FOR COMPOUND	336.6 55	POTASSIUM RICINELAIDATE/12 HYDROXY ELAIDATE/ 5.5 X10-3 M XB REFRACTIVE INDEX			KLEV	53010	T L
COMPOUND NO = 632 MOL WGT - 2 ENTRIES FOR COMPOUND	295.6 35 50	POTASSIUM N-DODECYL BETA-ALANINATE 2.6 X10-3 M XC REFRACTIVE INDEX 3.0 X10-3 M XC REFRACTIVE INDEX			KLEV	53010	T L
COMPOUND NO = 633 MOL WGT - 1 ENTRIES FOR COMPOUND	292.9 30	N-DODECYL BETA-ALANINE HYDROCHLORIDE 1.0 X10-2 M XC REFRACTIVE INDEX			KLEV	53010	T L
COMPOUND NO = 634 MOL WGT - 6.75 E-2 M NA CL 3 ENTRIES FOR COMPOUND	304.5 40 50 40	POTASSIUM DODECYL SULFATE 7.8 X10-3 M CB SPECFC CONDCTNCE GRAPH 5.2 X10-3 M DG VISUAL SPCTR CHNGE PNCN 1.5 X10-3 M CG FOTOMTR SPCTR CHNGE RHD6			MEGU KOND RAIS MEGU KOND	56020 52016 56020	T 3 T L T L
COMPOUND NO = 635 MOL WGT - 1 ENTRIES FOR COMPOUND	256.3 50	LITHIUM DODECYL SULFONATE 1.00 X10-2 M DG VISUAL SPCTR CHNGE PNCN			RAIS	52016	T L
COMPOUND NO = 636 MOL WGT - 1 ENTRIES FOR COMPOUND	304.4 50	SODIUM DODECYL THIOSULFATE 3.0 X10-3 M DG VISUAL SPCTR CHNGE PNCN			RAIS	52016	T L
COMPOUND NO = 637 MOL WGT - 1 ENTRIES FOR COMPOUND	300.4 50	LITHIUM TETRADECYL SULFATE 1.64 X10-3 M DG VISUAL SPCTR CHNGE PNCN			RAIS	52016	T L
COMPOUND NO = 638 MOL WGT - 1 ENTRIES FOR COMPOUND	328.4 50	LITHIUM HEXADECYL SULFATE 2.9 X10-4 M DG VISUAL SPCTR CHNGE PNCN			RAIS	52016	T L
COMPOUND NO = 639 MOL WGT - 1 ENTRIES FOR COMPOUND	290.3 50	SODIUM UNDECYL THIOSULFATE 7.0 X10-3 M DG VISUAL SPCTR CHNGE PNCN			RAIS	52016	T L

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Detailson page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives		Temp. °C	CMC		Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 640	MOL WGT -	325.5	HEXYL TRIMETHYLAMMONIUM HEXANE SULFATE							
1 ENTRIES FOR COMPOUND		25	1.1 X10-1 M	BC	METHOD NOT CITED		CORK GOOD	66014	T 3	
COMPOUND NO = 641	MOL WGT -	353.6	HEXYL TRIMETHYLAMMONIUM OCTANE SULFATE							
1.00 E 2 I NA BR		25	2.7 X10-2 M	BC	METHOD NOT CITED		CORK GOOD	66014	T 3	
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 642	MOL WGT -	409.7	OCTYL TRIMETHYLAMMONIUM DECANE SULFATE							
1 ENTRIES FOR COMPOUND		25	1.9 X10-3 M	BC	METHOD NOT CITED		CORK GOOD	66014	T 3	
COMPOUND NO = 643	MOL WGT -	437.8	DODECYL TRIMETHYLAMMONIUM OCTANE SULFATE							
1.00 E 2 I NA BR		25	4.5 X10-4 M	BC	METHOD NOT CITED		CORK GOOD	66014	T 3	
1 ENTRIES FOR COMPOUND										
COMPOUND NO = 644	MOL WGT -	309.5	HEXYL TRIMETHYLAMMONIUM HEXANE SULFONATE							
1 ENTRIES FOR COMPOUND		25	2.2 X10-1 M	BC	METHOD NOT CITED		CORK GOOD	66014	T L	
COMPOUND NO = 645	MOL WGT -	268.3	PARA/BETA-D-GLUCOSYL/ETHYLBENZENE							
1 ENTRIES FOR COMPOUND		25	5.0 X10-1 M	CE	SURFACE TENSION LOG PLOT	HUTC SHEA		64037	T L	
COMPOUND NO = 646	MOL WGT -	282.4	PARA/BETA-D-GLUCOSYL/PROPYL BENZENE							
1 ENTRIES FOR COMPOUND		25	1.3 X10-1 M	CC	SURFACE TENSION LOG PLOT	HUTC SHEA		64037	T L	
COMPOUND NO = 647	MOL WGT -	296.4	PARA/BETA-D-GLUCOSYL/BUTYLBENZENE							
1 ENTRIES FOR COMPOUND		25	5. X10-2 M	CD	SURFACE TENSION LOG PLOT	HUTC SHEA		64037	T L	
COMPOUND NO = 648	MOL WGT -	276.4	ALPHA-D-GLUCOSYL OCTANE							
2 ENTRIES FOR COMPOUND		25	7.9 X10-3 M	CC	SURFACE TENSION LOG PLOT	HUTC SHEA		64037	T L	
COMPOUND NO = 648	MOL WGT -	276.4	ALPHA-D-GLUCOSYL OCTANE							
2 ENTRIES FOR COMPOUND		50	7.1 X10-3 M	CC	SURFACE TENSION LOG PLOT	HUTC SHEA		64037	T L	
COMPOUND NO = 650	MOL WGT -	266.4	PARA/BETA-D-XYLOSYL/BUTYL BENZENE							
1 ENTRIES FOR COMPOUND		25	6. X10-4 M	CD	SURFACE TENSION LOG PLOT	HUTC SHEA		64037	T L	
COMPOUND NO = 651	MOL WGT -	.0	EMASOL 1120 /ALKYL POLYOXYETHYLENE SORBITAN ESTER/							
2 ENTRIES FOR COMPOUND		25	1.1 X10-2 M	CC	SURFACE TENSION LOG PLOT	HUTC SHEA		64037	T L	
COMPOUND NO = 651	MOL WGT -	.0	EMASOL 1120 /ALKYL POLYOXYETHYLENE SORBITAN ESTER/							
2 ENTRIES FOR COMPOUND		30	7.5 X10-1 D	HG	VISCOSITY MINIMUM	OKUY TYUZ		54008	T L	
COMPOUND NO = 652	MOL WGT -	.0	EMASOL 1130 /ALKYL POLYOXYETHYLENE SORBITAN ESTER/							
1 ENTRIES FOR COMPOUND		30	5.0 X10-1 D	HG	VISCOSITY MINIMUM	OKUY TYUZ		54008	T L	
COMPOUND NO = 653	MOL WGT -	.0	EMULGEN 120 /ALKYL POLYOXYETHYLENE ETHER/							
1 ENTRIES FOR COMPOUND		30	3.1 X10-1 D	HG	VISCOSITY MINIMUM	OKUY TYUZ		54008	T L	
COMPOUND NO = 654	MOL WGT -	374.7	OCTADECYL TRIMETHYLAMMONIUM NITRATE							
5.01 E 0 H	METHANOL	25	2.3 X10-4 M	BE	EQUIV CONDCTNCE GRAPH	GRIE KRAU		48010	T L	
1.227E 1 H	METHANOL	25	1.76 X10-4 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU		48028	P 2	
1.508E 1 H	METHANOL	25	3.28 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU		48028	P 3	
1.985E 1 H	METHANOL	25	4.37 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU		48028	P 3	
2.589E 1 H	METHANOL	25	5.93 X10-4 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU		48028	P 2	
3.463E 1 H	METHANOL	25	1.10 X10-3 M	BA	EQUIV CONDCTNCE GRAPH	EVER KRAU		48028	P 2	
7 ENTRIES FOR COMPOUND		25	3.03 X10-3 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU		48028	P 3	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality; counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l (or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives		Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 655 MOL WGT -		368.1	OCTADECYL PYRIDINIUM CHLORIDE						
		25	2.4 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	GRIE KRAU	48010	T L	
		25	2.40 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 3	
		25	2.54 X10-4 M	BC	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	T L	
3 ENTRIES FOR COMPOUND									
COMPOUND NO = 656 MOL WGT -		394.7	OCTADECYL PYRIDINIUM NITRATE						
2.0 E 1 H METHANOL		25	1.28 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	T 3	
2 ENTRIES FOR COMPOUND		25	5.76 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 3	
COMPOUND NO = 657 MOL WGT -		412.6	OCTADECYL PYRIDINIUM BROMIDE						
2.0 E 1 H METHANOL		25	6.10 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	EVER KRAU	48028	P 3	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 658 MOL WGT -		440.6	OCTADECYL TRIMETHYLLAMMONIUM BROMATE						
		25	3.1 X10-4 M	BG	EQUIV COND 1ST DEVIATION	GRIE KRAU	48010	T L	
		25	3.31 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	GRIE KRAU	48010	P 3	
2 ENTRIES FOR COMPOUND									
COMPOUND NO = 659 MOL WGT -		357.7	OCTADECYL TRIMETHYLLAMMONIUM FORMATE						
		25	4.4 X10-4 M	BC	EQUIV CONDCTNCE GRAPH	GRIE KRAU	48010	T 3	
1 ENTRIES FOR COMPOUND									
COMPOUND NO = 660 MOL WGT -		479.5	HEXADECYL PYRIDINIUM IODATE						
		25	1.6 X10-3 M	BG	EQUIV CONDCTNCE MAXIMUM	BROW GRIE	49014	T L	
7.9 E 0 H TERTIARY BUTANOL		25	9.9 X10-4 M	BG	EQUIV COND MAX BEGINING	BROW GRIE	49014	T L	
7.9 E 0 H TERTIARY BUTANOL		25	1.35 X10-4 M	BG	EQUIV COND MAX BEGINING	BROW GRIE	49014	T L	
4 ENTRIES FOR COMPOUND		25	9. X10-4 M	BG	EQUIV CONDCTNCE MAXIMUM	BROW GRIE	49014	T L	
COMPOUND NO = 661 MOL WGT -		507.6	OCTADECYL PYRIDINIUM IODATE						
		25	5. X10-4 M	BG	EQUIV CONDCTNCE MAXIMUM	BROW GRIE	49014	T L	
1.62 E 1 H METHANOL		25	9. X10-5 M	BG	EQUIV COND MAX BEGINING	BROW GRIE	49014	T L	
1.62 E 1 H METHANOL		25	8. X10-4 M	BG	EQUIV CONDCTNCE MAXIMUM	BROW GRIE	49014	T L	
4 ENTRIES FOR COMPOUND		25	2.9 X10-4 M	BG	EQUIV COND MAX BEGINING	BROW GRIE	49014	T L	
COMPOUND NO = 662 MOL WGT -		482.7	OCTADECYL TRIETHYLLAMMONIUM BROMATE						
1 ENTRIES FOR COMPOUND		25	2.5 X10-4 M	BC	EQUIV CONDCTNCE GRAPH	MCDO KRAU	51009	T 3	
COMPOUND NO = 663 MOL WGT -		524.8	OCTADECYL TRIPROPYLLAMMONIUM BROMATE						
1 ENTRIES FOR COMPOUND		25	1.25 X10-4 M	BG	EQUIV COND MAX BEGINING	MCDO KRAU	51009	T L	
COMPOUND NO = 664 MOL WGT -		566.9	OCTADECYL TRIBUTYLLAMMONIUM BROMATE						
1 ENTRIES FOR COMPOUND		25	5.3 X10-5 M	BG	EQUIV COND MAX BEGINING	MCDO KRAU	51009	TK L	
COMPOUND NO = 665 MOL WGT -		608.9	OCTADECYL TRIAMYLAMMONIUM BROMATE						
1 ENTRIES FOR COMPOUND		25	1.6 X10-5 M	BG	EQUIV COND MAX BEGINING	MCDO KRAU	51009	TK L	
COMPOUND NO = 666 MOL WGT -		538.8	HEXADECYL TRIBUTYLLAMMONIUM BROMATE						
1 ENTRIES FOR COMPOUND		25	3.3 X10-4 M	BG	EQUIV COND MAX BEGINING	MCDO KRAU	51009	TK L	
COMPOUND NO = 667 MOL WGT -		713.4	OCTADECYL TRIMETHYLLAMMONIUM OXALATE						
9.91 E 0 H ACETONE		25	6.4 X10-5 M	BE	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	T L	
9.91 E 0 H ACETONE		25	1.44 X10-4 M	BG	EQUIV COND 1ST DEVIATION	YOUN GRIE	49017	T L	
1.39 E 1 H ACETONE		25	1.61 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	P 3	
1.39 E 1 H ACETONE		25	2.56 X10-4 M	BG	EQUIV COND 1ST DEVIATION	YOUN GRIE	49017	T L	
2.07 E 1 H ACETONE		25	2.56 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	P 3	
		25	5.29 X10-4 M	BG	EQUIV COND 1ST DEVIATION	YOUN GRIE	49017	T L	

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source	Evaluation
2.07 E 1 H ACETONE	25	6.50 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	P	3
2.88 E 1 H ACETONE	25	1.30 X10-3 M	BG	EQUIV COND 1ST DEVIATION	YOUN GRIE	49017	T	L
2.88 E 1 H ACETONE	25	1.94 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	P	3
2.07 E 1 H METHANOL	25	2.50 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	P	3
2.07 E 1 H METHANOL	25	2.70 X10-4 M	BE	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	T	L
3.02 E 1 H METHANOL	25	6.8 X10-4 M	BE	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	T	L
3.02 E 1 H METHANOL	25	6.25 X10-4 M	BB	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	P	3
4.01 E 1 H METHANOL	25	1.76 X10-3 M	BE	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	T	L
4.01 E 1 H METHANOL	25	2.20 X10-3 M	BC	EQUIV CONDCTNCE GRAPH	YOUN GRIE	49017	P	3
15 ENTRIES FOR COMPOUND								
COMPOUND NO = 668 MOL WGT -	292.5	DIPOTASSIUM OCTYL MALONATE						
	20	3.0 X10-1 M	BC	SURFACE TENSION UNSPEC	SHIN	55007	T	3
	25	3.5 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	T	L
7.6 E-1 K K ION	25	3.2 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
7.7 E-1 K K ION	25	3.0 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
8.2 E-1 K K ION	25	2.8 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
9.2 E-1 K K ION	25	2.6 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
1.00 E 0 K K ION	25	2.3 X10-1 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
7 ENTRIES FOR COMPOUND								
COMPOUND NO = 669 MOL WGT -	320.5	DIPOTASSIUM DECYL MALONATE						
	25		BG	THEORETICALLY ESTIMATED	SHIN	55007	T	R
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 670 MOL WGT -	348.6	DIPOTASSIUM DODECYL MALONATE						
	20	4.8 X10-2 M	BC	SURFACE TENSION UNSPEC	SHIN	55007	T	L
	20	4.8 X10-2 M	BG	FOTOMTR SPCTR CHNGE PNCN	SHIN	55007	T	L
	25	4.8 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	T	L
	25	5.0 X10-2 M	BB	FOTOMTR SOLUBLZTN 2NPA	SHIN	55007	T	3
1.10 E-1 K K ION	25	4.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
1.16 E-1 K K ION	25	3.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
1.31 E-1 K K ION	25	3.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
1.58 E-1 K K ION	25	2.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
1.89 E-1 K K ION	25	2.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
2.17 E-1 K K ION	25	1.9 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
2.56 E-1 K K ION	25	1.6 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
3.08 E-1 K K ION	25	1.3 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
3.48 E-1 K K ION	25	1.1 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
13 ENTRIES FOR COMPOUND								
COMPOUND NO = 671 MOL WGT -	376.7	DIPOTASSIUM TETRADECYL MALONATE						
	20	1.9 X10-2 M	BC	SURFACE TENSION UNSPEC	SHIN	55007	T	L
	25	1.7 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	T	L
	25	1.8 X10-2 M	BC	FOTOMTR SOLUBLZTN 2NPA	SHIN	55007	T	3
3.7 E-2 K K ION	25	1.5 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
4.0 E-2 K K ION	25	1.4 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
4.5 E-2 K K ION	25	1.2 X10-2 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
5.4 E-2 K K ION	25	9.6 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
7.0 E-2 K K ION	25	7.1 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
9.7 E-2 K K ION	25	5.4 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
1.23 E-1 K K ION	25	4.4 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
10 ENTRIES FOR COMPOUND								
COMPOUND NO = 672 MOL WGT -	404.7	DIPOTASSIUM HEXADECYL MALONATE						
	20	9. X10-3 M	BD	SURFACE TENSION UNSPEC	SHIN	55007	T	L
	25	6.3 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	T	L
F.48 E-2 K K ION	25	5.5 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
1.86 E-2 K K ION	25	4.4 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
2.23 E-2 K K ION	25	3.2 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
2.97 E-2 K K ION	25	2.2 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
4.75 E-2 K K ION	25	1.5 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	G	L
7 ENTRIES FOR COMPOUND								
COMPOUND NO = 673 MOL WGT -	432.8	DIPOTASSIUM OCTADECYL MALONATE						
	25	2.3 X10-3 M	BG	VISUAL SPCTR CHNGE PNCN	SHIN	55007	T	L
1 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/l or kg; W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 674 MOL WGT - BRANCHED CHAIN, NATURAL OE DISTRIBUTION	470.7	DIISOBUTYL BENZENE/OXYETHYLENE/6 ALCOHOL				(DB)		
	20	1.8 X10-3 M	HG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	1.4 X10-3 M	HG	FOTOMTR SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 675 MOL WGT - NATURAL DISTRIBUTION OF HEAD GROUPS	647.0	OCTYL BENZENE /OXYETHYLENE/10 ALCOHOL						
	20	4.9 X10-4 M	HG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	3.8 X10-4 M	HG	FOTOMTR SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 676 MOL WGT -	292.3	SODIUM 1,1,3,3-TETRAMETHYL BUTYL BENZENE SULFONATE						
	20	8.4 X10-3 M	XG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	1.00 X10-2 M	XG	VISUAL SPCTR CHNGE RHD6	TAUB KONS	60033	T L	
	40	8.4 X10-3 M	XC	UNSPECIFIED CONDUCTANCE	TAUB KONS	60033	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 677 MOL WGT -	404.6	SODIUM DI-/1,1,3,3-TETRAMETHYL BUTYL/BENZENESULFONATE						
	20	1.0 X10-3 M	XG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	1.0 X10-3 M	XG	VISUAL SPCTR CHNGE RHD6	TAUB KONS	60033	T L	
	40	9. X10-4 M	XD	UNSPECIFIED CONDUCTANCE	TAUB KONS	60033	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 678 MOL WGT -	334.4	SODIUM 1,1-DIMETHYLNONYL BENZENE SULFONATE						
	20	3.0 X10-3 M	XG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	2.4 X10-3 M	XG	VISUAL SPCTR CHNGE RHD6	TAUB KONS	60033	T L	
	40	3.3 X10-3 M	XC	UNSPECIFIED CONDUCTANCE	TAUB KONS	60033	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 679 MOL WGT -	334.4	SODIUM 3-BUTYL HEPTYL BENZENE SULFONATE						
	20	2.7 X10-3 M	XG	VISUAL SPCTR CHNGE RHD6	TAUB KONS	60033	T L	
	20	3.3 X10-3 M	XG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	40	3.3 X10-3 M	XC	UNSPECIFIED CONDUCTANCE	TAUB KONS	60033	T L	
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 680 MOL WGT -	348.5	SODIUM 2-PROPYL-4-METHYL-OCTYL BENZENE SULFONATE						
	20	1.7 X10-3 M	XG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
	20	1.7 X10-3 M	XG	VISUAL SPCTR CHNGE RHD6	TAUB KONS	60033	T L	
	40	1.6 X10-3 M	XD	UNSPECIFIED CONDUCTANCE	TAUB KONS	60033	T L	
1.33 E 2 Q NA2 SO4	20	1.1 X10-3 M	XG	VISUAL SPCTR CHNGE PNCN	TAUB KONS	60033	T L	
1.33 E 2 Q NA2 SI03 META								
1.33 E 2 Q NA2 CO3								
1.33 E 2 Q NA2 SO4	20	1.1 X10-3 M	XG	VISUAL SPCTR CHNGE RHD6	TAUB KONS	60033	T L	
1.33 E 2 Q NA2 SI03 META								
1.33 E 2 Q NA2 CO3								
1.33 E 2 Q NA2 SO4	40	1.1 X10-3 M	XD	UNSPECIFIED CONDUCTANCE	TAUB KONS	60033	T L	
1.33 E 2 Q NA2 SI03 META								
1.33 E 2 Q NA2 CO3								
6 ENTRIES FOR COMPOUND								
COMPOUND NO = 681 MOL WGT -	292.3	SODIUM DIBUTYL BENZENE SULFONATE						
1 ENTRIES FOR COMPOUND	UNK	1.6 X10-4 M	CD	TURBIDITY PLT LITE SCATR	YURZ KUCH	52018	T L	
COMPOUND NO = 682 MOL WGT -	342.4	SODIUM DIBUTYL NAPHTHALENE SULFONATE					/NEKAL/	
1 ENTRIES FOR COMPOUND	UNK	2.9 X10-4 M	HC	TURBIDITY PLT LITE SCATR	YURZ KUCH	52018	T L	
COMPOUND NO = 683 MOL WGT -	460.7	SODIUM EICOSYLBENZENE SULFONATE						
1 ENTRIES FOR COMPOUND	UNK	1.7 X10-5 M	HE	TURBIDITY PLT LITE SCATR	YURZ KUCH	52018	T L	
COMPOUND NO = 684 MOL WGT -	178.3	HEXYL SULFINYLETHANOL						
1 ENTRIES FOR COMPOUND	25	2.5 X10-1 W	BD	VAPR PRESSURE LOWERING	CORK GOOD	66015	E L	

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/(l or kg); W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat.	Method	Authors	Reference	Source	Evaluation
COMPOUND NO = 685 MOL WGT -	192.3 25	HEXYL SULFINYLPROPANOL X10-1 W BD	VAPR PRESSURE LOWERING		CORK GOOD	66015	E L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 686 MOL WGT -	206.3 25	HEXYL SULFINYLBUTANOL X10-1 W BD	VAPR PRESSURE LOWERING		CORK GOOD	66015	E L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 687 MOL WGT -	220.4 25	HEXYL SULFINYLPENTANOL X10-1 W BD	VAPR PRESSURE LOWERING		CORK GOOD	66015	E L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 688 MOL WGT -	206.3 25 40	OCTYL SULFINYLETHANOL X10-2 W BD X10-2 W BD	SURFACE TENSION LOG PLOT SURFACE TENSION LOG PLOT		CORK GOOD CORK GOOD	66015 66015	E L E L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 689 MOL WGT -	220.4 25 40	OCTYL SULFINYLPROPANOL X10-2 W BD X10-2 W BD	SURFACE TENSION LOG PLOT SURFACE TENSION LOG PLOT		CORK GOOD CORK GOOD	66015 66015	E L E L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 690 MOL WGT -	234.4 25 40	OCTYL SULFINYLBUTANOL X10-2 W BD X10-2 W BD	SURFACE TENSION LOG PLOT SURFACE TENSION LOG PLOT		CORK GOOD CORK GOOD	66015 66015	E L E L	
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 691 MOL WGT -	176.3 25	OCTYL METHYL SULFOXIDE X10-2 W BD	SURFACE TENSION LOG PLOT		CORK GOOD	66015	E L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 692 MOL WGT -	234.4 25	DECYL SULFINYLETHANOL X10-3 W BD	SURFACE TENSION LOG PLOT		CORK GOOD	66015	E L	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 693 MOL WGT -	354.1 1 ENTRIES FOR COMPOUND	N-CETYL 2-METHYL PYRIDINIUM CHLORIDE QUESTIONABLE CRITERION			ANGE NICO	61029	R	
COMPOUND NO = 694 MOL WGT -	354.1 1 ENTRIES FOR COMPOUND	N-CETYL 3-METHYL PYRIDINIUM CHLORIDE QUESTIONABLE CRITERION			ANGE NICO	61029	R	
COMPOUND NO = 695 MOL WGT -	354.1 20 1.9	N-CETYL 4-METHYL PYRIDINIUM CHLORIDE X10-2 D GE X10-3 M	KRAFFT POINT SOLUBILITY		ANGE NICO	61029	T L M	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 696 MOL WGT -	445.5 65 8.9	N-CETYL 2-METHYL PYRIDINIUM IODIDE X10-3 D GE X10-5 M	KRAFFT POINT SOLUBILITY		ANGE NICO	61029	T L M	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 697 MOL WGT -	445.5 34 4.4	N-CETYL-3-METHYL PYRIDINIUM IODIDE X10-3 D GE X10-5 M	KRAFFT POINT SOLUBILITY		ANGE NICO	61029	T L M	
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 698 MOL WGT -	445.5 30 1.7	N-CETYL-4-METHYL PYRIDINIUM IODIDE X10-2 D GE X10-3 M	KRAFFT POINT SOLUBILITY		ANGE NICO	61029	T L M	
1 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture; D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values – Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation
COMPOUND NO = 699 MOL WGT – 1 ENTRIES FOR COMPOUND		110.1 UNK 3.5	SODIUM BUTYRATE X10 0 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 700 MOL WGT – 1 ENTRIES FOR COMPOUND		116.2 UNK 1.	HEXANOIC ACID X10-1 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 701 MOL WGT – 1 ENTRIES FOR COMPOUND		352.2 UNK 5.	POTASSIUM PERFLUROHEXANOATE X10-1 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 702 MOL WGT – 1 ENTRIES FOR COMPOUND		552.2 UNK 9.	POTASSIUM PERFLUORODECANOATE X10-4 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 703 MOL WGT – E 0 PH OF SOLUTION 2 ENTRIES FOR COMPOUND		194.3 UNK 4.	4-HEXYL RESORCINOL X10-3 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
		UNK		GRAPH DATA NOT RETRIEVED	KLEV RAIS	54004	R
COMPOUND NO = 704 MOL WGT – 1 ENTRIES FOR COMPOUND		232.4 UNK 3.8	POTASSIUM 4-HEXYL RESORCINOLATE X10-2 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 705 MOL WGT – 1 ENTRIES FOR COMPOUND		266.4 UNK 6.2	DODECYL SULFURIC ACID X10-3 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 706 MOL WGT – 1 ENTRIES FOR COMPOUND		185.1 UNK 1.32	PERFLUORO PROPYLAMINE X10-1 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 707 MOL WGT – 1 ENTRIES FOR COMPOUND		221.5 UNK 1.1	PERFLUORO PROPYLAMINE HYDROCHLORIDE X10 0 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 708 MOL WGT – 1 ENTRIES FOR COMPOUND		101.2 UNK 0.4	HEXYLAMINE X10-2 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 709 MOL WGT – 1 ENTRIES FOR COMPOUND		137.7 UNK 9.	HEXYLAMINE HYDROCHLORIDE X10-1 M XG	METHOD NOT CITED	KLEV RAIS	54004	T L
COMPOUND NO = 710 MOL WGT – 3 ENTRIES FOR COMPOUND		190.3 30 7.7 4.04 30 7.9 4.15 30 8. 4.2 X10-1 D X10-2 M X10-1 D X10-2 M X10-1 P X10-2 S	OCTYL DIMETHYL PHOSPHINE OXIDE BB TURBIDITY PLT LITE SCATR SURFACE TENSION LOG PLOT BD DENSITY	HERR BRUS HERR BRUS BENJ	66039 66039 66040	T 3 M T L M T L M	
COMPOUND NO = 711 MOL WGT – 3 ENTRIES FOR COMPOUND		218.4 30 1.0 4.57 30 8.2 3.75 30 7. 3.2 X10-1 D X10-3 M X10-2 D X10-3 M X10-2 P X10-3 S	DECYL DIMETHYL PHOSPHINE OXIDE BD TURBIDITY PLT LITE SCATR SURFACE TENSION LOG PLOT BD DENSITY	HERR BRUS HERR BRUS BENJ	66039 66039 66040	T L M T L M T L M	
COMPOUND NO = 712 MOL WGT – 3 ENTRIES FOR COMPOUND		246.4 1 2.0 8.11 30 1.4 5.68 X10-2 D X10-4 M X10-2 D X10-4 M	DODECYL DIMETHYL PHOSPHINE OXIDE BC TURBIDITY PLT LITE SCATR BD TURBIDITY PLT LITE SCATR	HERR BRUS HERR BRUS	66039 66039	T 3 M T 3 M	

Concentration units: A – mol %; B – vol % solvent; C – mol % surfactant mixture; D – wt/vol %; E – % saturation; H – wt % solvent; I – mol % surfactant; K – normality counterions; M – molar; N – normal; P – wt %; Q – wt % surfactant; R – varied; S – mol/kg; T – wt % surfactant mixture; U – mol/l or kg; W – molal; Y – atm. Details on page 222.

Complete Table of Critical Micelle Concentration Values—Continued

Additives	Temp. °C	CMC	Qual. Mat. Meth.	Method	Authors	Reference	Source Evaluation	
	30	8. 3.2	X10-3 D X10-4 M	BD	SURFACE TENSION LOG PLOT	HERR BRUS	66039	T L M
3 ENTRIES FOR COMPOUND								
COMPOUND NO = 713 MOL WGT —	215.4		UNDECYL DIMETHYL AMINE OXIDE					
	30	1.3 6.03	X10-1 P X10-3 S	BD	DENSITY	BENJ	66040	T L M
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 714 MOL WGT —	145.3		HEXYL DIMETHYL AMINE OXIDE					
	UNK	3.0 2.06	X10 1 P X10 0 S	BE	METHOD NOT CITED	BENJ	66040	T L M
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 715 MOL WGT —	159.3		HEPTYL DIMETHYL AMINE OXIDE					
	UNK	1.0 6.27	X10 1 P X10 1 S	BE	METHOD NOT CITED	BENJ	66040	T L M
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 716 MOL WGT —	891.3		DODECYL /OXYETHYLENE/16 ALCOHOL REDUCED POLYDISPERSITY OF HEAD GROUPS					
	25	2.205X10-2 2.473X10-4	D M	EE	FOTOMTR SOLUBLZTN OROT	SCHO	66036	T L M
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 717 MOL WGT —	378.4		DODECYLQUINOLINIUM BROMIDE					
	25	4.80	X10-3 M	BB	SURFACE TENSION LOG PLOT	FEW GILB	58031	T 3
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 718 MOL WGT —	395.7		TETRAETHYLAMMONIUM DODECYL SULFATE					
	30	4.5 4.1	X10-3 M X10-3 M	CB CG	EQUIV CONDCTNCE GRAPH FOTOMTR SPCTR CHNGE RHD6	MEGU KOND MEGU KOND	59026	T L T L
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 719 MOL WGT —	507.9		TETRABUTYLAMMONIUM DODECYL SULFATE					
	30	1.3 1.0	X10-3 M X10-3 M	CC CG	EQUIV CONDCTNCE GRAPH FOTOMTR SPECTR CHNGE RHD6	MEGU KOND MEGU KOND	59026	T L T L
2 ENTRIES FOR COMPOUND								
COMPOUND NO = 720 MOL WGT —	733.2		1-6-DITRIMETHYLAMMONIUM-HEXANE/DODECYL SULFATE/2					
	30	9.6	X10-4 M	CB	EQUIV CONDCTNCE GRAPH	MEGU KOND	59026	T 3
1 ENTRIES FOR COMPOUND								
COMPOUND NO = 721 MOL WGT —	1,542.1		NONYL BENZENE /OXYETHYLENE/30 ALCOHOL BRANCHED CHAIN REDUCED OE DISTRIBUTION					
	25	4.22 2.736X10-4	X10-2 D M	EC	ULTRAFILTRATION	SCHO	64004	T L M
1 ENTRIES FOR COMPOUND								

Concentration units: A—mol %; B—vol % solvent; C—mol % surfactant mixture;
D—wt/vol %; E—% saturation; H—wt % solvent; I—mol % surfactant; K—normality

counterions; M—molar; N—normal; P—wt %; Q—wt % surfactant; R—varied; S—mol/
kg; T—wt % surfactant mixture; U—mol/(l or kg); W—molal; Y—atm. Details on page 222.

Index of Additives Giving Compound Numbers

(CH ₃) ₂ N BR		DIOXANE	
99		1	4
(CH ₃) ₂ N CL		5	38
116 321 324		95	166
(CH ₃) ₂ N I		320	322
1		323	
(C ₂ H ₅) ₂ N BR		ETHANOL	
99		1	21
(C ₃ H ₇) ₂ N BR /NORMAL		38	44
99		90	91
(C ₄ H ₉) ₂ N I		92	95
1		166	263
(C ₄ H ₉) ₂ N BR /NORMAL		ETHYLENE GLYCOL	
99		38	95
ACETIC ACID		FORMIC ACID	
38 482		38	
ACETONE		GLYCEROL	
38 667		38	245
ACETONITRILE		GUANIDINIUM CL	
38		21	320
AG NO ₃		322	323
1 23		GUANIDINIUM CO ₃	
AL CL ₃		21	
1		H BR	
BA CL ₂		99 293	
38 95 321		H CL	
BENZENE		1	22
38 40 44 90 91 92 179 182 183		38	168
BROMPHENOL BLUE		245	320
99		321	322
BUTANOL-1		323	
21 38 44 90 91 92 95		417	482
BZL*1C ₆ H ₅ /CH ₃ / N I		H CL ₀₄	
CA ACETATE		206	528
38		H NO ₃	
CA BR ₂		38	97
97		99 168	206
CA CL ₂		HEPTANOL-1	
1 18 24 132 163 164 165 166 205 206 321		38 44	
CA FORMATE		90 91	92
38		HEXANOL-1	
CA NO ₃		1 3 38 44	
321		90 91 92 273	
CALGON (NA HXMTP*)		HEXYL AMMONIUM CL	
273		38	
CAPRYLAMIDE		H ₂ SO ₄	
171		491 529 530 531 532 533 534	
CARBOXYMETHYLCELLULOSE		H ₃ PO ₄	
273		460	
CL- ION		IONIC STRENGTH	
38		1 206	
CO SO ₄		ISOC ₅ GLYCEROL ETHER	
572		171	
CS CL		K BR	
1		91 93 94 95 96 97 98 99 100 101 102 290	
CS ₂ SO ₄		427	
1		K CL	
CU SO ₄		1 38 41 44 90 91 92 97 99 206 278 296	
1 572 573		297 321 350 351 353 399 417 550 551 552 553	
CYCLOHEXANE		K CNS	
38		206 278	
C ₁₂ CLORHYDRIN GLET*		K H SO ₄	
171		531 534	
C ₁₂ DIETHANOLAMIDE		K I	
139		91 290 376 550 551 552 553	
C ₁₂ ETHANOL AMIDE		K I ₀₃	
171 172 173 174		290	
C ₁₂ GLYCEROL ETHER		K NO ₃	
171		40 91 97 99 206 321 456 550 551 552 553	
C ₁₂ SULFOLANYLAMIDE		K OH	
171		1 44 90 91 92 97 99 188 206 296 297 417	
C ₆ H ₅ (CH ₃) ₂ N OH		K+ ION	
277		91 185 668 670 671 672	
C ₈ GLYCEROL ETHER		K ₂ SO ₄	
171 172		1 90 91 290	
DC ANTIFOAM A (PMS*)		K ₄ P207 PYRO	
451		91	
DECANOL-1		LA BR ₃	
38 44 90 91 92 171 201 202 204 263 273 451		97	
DEUTERIUM OXIDE		LA CL ₃	
1 3		38	
		LAURYL ALCOHOL	
		1 38 263 451	
		LAZ (SO ₄) ₃	
		308	
		LI BR	
		290	
		LI CL	
		1 116 321 324	
		LI NO ₃	
		321	
		LI ₂ SO ₄	
		1	
		METHANOL	
		21 38 95 270 427 654 656 657	
		661 667	
		MG (NO ₃) ₂	
		321	
		MG CL ₂	
		1 321	

Index of Additives Giving Compound Numbers—Continued

MG S04	NH4 BR
308 568	99
N-C10 GLYCEROL ETHER	NH4 CL
1 171 172 175 176 177 178 179	97 99 206
N-C10 SULFOLANYL ETH	NI S04
171	575
N-DECANE	NITROBENZENE
201 202 204	274
N-HEPTANE	NONANOL-1
1	44 90 91 92
N-HEXANE	OCTANOL-1
38 91	1 44 90 91 92 273
N-3SOA*	OLEIC ACID
171	263
NA ACETATE	PENTADECANOL V. BR*
38 482	1
NA BR	PENTAMINE
1 41 97 98 102 127 128 129	1
130 131 168 278 290 291 293 321	PENTANOL-1
346 347 643	1 245
NA BRO3	PH OF SOLUTION
129 168 321	1 21 38 91 92 99 185 206
NA CITRATE	263 273 298 299 320 322 323 416
163 164 165 166 321	417 485 487 529 703
NA CL	PHENOL
1 2 3 4 5 18 21 22	97
24 38 41 42 91 93 95 97	PHENYL (CH3)3 N I
111 116 132 133 134 135 136 137	1
139 156 163 164 165 166 167 168	PINACYANOL CL (DYE)
169 179 182 205 206 258 273 274	1 44 91 188 296
278 290 295 311 321 324 329 337	PRESSURE
345 418 419 432 433 434 459 462	1 38 41 95 97 279
482 492 588 634	PROPANOL-1
NA CLO4	1 21 38 44 90 91 92 95
3	PROPANOL-2
NA CNS	1 38
116 321 324	PROPIONIC ACID
NA F	1
1 130 321	RB BR
NA HC02 FORMATE	290
38 128	RHODAMINE 6GPC
NA HC03	1 111
273	SIO2/NA20=1.60
NA I	91 273
1 168 278 321	SIO2/NA20=2.46
NA IO3	273
127 168 321	SIO2/NA20=3.93
NA NO3	273
1 23 24 38 131 273 278	SR CL2
NA OH	321
132 139 168 263 273 298 300 321	SUCCINIC ACID
448	38
NA P-TOLUENE-S03	SUCROSE
295	1 166
NA PO4	TARTARIC ACID
139 273	38
NA SUCCINATE	TERTIARY BUTANOL
38	38 660
NA+ ION	TETRADECANOL-2
1	171
NA14 P12037 POLY	TMCHCGLET*
139	171
NA2 B407	TRIBUTYL PHOSPHATE
273	1 263 451
NA2 C03	TRIETHYL CARBINOL
139 273 680	38
NA2 S103 META	UNDECANOL-1
91 139 273 680	38
NA2 S04	UREA
1 18 91 116 132 163 164 165	1 3 5 21 97 111 112 113
166 171 182 273 278 307 308 309	114 115 116 117 320 321 322 323
321 324 573 680	376
NA2 S203 THIOSULF	1,10 DECANE DIOL
376	273
NA4 P207 PYRO	1,2 DECANE DIOL
1 91 139 149 273 278	273
NA5 P3010 TRIPOLY	2-ETHYL HEXANOL
1 139 141 142	451
NA53 P500154 POLY	3-METHYL BUTANOL-1
139	44 90 91 92
NH3	3,5(CH3)2 C6H3 GLET*
97 99 206	1

Index to References by Additives

(CH3) 4 N BR	C6H5 (CH3) 3 N OH
65012	48009
(CH3) 4 N CL	C8 GLYCEROL ETHER
62019 64020	57014
(CH3) 4 N I	DC ANTIFOAM A (PMS*)
53012	57031
(C2H5) 4 N BR	DECANOL-1
65012	50008 55004 56018 57014 57031 60006 62009
(C3H7) 4 N BR /NORMAL	DEUTERIUM OXIDE
65012	66002
(C4H9) CH3 3 N I	DIOXANE
53012	46004 49006 65011 65013 65020
(C4H9) 4 N BR /NORMAL	ETHANOL
65012	40004 46001 48023 49006 50008 55004 57013
ACETIC ACID	58007 65013 66012
49008	ETHYLENE GLYCOL
ACETONE	46004 49006
48023 49017	FORMIC ACID
ACETONITRILE	49008
48023	GLYCEROL
AG NO3	36002 46004
62006	GUANIDINIUM CL
AL CL3	65011 66012
51005	GUANIDINIUM CO3
BA CL2	66012
47010 49008 62019	H BR
BENZENE	64006 65012
48021 49006 50003 55008	H CL
BROMPHENOL BLUE	36002 49008 54010 56014 57021 62005 62019
51008	64016 65011
BUTANOL-1	H CL04
49006 50008 55004 66012	50012 65019
BZL* C6H5 /CH3/2 N I	H NO3
53012	49008 50012 56014
CA ACETATE	HEPTANOL-1
49008	50008 55004 56018
CA BR2	HEXANOL-1
64043	50008 55004 56018 59018
CA CL2	HEXYL AMMONIUM CL
51005 61008 62002 62011 62019 63008	48020
CA FORMATE	H2 SO4
49008	65019 65025
CA NO3	IONIC STRENGTH
62019	65030
CALGON (NA HXMT*)	ISOC5 GLYCEROL ETHER
48024	57014
CAPRYLAMIDE	K BR
57014	48005 49001 55009 59010 59024 64043 66028
CARBOXYMETHYLCELLULO	K CL
48024	47010 48005 49005 50008 50012 51005 52017
CL- ION	53002 54010 56001 56002 62019 62037 65026
54013	66023 66028
CO S04	K CNS
63034	50012 51005
CS CL	K H SO4
53012	65025
CS2 S04	K I
53012	48005 59024 60015 66006 66023 66028
CU S04	K IO3
63034	59024
CYCLOHEXANE	K NO3
48021	48005 48025 49005 50012 61026 62019 62037
C12 CLORHYDRIN GLET*	64011 66023
57014	K OH
C12 DIETHANOLAMIDE	48004 48009 48011 48012 49005 50012 51004
58008	53006 54003 54005 54010 55004 56001 59008
C12 ETHANOL AMIDE	59012 60002 61025 65012
57014	K+ION
C12 GLYCEROL ETHER	54013 55007
57014	K2 S04
C12 SULFOLANYLAMIDE	47010 48005 49005 53012 59024 62037
57014	

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K4 P207 PYRO	NA P04
48005	48024 59009 62038
LA BR3	NA SUCCINATE
64043	49008
LA CL3	NA14 P12037 POLY
47010	59009
LAURYL ALCOHOL	NA2 B407
49014 57014 57031 58023	48024 62038
LA2(SO4)3	NA2 C03
64043	48024 59009 60033 62038
LI BR	NA2 S103 META
66028	48024 59009 60033
LI CL	NA2 S04
56002 62019 64020	47010 51005 53012 57014 60033 61008 61030
LI N03	62002 62019 62037 63008 63026 63034 64020
62019	64043 65019
LI2 S04	NA2 S203 TH10SULF
53012	63032
METHANOL	NA4 P207 PYRO
48010 48023 48028 49006 49008 49014 49017	47010 48024 51005 59009
66012	NA5 P3010 TRIPOLY
MG (N03)2	58008 59009 61015
62019	NA53 P500154 POLY
MG CL2	59009
51005 62019	NH3
MG S04	50012
63034 64043	NH4 BR
N-C10 GLYCEROL ETHER	65012
57014	NH4 CL
N-C10 SULFOLANYL ETH	50012
57014	NI S04
N-DECANE	63034
60006 62009	NITROBENZENE
N-HEPTANE	49018
56001	NONANOL-1
N-HEXANE	55004
48011 48021	OCTANOL-1
N-3SOA*	55004 56001 56018
57014	OLEIC ACID
NA ACETATE	53004
49008	PENTADECANOL V.BR*
NA BR	57014
51005 56014 62019 63014 63016 63026 64001	PENTAMINE
64006 64017 64047 65005 66014 66025	56001
NA BR03	PENTANOL-1
56014 62019 63016	36002 56001
NA CITRATE	PH OF SOLUTION
62002 62019	51008 54004 54013 56001 56018 60025 64051
NA CL	65011 65030 66012 66027
38001 39007 47010 48024 49005 49008 51005	PHENOL
53005 55003 55005 55021 56002 56011 56014	62035
56020 57006 58009 58020 59001 59009 59024	PHENYL (CH3)3 N I
60005 60011 60025 60027 60028 61005 61008	53012
61030 62002 62004 62005 62009 62010 62011	PINACYANOL CL (DYE)
62019 62020 63008 63026 64020 64047 65018	46010 52015 52017 55015
66010 66013 66027 66036	PRESSURE
NA CL04	62035 62036 65036
62040	PROPANOL-1
NA CNS	49006 50008 53008 55004 66012
62019 63026 64020	PROPANOL-2
NA F	49014 50008 53008
62019 63016 64047	PROPIONIC ACID
NA HC02 FORMATE	53008
49008 63016	RB BR
NA I	66028
51005 56014 62019 64047	RHODAMINE 6GPC
NA IO3	56020
63016	SI02/NA20 = 1.60
NA N03	48024
49008 51005 56014 61026 61030 62006 62019	SI02/NA20 = 2.46
63016	48024
NA OH	SI02/NA20 = 3.93
38006 48011 48024 51004 53004 56014 59009	48024
62019 62038 63008	SR CL2
NA P-TOLUENE-S03	62019
66013	

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SUCCINIC ACID	UNDECANOL-1
49008	50008
SUCROSE	UREA
63020 64034	47010 61016 62020 63032 64020 65011 66012
TARTARIC ACID	47010 48011 53010
49008	1,10 DECANE DIOL
TERTIARY BUTANOL	56018
49014	1,2 DECANE DIOL
TETRADECANOL-2	56018
57014	2-ETHYL HEXANOL
TMCHCGLET*	57031
57014	3-METHYL BUTANOL-1
TRIBUTYL PHOSPHATE	55004
57031	3,5 (CH ₃) 2 C ₆ H ₃ GLET*
TRIETHYL CARBINOL	57014
50008	

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27001	273	298					48023	38					
27002	43						48024	91	273	300			
29001	43						48025	91					
30001	44						48027	38					
32001	298						48028	427	654	655	656	657	
34001	274						49001	95	97	98			
35001	408						49004	271	272				
35008	91						49005	38	90	91	92	258	273
36001	99	427						305					
36002	245	274					49006	91	95	182			
38001	274						49008	38	482				
38006	273	298	300	448			49009	500					
39002	179	182	183	184	349		49013	38	41	449	450		
39006	273	298	300	448			49014	38	660	661			
39007	179	182	183				49017	667					
39009	1	263	273	299	476	484	49018	274					
	485	486	487				50003	44	90	91	92	179	182
39011	200	243	536	537				183					
40003	273	298	299	300			50008	38					
40004	1						50012	97	99	206			
41003	299						51001	99					
41004	273						51003	181	339	340	341	342	343
42002	37	38	39	186	187			344					
42003	1	2	3	260	261	262	51004	1	38	90	91	92	273
	352							278	554	556	557	558	559
42004	273	300						560	561	562	563		
43001	49	51					51005	1	24	278			
43003	93	95	97	99			51006	38					
43004	287	288					51008	99					
43006	286	304					51009	662	663	664	665	666	
46001	38						51010	376	458				
46002	91	92					52001	41	42	265	270	279	399
46004	38	95						400	401	402	403	404	405
46005	1	38	91	95	182			406	407				
46006	92	273	299				52011	18	100				
46007	1	44	90	91	92		52013	278	554	555			
46008	285						52015	44	90	91	92	188	296
46010	5	91	92	305	494			297	350	351			
46012	38	91					52016	1	91	111	273	634	635
46015	1	90	91	92				636	637	638	639		
46016	283						52017	44	90	296	297	350	351
46019	284	303					52018	681	682	683			
47003	265	266	267	268	269	270	53001	263	264				
47004	1	38	91	92	179	181	53002	287	288	353			
	182	183	184				53003	91	92	355	372	373	374
47005	91	92	179	182				375					
47006	1	38	91	95	182		53004	263					
47007	206	276					53005	1	4	5	64	99	290
47010	1	38	91	95	182			427					
47011	259						53006	1	38	90	259	273	284
47013	92	273	299					299	476	487			
48004	91	200					53007	275	279	354	356	357	358
48005	1	37	38	39	44	90		359	360	361	362	363	364
	91	92	93	95	97	99		365	366	367	368	369	370
	179	181	182	183	184	185		371					
48007	200						53008	1	4				
48008	4	16	17	72	73	77	53010	37	38	39	41	42	91
	84	525	526	527				95	185	186	203	255	256
48009	42	91	277	278	279			273	278	290	305	392	627
48010	270	654	655	658	659			628	629	630	631	632	633
48011	91	92	99	263	273	274	53012	1					
	275						53015	139	279				
48012	255	256					54003	44	90	91	92	188	296
48014	483							297					
48016	38	90	91	92	258	273	54004	1	43	44	179	188	200
	299	305						416	417	455	456	529	699
48018	263							700	701	702	703	704	705
48020	38	39						706	707	708	709		
48021	38						54005	44	90	91	92	188	296
48022	1	200						297					

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54006	38	263	273	298	300	448	58017	578	579
	564	565	566	567			58020	1	
54008	651	652	653				58021	409	410
54010	1	416	417	428	430	455	58023	1	
	456						58028	586	
54013	1	38	91	185			58031	717	
55003	1						59001	1	492
55004	44	90	91	92			59002	111	493
55005	1						59004	1	5
55006	97	307						52	53
55007	668	669	670	671	672	673		56	57
55008	40							62	63
55009	93	94	95	96	97	98	59005	166	
	99	100	101	102			59006	163	164
55015	1							433	435
55018	1							439	440
55021	1	3	4	179	181	182	59007	43	
	183	340	341	342	345		59008	44	
55028	492	501	502	503	504	505	59009	139	149
	506	507	508	509	510	511	59010	99	273
	512	513	514				59012	44	
56001	1	38	49	50	91	92	59013	18	423
	259	416	417				59015	1	424
56002	1						59016	1	273
56003	420	421	422	425	426		59017	409	
56005	171	301	302				59018	1	3
56006	1	2	4	5	15	16	59020	166	
	17	64	66	68	69	70	59023	430	
	71	72	73	74	75	76	59024	278	290
	77	78	79	80	81	82	59026	112	376
	83	84	85	86	87	88	60001	37	718
	89						60002	38	719
56008	36	189	190	191	192	193	60004	1	720
	194	195	196	197	198	199	60005	38	
56010	416	417	428	429	430		60006	201	202
56011	1	2	3	4	5	295	60008	6	
								7	9
56014	156	168	169	335	336	337	60010	1	10
	338							138	141
56016	51							144	142
56018	273							145	148
56019	102	274	376	427	479	480		149	151
	481							155	152
56020	1	111	634					156	153
57004	37	38	39					157	154
57005	1	40							160
57006	38	41	42						
57009	257	258	259	260	261	262	60018	1	57
57010	451							4	55
57011	1	2	3					5	54
57012	45	46	47	48	49	50	60020	1	541
	51						60021	1	542
57013	1						60024	1	543
57014	1	67	171	172	173	174	60025	1	544
	175	176	177	178	179		60026	91	545
57016	186						60027	295	546
57017	416	417	428	429	430		60028	446	547
57020	578	579	580	581	582	583	60029	1	
	584	585						24	568
57021	38							572	569
57022	1	2	3	4	5	64	60032	91	570
57024	495						60033	273	571
57025	1							674	
57031	1	263	451				60034	91	
58001	257	258	259	260	262		61001	280	
58003	5	52	53	54	55	56	61002	41	
	57	58	59				61003	164	166
58004	263							165	167
58007	263	264					61004	105	168
58008	1	139	263				61005	3	
58009	274						61007	1	15
58011	44	90	91	92	188	296	61008	18	16
	297	350	351	374	375	416	61014	1	17
	417	428	429	430	452	453		99	
	454	455					61015	1	145
58012	1	97						98	149
								139	
								141	
								142	
									149

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61016	97						64004	170	721					
61017	1	3					64006	21	22	293				
61025	90						64007	280						
61026	40	1					64009	108	110	282	331	377	378	
61027	99	451					64010	139	206					
61029	693	694	695	696	697	698	64011	456						
61030	273						64012	179						
61031	1	2	3	4	5	64	64014	207	208	209	210	211	212	
	273	298	299	300	448	501		213	214	215	216			
	502	503	505	506	507	508		64016	21	22	251	252	254	
	509							64017	1	97				
62001	166	168					64020	1	3	5	111	112	113	
62002	163	164	165	166	167	168		114	115	116	117			
	169	432	433	434	435	436		64023	105	108	282	289		
62004	2	3	93	95				64024	468	469	470	471	472	473
62005	21	22	251	252	253				474	475				
62006	1	23	24					64025	1					
62008	7	25	26	27	28	29		64027	294					
	30	31	32	33	34	35		64032	1					
	36							64034	1					
62009	1	204						64035	411	412	413	414	415	
62010	1	205						64037	645	646	647	648	650	651
62011	205	206						64043	95	97	306	307	308	309
62015	103	294	380	381				64047	1	2	3	295	311	
62019	115	116	312	313	314	315		64049	294	393	394	395	396	397
	316	317	318	319	320	321			398					
	322	323	324	325	326	327		64050	290					
	328	329						64051	1	5	263	273	298	299
62020	115	116	320	321	322	323			485	487				
	324	326	327	328	329			65003	119	120	121	122	123	
62023	495	496	497	498	499			65005	346	347	348			
62027	282	330	331	332	333	334		65007	179					
62035	95	97						65011	320		322	323		
62036	1							65012	99					
62037	91							65013	166					
62038	273							65018	1	3				
62040	3							65019	460	491	528			
63001	1	3	4					65020	1	4	5			
63008	132	133	134	135	136	137		65022	1	2	4			
63009	118	132						65024	273	298	299	476		
63010	132	133	134					65025	529	530	531	532	533	534
63012	133	134						65026	41	278	399			
63013	1	4	5	9	10	64		65028	1	91	99	596		
	179	183	184	200	229	230		65030	1	206				
	231	232	233	234	235	236		65031	38					
	237	238	239	240	241	242		65036	38	41	279			
	243	244	245	246	247	248		65037	427					
	249	250						66001	1	115	116	325	327	
63014	346							66002	1	3				
63015	118	119	120	121	122	123		66003	597	598	599	600	601	
	124	125	126					66006	290	376				
63016	41	97	127	128	129	130		66007	1					
	131							66010	1					
63017	207	208	209	210	211	212		66011	51	274	275	358	477	478
	213	214	215	216	217	218		66012	21					
	219	220	221	222	223	224		66013	587	588	589	590	591	592
63020	166								593	594	595			
63021	108	110	377	378	380	381		66014	287	288	346	347	385	640
63026	1	97	113	114	115	116			641	642	643	644		
	117							66015	684	685	686	687	688	689
63030	112	280	382	383	384	385			690	691	692			
	386	387	388	389	390	391		66018	1	97	98	99	459	
63032	376							66019	461					
63034	1	568	572	573	575	576		66020	105					
	577							66021	139	463	464	465	466	467
63037	457							66022	310	336	515	516	517	518
64001	98	102	290	291	292				519	520	521	522	523	524
64002	26	189	190	191	194	602		66023	550	551	552	553		
	603	604	605	606	607	608		66025	97	99	115	116	117	325
	609	610	611	612	613	614		66027	327	535				
	615	616	617	618	619	620			21					
64003	621	622	623	624	625	626								
	103	104	105	106	107	108								
	109	110												

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66028	278	290	376		66038	1	4	5	102	290	427
66030	99				66039	710	711	712			
66036	1	170	182	716	66040	21	251	252	254	587	588
66037	331					710	711	713	714	715	

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AVER COND BEGINING MAXIM		GRAPH DATA NOT RETRIEVED
53008		46007 53005 54004 54005 61027 61030 62037
AVER SP EQUIV COND		62038 63034
53008 58007 61007		HEAT OF DILUTION
DEBYE PLT LIGHT SCATTER		64016 66012
49001 53002 55005 63016 64007		INTERFACIAL TENSION LOGMI
DENSITY		38006 57022 58012 60011 60025 62004 62023
35008 47011 60005 63009 66020 66040		INTERFACIAL TNSN UNSPEC
DIFFUSION COEFFICIENT		56019
61025		KRAFFT POINT SOLUBILITY
ELECTROMOTIVE FORCE		39002 49004 56019 57012 61029
47013 48007 48025 57021 59016 60021 63001		METHOD NOT CITED
64012 64032		49006 51003 53003 54004 56001 61014 62037
EMF ALONG CONC GRADIENT		63020 63034 64035 64043 66013 66014 66040
59015		MICELLAR SPECTRAL CHANGE
EQUIV COND MAX BEGINING		51010 63032 66006
49014 49018 51009 58003		PARTIAL VOLUME
EQUIV COND 1ST DEVIATION		29001 30001
36002 40004 43001 48010 49014 49017 53005		PH OR HYDROLYSIS
55018		27001 39006 41003 60001
EQUIV CONDCTNCE GRAPH		POLAROGRAPHIC MAXIMUM
27001 32001 34001 35001 36001 39011 40004		50012 66023
41003 42002 42004 43001 43003 47003 48009		QUESTIONABLE CRITERION
48010 48012 48014 48020 48021 48023 48028		39006 39009 41004 42003 51001 52013 53001
49004 49008 49009 49013 49014 49017 49018		53015 54010 57010 57017 58011 60008 60017
51009 53004 53005 53008 55005 55006 55018		61029 62002 62023 63013 63015 63017 64043
56006 56008 57013 57017 57025 58001 58003		65012
58007 59002 59026 61005 61016 61017 62040		REACTN RATE SULUBILIZATE
64034 65020		65030
EQUIV CONDCTNCE MAXIMUM		REFRACTIVE INDEX
49014 56006		46012 47004 47005 48005 53010 56011 57005
FLOCCULATION RATE		63015 63021 63037 64047 65003 65012 66019
66037		SEE CMPD NMBR IN ADDITV
FOAMING POWER		46015 48005 48020 51005 53003 53005 54003
59017		54005 54010 56014 58017 61002 61007 61008
FOTOMTR SOLUBLZTN AZBZ		61017 62010 63001 63017 64010 66001 66025
38001 49005		SOLUBLZTN TOLUENE
FOTOMTR SOLUBLZTN DMYL		60032 60034
60027		SPECFC CONDCTNCE EQUATNS
FOTOMTR SOLUBLZTN OROT		57011 59012 66002
48016 48024 49005 51010 55005 59009 60010		SPECFC CONDCTNCE GRAPH
61015 64010 66036		36001 36002 39007 39011 42003 43003 43004
FOTOMTR SOLUBLZTN PDMAB		46001 46010 48004 48027 51003 51006 53004
46006 47013 48016 49005 59009		54006 55005 55008 56005 56006 56020 57004
FOTOMTR SOLUBLZTN SDN 4		57012 57016 57025 57031 58001 58004 58007
57020 58028 63008 63010 63037 64024		58008 59002 59024 60020 60028 61005 61017
FOTOMTR SOLUBLZTN YLOB		62006 62035 62036 63030 64050 64051 65018
64024 66027		65024 65031 65036 65037 66007 66030 66038
FOTOMTR SOLUBLZTN 2NPA		SPECIFIC HEAT
55007		60002
FOTOMTR SPCTR CHNG BZP4		STREAMING CURRENT
62001 65013		66011
FOTOMTR SPCTR CHNG ERTS		SUMMARIZING EQN ONLY
57020		62002
FOTOMTR SPCTR CHNG I2		SURFACE TENSION LOG PLOT
59006 59020 60012 60026 60027 62001 62002		38006 48022 52011 54010 54013 56014 56016
63015 64009 66023		56018 57009 57012 57022 57024 58003 58031
FOTOMTR SPCTR CHNGE BRPB		59001 59013 59020 60004 60012 60027 60029
51008 63015		61004 61008 61026 62015 62019 62020 62023
FOTOMTR SPCTR CHNGE EOSN		62027 63008 63009 63012 63013 63014 63015
59024		63017 63026 63034 63037 64001 64003 64009
FOTOMTR SPCTR CHNGE INPX		64010 64014 64020 64023 64024 64037 64047
47004		64049 65003 65005 65007 65011 65019 65024
FOTOMTR SPCTR CHNGE PNCN		65025 65026 66001 66007 66013 66015 66019
46002 46010 47004 50003 52015 53012 55007		66021 66022 66025 66039
55015 56008 56018 57020 57025 58007 58008		SURFACE TENSION MINIMUM
58017 58028 60033		50012 54010 57012 63017 65024
FOTOMTR SPCTR CHNGE RHD6		SURFACE TENSION UNSPEC
56020 59026 65030		54004 55007 56008 58008 60018 60026 62001
FREEZING POINT		62002 64002 66018 66028
27002 39011 43006 46008 46016 46019 47007		SURFACE TNSN LINEAR PLOT
48011 59023		50012 57025 58020 59017 61001 66007

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THEORETICALLY ESTIMATED	VELOCITY OF SOUND
55007 56003 56011	65022 65028 66010
TURBIDITY PLT LITE SCATR	VIISCOSITY
52018 53002 55003 55005 55009 56011 57006	39009 47011 57025 64025
58009 59005 59006 59010 60006 60015 61003	VIISCOSITY MINIMUM
62002 62005 62009 62010 62011 63009 63012	53006 54008
63016 64006 64017 64047 65025 66013 66028	VISUAL FLUOR CHNGE RHD6
66039	56002
TURBIDMTR SOLUBLZTN LOH	VISUAL SPCTR CHNGE
58023	VISUAL SPCTR CHNGE DCFL
ULTRACENTRIFUGATION	46004
64017	VISUAL SPCTR CHNGE EOSN
ULTRAFILTRATION	47006 51005
27001 59018 64004	VISUAL SPCTR CHNGE FL
UNSPEC LIGHT SCATTER	47006
61004	VISUAL SPCTR CHNGE INPX
UNSPEC SOLUBLZTN PDMAB	47006 47010 50008 53007
48025 59007	VISUAL SPCTR CHNGE PNCN
UNSPEC SOLUBLZTN SDN 4	46015 47006 47010 48008 48024 50003 51003
61001 61002	51005 52015 52016 52017 54003 54005 55004
UNSPEC SPCTR CHNG PNCN	55007 56003 56008 56016 57014 58003 58008
49004 58011	59004 60008 60029 60033 60034 61030 61031
UNSPECIFIED CONDUCTANCE	62008 63013 64002 66003
42003 48008 52001 53010 55021 56002 57014	VISUAL SPCTR CHNGE RHD6
58023 59015 59016 60033 61002 62040 63034	47006 48011 55028 56016 60021 60028 60029
64011	60033
VALUES FRM REF IN CMC	VISUAL SPCTR CHNGE SKYB
40003 46004 46005 48005 49005 56010 57022	47006 47010 49006 51010
58021 59006 59013 60024 61017 62020 63026	WIEN EFFECT
64006 65005 65011 66028	48018
VAPR PRESURE LOWERING	X-RAY DIFFRACTION
48011 51004 59008 64027 66015	39009

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	TRANS FARADAY SOC	60	996	1964	64027
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GREENFIELD A	J PHYS CHEM	61	818	1957	57004
GREGORY NW	J AM CHEM SOC	70	1992	1948	48012
GRIEGER PF	J AM CHEM SOC	68	1137	1946	46018
	J AM CHEM SOC	69	1835	1947	47001
	J AM CHEM SOC	70	3803	1948	48010
	J AM CHEM SOC	71	95	1949	49014
	J AM CHEM SOC	71	309	1949	49017
	J AM CHEM SOC	71	1455	1949	49018
GRIESS W	FETTE, SEIFEN, ANSTRICHMI	57	236	1955	55026
	FETTE, SEIFEN, ANSTRICHMI	57	168	1955	55027
	FETTE, SEIFEN, ANSTRICHMI	57	24	1955	55028
GRINDLEY J	J CHEM SOC		679	1929	29001
GUENTHNER RA	J PHYS CHEM	57	923	1953	53014
HAFFNER FD	J PHYS CHEM	46	662	1942	42003
HALL NA	J PHARM SCI	54	1529	1965	65027
HAMANN SD	J PHYS CHEM	66	1359	1962	62036
HARKINS WD	J AM CHEM SOC	59	2197	1937	37005
	J AM CHEM SOC	62	1496	1940	40005
	J CHEM PHYS	14	216	1946	46002
	J CHEM PHYS	14	640	1946	46004
	J CHEM PHYS	14	641	1946	46005
	J CHEM PHYS	14	480	1946	46010
	J CHEM PHYS	14	215	1946	46011
	J AM CHEM SOC	68	220	1946	46013
	J COLLOID SCI	1	105	1946	46014
	J COLLOID SCI	1	469	1946	46015
	J CHEM PHYS	14	214	1946	46017
	J AM CHEM SOC	69	679	1947	47006
	J CHEM PHYS	15	763	1947	47008

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HARRIS JC	J AM CHEM SOC	69	683	1947	47010
	J CHEM PHYS	15	496	1947	47012
	J PHYS COLLOID CHEM	53	1350	1949	49006
	J COLLOID SCI	4	367	1949	49007
	J AM CHEM SOC	71	808	1949	49011
	J PHYS COLLOID CHEM	54	271	1950	50008
	SCI MONTHLY	70	220	1950	50011
	J COLLOID SCI	6	576	1951	51010
	J AM CHEM SOC	74	2061	1952	52001
	SOAP CHEM SPECIALTIES	1958		1958	58002
	J AM OIL CHEMISTS SOC	35	670	1958	58004
	J PHYS CHEM	62	1554	1958	58008
	J AM OIL CHEMISTS SOC	35	428	1958	58010
	J AM OIL CHEMISTS SOC	36	332	1959	59009
	J AM OIL CHEMISTS SOC	37	183	1960	60010
	J AM OIL CHEMISTS SOC	38	169	1961	61006
	J AM OIL CHEMISTS SOC	38	605	1961	61014
	J AM OIL CHEMISTS SOC	38	138	1961	61015
	J AM OIL CHEMISTS SOC	38	361	1961	61019
HARROLD SP	J PHYS CHEM	63	317	1959	59001
	J COLLOID SCI	15	280	1960	60004
	TRANS FARADAY SOC	60	202	1964	64003
	TRANS FARADAY SOC	62	994	1966	66014
HARTLEY GS	Z PHYSIK CHEM (LEIPZIG)	170	321	1934	34001
	TRANS FARADAY SOC	31	183	1935	35001
	TRANS FARADAY SOC	32	795	1936	36001
	J AM CHEM SOC	58	2347	1936	36002
	J CHEM SOC			1968	38001
	NATURE	142	161	1938	38003
	TRANS FARADAY SOC	34	1288	1938	38005
	TRANS FARADAY SOC	35	1109	1939	39001
	KOLLOID-Z	88	22	1939	39005
	J CHEM SOC			1828	1939
	TRANS FARADAY SOC	37	130	1941	41001
	ANN REP PROGR CHEM (CH. S. LONDON)	45	33	1948	48002
	NATURE	163	767	1949	49010
	CHEM IND (LONDON)	24	1012	1964	64030
HARVA O	FINSKA KEMISTSAMFUNDETS MEDD	25	257	1943	43005
	TRANS FARADAY SOC	49	980	1953	53012
	REC TRAV CHIM	75	112	1956	56018
HARWOOD HJ	J AM CHEM SOC	69	2095	1947	47003
	J AM CHEM SOC	71	672	1949	49009
	J AM CHEM SOC	71	671	1949	49013
	J AM CHEM SOC	73	3353	1951	51006
	J AM CHEM SOC	74	2061	1952	52001
HASAN A	ACTA CHEM SCAND	6	440	1952	52007
HATTORI K	BULL CHEM SOC JAPAN	36	1250	1963	63006
HAYANO S	KOLLOID-Z	181	139	1962	62023
HAYDON DA	TRANS FARADAY SOC	54	698	1958	58012
	PHIL TRANS ROY SOC LONDON, SER A	252	225	1960	60011
	TRANS FARADAY SOC	58	1233	1962	62004
HEALY TW	J PHYS CHEM	68	3562	1964	64035
	TRANS SME AIME			321	1964
HENNE AL	J AM CHEM SOC	73	2323	1951	51011
HERMANS JJ	KONINKI NED AKAD WETEN. PROC SER B	58	91	1955	55001
	J PHYS CHEM	59	576	1955	55002
	KONINKI NED AKAD WETEN. PROC SER B	58	97	1955	55009
	KONINKI NED AKAD WETEN. PROC SER B	59	190	1956	56002
	KONINKI NED AKAD WETEN. PROC SER B	59	162	1956	56007
	KONINKI NED AKAD WETEN. PROC SER B	59	298	1956	56011
HERRMANN KW	J PHYS CHEM	66	295	1962	62005
	J PHYS CHEM	67	935	1963	63018
	J PHYS CHEM	68	1540	1964	64006
	J COLLOID INTERFACE SCI	22	352	1966	66013
HERRMANNWK	J PHYS CHEM	70	2909	1966	66039
HERZFELD SH	J PHYS COLLOID CHEM	54	271	1950	50008
	J PHYS CHEM	56	953	1952	52015
	J PHYS CHEM	56	959	1952	52017
HESS K	KOLLOID-Z	88	40	1939	39009
HICKSON J	PROC INTERN CONGR SURFACE ACTIVITY	2ND		1957	57024
HIEBERT EN	J COLLOID SCI	1	385	1946	46020
HIGHAM EH	RES CORRESPONDENCE	7	1	1955	55018
HIGUCHI T	J AM PHARM ASSOC	43	465	1954	54015
HISKEY CF	J COLLOID SCI	9	243	1954	54011
HOERR CW	J AM CHEM SOC	64	2824	1942	42001
	J AM CHEM SOC	64	772	1942	42002
	J AM CHEM SOC	64	97	1942	42005
	J AM CHEM SOC	65	976	1943	43002
	J AM CHEM SOC	65	328	1943	43009

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	J AM CHEM SOC	68	2460	1946	46001
	J AM CHEM SOC	69	883	1947	47002
	J COLLOID SCI	15	427	1960	60001
HOEVE CAJ	J PHYS CHEM	61	593	1957	57002
HOFFMAN EJ	J AM CHEM SOC	64	97	1942	42005
	J AM CHEM SOC	64	2067	1942	42008
	J AM CHEM SOC	64	498	1942	42009
HOLLAHAN JR	J PHYS CHEM	63	757	1959	59023
HOLMBERG P	ACTA CHEM SCAND	19	573	1965	65023
HOLTZER A	J AM CHEM SOC	83	4865	1961	61016
	J PHYS CHEM	69	3718	1965	65006
HONIG JG	J PHYS CHEM	58	201	1954	54009
	J PHYS CHEM	60	1108	1956	56012
	J PHYS CHEM	60	1114	1956	56013
HORI R	BULL CHEM SOC JAPAN	34	237	1961	61008
HOSOKAWA S	BULL CHEM SOC JAPAN	35	1050	1962	62033
	BULL CHEM SOC JAPAN	36	204	1963	63034
HOULTON HG	J AM CHEM SOC	60	544	1938	38008
HOYER HW	J PHYS CHEM	61	818	1957	57004
	J PHYS CHEM	65	1804	1961	61001
	J PHYS CHEM	65	1807	1961	61002
	J PHYS CHEM	68	3494	1964	64007
HSIAO L	J PHYS CHEM	59	362	1955	55020
	J PHYS CHEM	60	657	1956	56014
HUBBARD HM	J AM CHEM SOC	76	4300	1954	54014
HUBBARD WD	J PHYS CHEM	57	808	1953	53009
	J PHYS CHEM	58	1163	1954	54002
	J COLLOID SCI	10	428	1955	55003
	J PHYS CHEM	61	371	1957	57003
	J RES NAT BUR STD A	59	113	1957	57006
	J RES NAT BUR STD A	68	359	1964	64043
HUDSON JB	J COLLOID SCI	12	523	1957	57010
HUFF H	J COLLOID SCI	3	511	1948	48004
HUGO WB	J PHYS COLLOID CHEM	55	311	1951	51004
HUISMAN HF	J PHARM PHARMACOL	12	447	1960	60026
	KONINKI NED AKAD WETEN. PROC SER B	67	367	1964	64045
	KONINKI NED AKAD WETEN. PROC SER B	67	376	1964	64046
	KONINKI NED AKAD WETEN. PROC SER B	67	388	1964	64047
	KONINKI NED AKAD WETEN. PROC SER B	67	407	1964	64048
HUTCHINSON E	J PHYS CHEM	58	1124	1954	54012
	Z PHYSIK CHEM (FRANKFURT)	11	165	1957	57021
	Z PHYSIK CHEM (FRANKFURT)	21	38	1959	59018
	Z PHYSIK CHEM (FRANKFURT)	31	397	1962	62016
	J CHEM EDUC	40	472	1963	63029
	J PHYS CHEM	68	2818	1964	64037
	J PHYS CHEM	70	3502	1966	66026
IDA O	J CHEM SOC JAPAN, PURE CHEM SEC	77	905	1966	66032
IFUKU N	REV PHYS CHEM JAPAN	35	32	1965	65036
	J CHEM SOC JAPAN, PURE CHEM SEC	87	329	1966	66033
IKEDA S	BULL CHEM SOC JAPAN	34	1236	1961	61012
	BULL CHEM SOC JAPAN	35	240	1962	62022
INO T	BULL CH SOC JAPAN	30	760	1957	57033
INOUE H	J CHEM SOC JAPAN, PURE CHEM SEC	78	636	1957	57019
	J CHEM SOC JAPAN, PURE CHEM SEC	79	348	1958	58018
	J CHEM SOC JAPAN, PURE CHEM SEC	79	345	1958	58019
	J CHEM SOC JAPAN, PURE CHEM SEC	79	48	1958	58028
	J COLLOID SCI	15	268	1960	60006
	KOLLOID-Z	183	68	1962	62010
	KOLLOID-Z	196	1	1964	64019
	KOLLOID-Z	195	93	1964	64021
	PROC INTERN CONGR SURFACE ACTIVITY	47H	N	1964	64040
	J PHYS CHEM	70	1108	1965	65033
ISEMURA T	BULL CHEM SOC JAPAN	34	1236	1961	610
	BULL CHEM SOC JAPAN	35	1737	1962	62012
	BULL CHEM SOC JAPAN	35	240	1962	62022
	BULL CHEM SOC JAPAN	36	1250	1963	63006
IWAMATSU I	BULL CHEM SOC JAPAN	35	1050	1962	62033
	BULL CHEM SOC JAPAN	36	204	1963	63034
JACOBS J	J PHARM PHARMACOL	18	925	1966	66019
JAKOB CW	J PHYS CHEM	67	2075	1963	63001
JAMES JW	PROC INTERN CONGR SURFACE ACTIVITY	3RD	227	1960	60020
JAN ZA	J PHARM PHARMACOL	15	825	1963	63021
JOHNSON JS	J PHYS CHEM	68	81	1964	64017
JOHNSON KE	J AM CHEM SOC	66	9	1944	44003
JOHNSON WF	J PHYS CHEM	50	440	1946	46006
	J PHYS COLLOID CHEM	51	636	1947	47013
	J PHYS COLLOID CHEM	52	22	1948	48025
	J AM CHEM SOC	73	4563	1951	51002
	J AM CHEM SOC	74	20	1952	52002
	J AM CHEM SOC	74	22	19	52003

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JONES E	PHIL MAG	4	841	1927	27002
JONES TG	TRANS FARADAY SOC	49	980	1953	53012
	RES CORRESPONDENCE	8	1	1955	55011
KAKIUCHI K	BULL CHEM SOC JAPAN	36	1250	1963	63006
KAPAUAN P	J COLLOID SCI	16	481	1961	61005
	J PHYS CHEM	70	783	1966	66002
KARNAUKH AM	MASLOB ZHIR PROM	29	22	1963	63039
KARTZMARK EM	CAN J CHEM	40	839	1962	62034
KASHIWAGI KM	J COLLOID SCI	13	618	1958	58021
KASHIWAGI M	BULL CHEM SOC JAPAN	32	624	1959	59017
KATO Y	CHEM PHARM BULL (TOKYO)	11	1202	1963	63037
KATSURA K	J PHYS CHEM	68	1568	1964	64011
KAUFMAN S	J COLLOID SCI	12	465	1957	57026
	J PHYS CHEM	62	1257	1958	58025
	WORLD PETROL CONGR, PROC	V 6	18	1959	59021
	J COLLOID SCI	17	231	1962	62025
	J PHYS CHEM	68	2814	1964	64041
KAWAMURA S	YAKUGAKU ZASSHI	84	246	1964	64034
KAZUO T	J CHEM SOC JAPAN, PURE CHEM SEC	79	48	1958	58028
KEIM GI	IND ENG CHEM	36	610	1944	44001
KIESSIG H	KOLLOID-Z	88	40	1939	39009
KINNEY FB	J AM OIL CHEMISTS SOC	36	332	1959	59009
	J AM OIL CHEMISTS SOC	37	18	S1960	60010
	J AM OIL CHEMISTS SOC	38	138	1961	61015
KINOSHITA K	J PHYS CHEM	63	648	1959	59013
KITAHARA A	BULL CHEM SOC JAPAN	28	234	1955	55019
	BULL CHEM SOC JAPAN	29	1	S1956	56015
	J COLLOID SCI	12	342	1957	57027
	BULL CHEM SOC JAPAN	30	586	1957	57028
	BULL CHEM SOC JAPAN	31	288	1957	57029
	BULL CHEM SOC JAPAN	31	653	1958	58022
	J PHYS CHEM	66	363	1962	62024
	KOGYO KAGAKU ZASSHI	68	2058	1965	65002
KLAMANN D	PROC INTERN CONGR SURFACE ACTIVITY	3RD	1	1960	60016
	PROC INTERN CONGR SURFACE ACTIVITY	3RD	27	1960	60017
KLEVENS HB	J CHEM PHYS	14	216	1946	46002
	J CHEM PHYS	14	742	1946	46007
	J CHEM PHYS	14	480	1946	46010
	J PHYS COLLOID CHEM	51	567	1946	46012
	J COLLOID SCI	2	301	1947	47005
	J PHYS COLLOID CHEM	52	130	1948	48005
	J AM OIL CHEMISTS SOC	26	456	1949	49003
	J PHYS COLLOID CHEM	54	1012	1950	50003
	J AM CHEM SOC	72	3780	1950	50004
	ANAL CHEM	22	1141	1950	50006
	CHEM REV	47	1	1950	50007
	MEM SERV CHIM ETAT (PARIS)	37	13	1952	52004
	J AM CHEM SOC	74	4624	1952	52005
	KOLLOID-Z	128	61	1952	52008
	J AM OIL CHEMISTS SOC	30	74	1953	53010
	PROC INTERN CONGR SURFACE ACTIVITY	1ST	1	1954	54004
	J CHIM PHYS	51	1	1954	54010
	NATURE	176	879	1955	55017
	J PHYS CHEM	60	1245	1956	56001
	MEM SERV CHIM ETAT (PARIS)	41	363	1956	56010
	PROC INTERN CONGR SURFACE ACTIVITY	2ND	395	1957	57017
	KOLLOID-Z	158	53	1958	58011
KLING W	PROC INTERN CONGR SURFACE ACTIVITY	2ND	295	1957	57022
KOBAYASHI T	J PHYS CHEM	66	363	1962	62024
KOLBEL H	ANGEW CHEM	71	211	1959	59022
	PROC INTERN CONGR SURFACE ACTIVITY	3RD	1	1960	60016
	PROC INTERN CONGR SURFACE ACTIVITY	3RD	27	1960	60017
KOLTHOFF IM	J PHYS CHEM	50	440	1946	46006
	J PHYS COLLOID CHEM	51	636	1947	47013
	J PHYS COLLOID CHEM	52	915	1948	48016
	J PHYS COLLOID CHEM	52	22	1948	48025
	J PHYS COLLOID CHEM	53	424	1949	49005
	J POLYMER SCI	5	201	1950	50002
	J POLYMER SCI	5	191	1950	50005
	J AM CHEM SOC	73	4563	1951	51002
	J AM CHEM SOC	74	20	1952	52002
	J AM CHEM SOC	74	22	1952	52003
KOMOR JA	J AM OIL CHEMISTS SOC	43	435	1966	66022
KONDO A	J CHEM SOC JAPAN, PURE CHEM SEC	77	905	1966	66032
KONDO T	J CHEM SOC JAPAN, PURE CHEM SEC	77	1236	1956	56020
	BULL CHEM SOC JAPAN	30	905	1957	S7032
	BULL CHEM SOC JAPAN	30	760	1957	57033
	J CHEM SOC JAPAN, PURE CHEM SEC	80	818	1959	59024

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KONSTANTIN. VV	J CHEM SOC JAPAN, PURE CHEM SEC	80	823	1959	59026
KOSTOVA NZ	KOGYO KAGAKU ZASSHI	68	2058	1965	65002
KRAUS CA	KHIM TEKHNOL	5	61	1960	60033
	KOLOOID ZH	26	76	1964	64051
	J AM CHEM SOC	68	1137	1946	46018
	J AM CHEM SOC	69	1835	1947	47001
	J AM CHEM SOC	70	3803	1948	48010
	J AM CHEM SOC	70	3049	1948	48028
	J AM CHEM SOC	71	95	1949	49014
	J AM CHEM SOC	71	309	1949	49017
	J AM CHEM SOC	71	1455	1949	49018
	J AM CHEM SOC	72	3676	1950	50013
	J AM CHEM SOC	73	2173	1951	51009
	J AM CHEM SOC	73	799	1951	51013
	J AM CHEM SOC	73	1129	1951	51014
	J AM CHEM SOC	73	2170	1951	51016
	PROC NAT ACAD SCI U S	39	1213	1953	53011
KRIUENTSOV WI	ZAVODSKAYA LAB	24	158	1958	58030
KRIZEK H	J COLLOID SCI	6	576	1951	51010
KRYUKOVA AS	KHIM TEKHNOL	5	61	1960	60033
KUCHER RV	COLLOID J (USSR)	14	243	1952	52018
KUHN DW	COLLOID J (USSR)	14	311	1952	52019
KUHN P	J AM CHEM SOC	72	3676	1950	50013
KUPPUSAMI J	ANGEW CHEM	71	211	1959	59022
KURIYAMA K	NATURE	208	780	1965	65028
	J CHEM SOC JAPAN, PURE CHEM SEC	78	1568	1957	57018
	J CHEM SOC JAPAN, PURE CHEM SEC	78	1573	1957	57020
	J CHEM SOC JAPAN, PURE CHEM SEC	79	348	1958	58018
	J CHEM SOC JAPAN, PURE CHEM SEC	79	345	1958	58019
	J CHEM SOC JAPAN, PURE CHEM SEC	79	48	1958	58028
	J COLLOID SCI	15	268	1960	60006
	KOLLOID-Z	180	55	1962	62009
	KOLLOID-Z	183	68	1962	62010
	KOLLOID-Z	181	144	1962	62011
	KOLLOID-Z	191	48	1963	63012
KURZ J L	J PHYS CHEM	66	2239	1962	62040
KURZENDORFER P	PROC INTERN CONGR SURFACE ACTIVITY	3RD	1	1960	60016
KUSHNER LM	J PHYS CHEM	57	808	1953	53009
	J PHYS CHEM	58	1163	1954	54002
	J COLLOID SCI	10	428	1955	55003
	J PHYS CHEM	61	371	1957	57003
	J RES NAT BUR STD A	59	113	1957	57006
KWARTLER CE	J COLLOID SCI	8	385	1953	53007
LACH JL	J AM PHARM ASSOC	43	465	1954	54015
LAKSHMINAR. GR	CAN J CHEM	40	839	1962	62034
	CAN J CHEM	43	1729	1965	65024
LAL H	J COLLOID SCI	8	414	1953	53013
LANGE H	KOLLOID-Z	121	66	1951	51005
	KOLLOID-Z	131	96	1953	53005
	PROC INTERN CONGR SURFACE ACTIVITY	2ND	295	1957	57022
	PROC INTERN CONGR SURFACE AOIVITY	3RD	279	1960	60012
	FETTE, SEIFEN, ANSTRICHMI	64	457	1962	62018
	PROC INTERN CONGR SURFACE ACTIVITY	4TH	1	1964	64039
LARSEN EC	Z ELEKTROCHEM	44	651	1938	38004
LELONG ALM	J AM CHEM SOC	73	5411	1951	51003
	J PHYS CHEM	59	1185	1955	55021
	ANALES ASOC QUIM ARGENT	53	39	1965	65034
	ANALES ASOC QUIM ARGENT	53	11	1965	65035
LESHCHENKO ZY	MASLOB ZHIR PROM	26	24	1960	60030
	MASLOB ZHIR PROM	28	20	1962	62039
	MASLOB ZHIR PROM	29	19	1963	63038
LESYUIS AA	MASLOB ZHIR PROM	29	22	1963	63039
	BULL CHEM SOC JAPAN	28	227	1955	55008
	J CHINESE CHEM SOC	4	28	1957	57005
	J CHINESE CHEM SOC	4	21	1957	57008
LINDSTROM RE	MEDD.ABO AKAD.FYS.KEM.IN.	SP NO	3	1941	41002
	KOLLOID-Z	94	42	1941	41004
LINGAFELTER EC	J AM CHEM SOC	65	686	1943	43001
	J AM CHEM SOC	65	698	1943	43004
	J AM CHEM SOC	68	1490	1946	46003
	J AM CHEM SOC	70	1989	1948	48007
	J AM CHEM SOC	71	1325	1949	49004
	J AM CHEM SOC	73	5411	1951	51003
	J COLLOID SCI	10	71	1955	55025
LIQUORI AM	TRANS FARADAY SOC	55	1975	1959	59015
	PROC INTERN CONGR SURFACE ACTIVITY	3RD	302	1960	60024
	J MOL BIOL	3	202	1961	61023
	J PHYS CHEM	65	1991	1961	61C24
	PONTIF ACAD SCIENTIAR. SCRIPTA VAR.	22	/12/1	1962	62021
	RIC SCI	6	71	1964	64033

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	J PHYS CHEM	68	2709	1964	64044
LONG FA	J AM CHEM SOC	59	2197	1937	37005
	J AM CHEM SOC	62	1496	1940	40005
	J AM CHEM SOC	63	84	1941	41006
LORENZ PB	J PHYS CHEM	60	657	1956	56014
LOTTERMOSER A	KOLLOID-Z	63	175	1933	33003
	KOLLOID-Z	63	49	1933	33004
	KOLLOID-Z	73	276	1935	35002
	TRANS FARADAY SOC	31	200	1935	35004
	KOLLOID-BEIH.	45	303	1937	37004
LOVELL VM	ANAL CHEM	38	1926	1966	66024
LUCK W	PROC INTERN CONGR SURFACE ACTIVITY	3RD	264	1960	60023
LUDLUM DB	J PHYS CHEM	60	1240	1956	56005
MACFARLANE CB	J PHARM PHARMACOL	14	100	1962	62027
	J PHARM PHARMACOL	17	65	1965	65016
MALIK WU	INDIAN J CHEM	3	441	1965	65021
	J AM OIL CHEMISTS SOC	43	446	1966	66023
MALSCH J	Z PHYSIK CHEM (LEIPZIG)	170	321	1934	34001
MANCHESTER KE	J PHYS CHEM	58	1124	1954	54012
MANDELL L	ACTA CHEM SCAND	17	111	1963	63033
MANKOWICH AM	J AM OIL CHEMISTS SOC	41	449	1964	64010
	J AM OIL CHEMISTS SOC	43	615	1966	66021
MANNING DJ	J AM OIL CHEMISTS SOC	43	133	1966	66001
MARKAN AL	ZAVODSKAYA LAB	24	158	1958	58030
MARKINA ZN	KOLLOID ZH	26	76	1964	64051
MARMO A	J PHYS CHEM	65	1804	1961	61001
	J PHYS CHEM	65	1807	1961	61002
MARON SH	J COLLOID SCI	9	382	1954	54006
MARUTA I	J CHEM SOC JAPAN, PURE CHEM SEC	82	1657	1961	61009
	J CHEM SOC JAPAN, PURE CHEM SEC	82	1512	1961	61010
	J CHEM SOC JAPAN, PURE CHEM SEC	82	1660	1961	61011
	J CHEM SOC JAPAN, PURE CHEM SEC	83	732	1962	62007
	J CHEM SOC JAPAN, PURE CHEM SEC	83	786	1962	62026
	J CHEM SOC JAPAN, PURE CHEM SEC	83	395	1962	62028
	J CHEM SOC JAPAN, PURE CHEM SEC	83	782	1962	62029
	J CHEM SOC JAPAN, PURE CHEM SEC	83	788	1962	62030
	J CHEM SOC JAPAN, PURE CHEM SEC	83	858	1962	62031
	J CHEM SOC JAPAN, PURE CHEM SEC	83	861	1962	62032
	KOGYO KAGAKU ZASSHI	68	1090	1965	65017
MATHAI K	TRANS FARADAY SOC	62	759	1966	66031
	TRANS FARADAY SOC	62	750	1966	66037
MATIJEVIC E	CROAT CHEM ACTA	29	431	1957	57001
	TRANS FARADAY SOC	54	587	1958	58020
MATSUMOTO T	J SCI HIROSHIMA UNIV, SER A-II	21	51	1957	57025
MATTOON RW	J AM CHEM SOC	68	220	1946	46013
	J COLLOID SCI	1	105	1946	46014
	J CHEM PHYS	15	763	1947	47008
MATUURA R	BULL CHEM SOC JAPAN	35	1050	1962	62033
	BULL CHEM SOC JAPAN	36	204	1963	63034
	BULL CHEM SOC JAPAN	36	813	1963	63035
	BULL CHEM SOC JAPAN	38	373	1965	65020
MAURER EW	J AM OIL CHEMISTS SOC	36	241	1959	59004
	J AM OIL CHEMISTS SOC	41	205	1964	64002
MCBAIN JW	J AM CHEM SOC	57	1905	1935	35007
	J PHYS CHEM	40	493	1936	36005
	J AM CHEM SOC	63	670	1941	41005
	J PHYS CHEM	46	10	1942	42006
	IND ENG CHEM	34	915	1942	42007
	J AM CHEM SOC	65	2072	1943	43006
	J PHYS CHEM	47	94	1943	43008
	J AM CHEM SOC	66	9	1944	44003
	J COLLOID SCI	1	127	1946	46008
	REC TRAV CHIM	65	601	1946	46016
	J AM CHEM SOC	68	731	1946	46022
	J AM CHEM SOC	69	334	1947	47007
	J COLLOID SCI	3	425	1948	48009
	J PHYS COLLOID CHEM	52	881	1948	48011
	J AM CHEM SOC	70	3838	1948	48015
	J PHYS COLLOID CHEM	52	12	1948	48026
	J PHYS COLLOID CHEM	55	311	1951	51004
MCBAIN MEL	J AM CHEM SOC	61	3210	1939	39011
	J COLLOID SCI	10	223	1955	55023
	J PHYS CHEM	47	196	1943	43007
MCCORKLE MR	J AM CHEM SOC	65	328	1943	43009
MCDOWELL MJ	J AM CHEM SOC	73	2173	1951	51009
	J AM CHEM SOC	73	2170	1951	51016
MCHAN H	J AM CHEM SOC	70	3838	1948	48015
MCNEILL W	J PHYS CHEM	56	701	1952	52013
MEADER AL	IND ENG CHEM	44	1636	1952	52014

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MEEHAN EJ	J POLYMER SCI	5	201	1950	50002
	J POLYMER SCI	5	191	1950	50005
MEGURO K	J CHEM SOC JAPAN, PURE CHEM SEC	77	1236	1956	56020
	BULL CHEM SOC JAPAN	30	905	1957	57032
	BULL CHEM SOC JAPAN	30	760	1957	57033
	J CHEM SOC JAPAN, PURE CHEM SEC	80	818	1959	59024
	J CHEM SOC JAPAN, PURE CHEM SEC	80	821	1959	59025
	J CHEM SOC JAPAN, PURE CHEM SEC	80	823	1959	59026
	J CHEM SOC JAPAN, PURE CHEM SEC	77	905	1966	66032
MELE A	NATURE	184	1482	1959	59014
	TRANS FARADAY SOC	55	1975	1959	59015
	J PHYS CHEM	63	650	1959	59016
MERRILL RC	J PHYS COLLOID CHEM	52	774	1948	48024
MERRILL RC JR	J AM CH M S	63	670	1941	41005
	J PHYS CHEM	46	10	1942	42006
	IND ENG CHEM	34	915	1942	42007
	J PHYS COLLOID CHEM	52	12	1948	48026
METCALF AD	J COLLOID SCI	17	523	1962	62015
MEYER HG	J PHYS CHEM	70	783	1966	66002
MIJNLIEFF PF	NATURE	208	889	1965	65029
MILES GD	IND ENG CHEM	36	610	1944	44001
	J PHYS CHEM	48	57	1944	44002
	J PHYS CHEM	49	71	1945	45002
MILLER ML	J COLLOID SCI	13	411	1958	58001
MITTELMANN R	J CHEM PHYS	15	763	1947	47008
	J PHYS COLLOID CHEM	53	1350	1949	49006
	J COLLOID SCI	4	367	1949	49007
MIURA M	J SCI HIROSHIMA UNIV, SER A-II	21	51	1957	57025
	J SCI HIROSHIMA UNIV, SER A-II	22	57	1958	58023
	J SCI HIROSHIMA UNIV, SER A-II	28	41	1964	64036
MIYAMOTO S	BULL CHEM SOC JAPAN	33	375	1960	60029
	BULL CHEM SOC JAPAN	33	371	1960	60035
MODI HJ	J ELECTROCHEM SOC	106	336	1959	59019
MOLYNEUX P	TRANS FARADAY SOC	61	1043	1965	65003
MOULE D	TRANS FARADAY SOC	54	1638	1958	58006
	CAN J CHEM	37	2083	1959	59007
	CAN J CHEM	37	2086	1959	59008
MUKERJEE P	J AM CHEM SOC	77	2937	1955	55015
	J PHYS CHEM	62	1390	1958	58013
	J PHYS CHEM	62	1397	1958	58014
	J PHYS CHEM	62	1400	1958	58015
	J PHYS CHEM	62	1404	1958	58016
	J PHYS CHEM	67	190	1963	63032
	J PHYS CHEM	70	783	1966	66002
	J PHYS CHEM	70	2138	1966	66004
	J PHYS CHEM	70	2144	1966	66005
	J PHYS CHEM	70	2150	1966	66006
MUKHERJI BK	J PHYS CHEM	64	1	1960	60028
MULLEY BA	J COLLOID SCI	17	523	1962	62015
	J COLLOID SCI	19	201	1964	64009
MURRAY RC	TRANS FARADAY SOC	31	183	1935	35001
MYSELS EK	J COLLOID SCI	20	315	1965	65018
MYSELS KJ	TRANS FARADAY SOC	51	728	1955	55005
	J PHYS CHEM	59	325	1955	55014
	J AM CHEM SOC	77	2937	1955	55015
	J COLLOID SCI	10	507	1955	55016
	J PHYS CHEM	62	1390	1958	58013
	J PHYS CHEM	62	1400	1958	58015
	J PHYS CHEM	63	1696	1959	59002
	J PHYS CHEM	63	1781	1959	59003
	J COLLOID SCI	16	481	1961	61005
	J COLLOID SCI	16	462	1961	61017
	J COLLOID SCI	16	474	1961	61018
	J COLLOID SCI	20	315	1965	65018
	J PHYS CHEM	69	1466	1965	65019
	J COLLOID SCI	21	331	1966	66007
NAKADATE S	REP INST SCI TECH UNIV TOKYO	7	401	1953	53001
NAKAGAKI M	J CHEM SOC JAPAN, PURE CHEM SEC	72	113	1951	51018
	BULL CHEM SOC JAPAN	37	817	964	64025
	YAKUGAKU ZASSHI	84	246	1964	64034
NAKAGAWA T	J CHEM SOC JAPAN, PURE CHEM SEC	78	1568	1957	57018
	J CHEM SOC JAPAN, PURE CHEM SEC	78	636	1957	57019
	J CHEM SOC JAPAN, PURE CHEM SEC	78	1573	1957	57020
	SHINOGI KENKYUSHO NEMPO	8	805	1958	58017
	J CHEM SOC JAPAN, PURE CHEM SEC	79	348	1958	58018
	J CHEM SOC JAPAN, PURE CHEM SEC	79	345	1958	58019
	J CHEM SOC JAPAN, PURE CHEM SEC	79	48	1958	58028
	J COLLOID SCI	15	268	1960	60006
	PROC INTERN CONGR SURFACE ACTIVITY	3RD	636	1960	60007
	KOLLOID-Z	168	132	1960	60009
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	KOLLOID-Z	188	47	1963	63009
	KOLLOID-Z	187	44	1963	63010
	KOLLOID-Z	191	48	1963	63012
	KOLLOID-Z	194	143	1964	64015
	KOLLOID-Z	195	93	1964	64021
	PROC INTERN CONGR SURFACE ACTIVITY	4TH	N	1964	64040
	J PHYS CHEM	70	1108	1965	65033
NAKAYAMA H	J COLLOID SCI	18	705	1963	63036
	J PHYS CHEM	70	3502	1966	66026
NASH T	CHEM IND (LONDON)		590	1958	58027
NATALE I	ANALES ASOC QUIM ARGENT	53	11	1965	65035
NAUMAN RV	J PHYS CHEM	68	3498	1964	64001
NEFF LL	J AM CHEM SOC	70	1989	1948	48007
NEWTON JM	J PHARM PHARMACOL	12	447	1960	60026
NICOLESU A	REV CHIM AC REP POP ROM.	6	309	1961	61029
NINOMIYA Y	BULL CHEM SOC JAPAN	37	817	1964	64025
NOEL DR	J AM CHEM SOC	74	2061	1952	52001
NOGUCHI J	BULL CHEM SOC JAPAN	34	1236	1961	61012
NUTTING GC	J AM CHEM SOC	59	2197	1937	37005
	J AM CHEM SOC	62	1496	1940	40005
	J AM CHEM SOC	63	84	1941	41006
OGDEN CP	PROC ROY SOC	273	84	1963	63014
	TRANS FARADAY SOC	61	583	1965	65005
OHBA N	BULL CHEM SOC JAPAN	30	905	1957	57032
	BULL CHEM SOC JAPAN	30	760	1957	57033
OHKI K	J PHYS CHEM	70	3437	1966	66027
OKUYAMA H	BULL CHEM SOC JAPAN	27	259	1954	54008
	BULL CHEM SOC JAPAN	30	186	1957	57023
OLENIKOVA ZV	MASLOB ZHIR PROM	26	26	1960	60031
OLIVIER JP	J PHYS CHEM	63	1671	1959	59020
OLOANE JK	J AM CH M SOC	73	5411	1951	51003
OPPENHEIMER H	J CHEM PHYS	15	496	1947	47012
	J AM CHEM SOC	71	808	1949	49011
OSIPOW L	PROC INTERN CONGR SURFACE ACTIVITY	2ND		1957	57024
OSTER G	J COLLOID SCI	9	243	1954	54011
OSUGI J	REV PHYS CHEM JAPAN	35	32	1965	65036
	J CHEM SOC JAPAN, PURE CHEM SEC	87	329	1966	66033
OTTER RJ	J COLLOID SCI	16	462	1961	61017
	J COLLOID SCI	16	474	1961	61018
OTTEWILL RH	TRANS FARADAY SOC	62	750	1966	66037
	J CHEM SOC		1712	1958	58031
	TRANS FARADAY SOC	57	1627	1961	61004
	J COLLOID INTERFACE SCI	21	522	1966	66028
	TRANS FARADAY SOC	62	759	1966	66031
OYAMA T	J CHEM SOC JAPAN, PURE CHEM SEC	79	348	1958	58018
	J CHEM SOC JAPAN, PURE CHEM SEC	79	345	1958	58019
PACKTER A	J PHARM PHARMACOL	15	317	1963	63030
PANKHURST KGA	TRANS FARADAY SOC	42	523	1946	46021
PAQUETTE RG	J AM CHEM SOC	65	686	1943	43001
PARKER RA	J RES NAT BUR STD A	59	113	1957	57006
PARREIRA HC	ANALIS ACAD BRASIL CIENC	32	207	1960	60015
	J COLLOID INTERFACE SCI	21	522	1966	66028
PARRY GA	J CHEM SOC		626	1935	35008
PATEL RM	J PHARM SCI	55	1345	1966	66017
PATTERSON GD	J PHYS CHEM	57	247	1953	53003
PETHICA BA	CROAT CHEM ACTA	29	431	1957	57001
	TRANS FARADAY SOC	54	587	1958	58020
PHILIPPOFF W	PROC INTERN CONGR SURFACE ACTIVITY	3RD	227	1960	60020
PHILLIPS JN	KOLLOID-Z	88	40	1939	39009
	TRANS FARADAY SOC	51	728	1955	55005
	J PHYS CHEM	59	325	1955	55014
	TRANS FARADAY SOC	51	561	1955	55024
	TRANS FARADAY SOC	54	698	1958	58012
PICCIONE GA	J PHYS CHEM	46	662	1942	42003
PILPEL N	J COLLOID SCI	9	285	1954	54001
	J PHYS CHEM	60	779	1956	56004
	TRANS FARADAY SOC	57	1426	1961	61013
	CHEM REV	63	221	1963	63002
	NATURE	204	378	1964	64022
POWNEY J	TRANS FARADAY SOC	31	1510	1935	35005
	TRANS FARADAY SOC	33	851	1937	37003
	TRANS FARADAY SOC	34	372	1938	38006
PRINCEN LH	J PHYS CHEM	63	1696	1959	59002
	J PHYS CHEM	63	1781	1959	59003
PRINS W	J PHYS CHEM	59	576	1955	55002
	KONINKI NED AKAD WETEN. PROC SER B	59	190	1956	56002
	KONINKI NED AKAD WETEN. PROC SER B	59	162	1956	56007
	KONINKI NED AKAD WETEN. PROC SER B	59	298	1956	56011

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	ZESZTY NAUK UNIW JAGIEL	211	209	1966	66035
RADLEY JA	PROC INTERN CONGR SURFACE ACTIVITY	3RD	246	1960	60021
RAISON M	COMPT REND	235	1129	1952	52016
	PROC INTERN CONGR SURFACE ACTIVITY	1ST	1	1954	54004
	J CHIM PHYS	51	1	1954	54010
RALSTON AW	J AM CHEM SOC	64	2824	1942	42001
	J AM CHEM SOC	64	772	1942	42002
	J AM CHEM SOC	64	97	1942	42005
	J AM CHEM SOC	64	2067	1942	42008
	J AM CHEM SOC	64	498	1942	42009
	J AM CHEM SOC	65	976	1943	43002
	J AM CHEM SOC	65	328	1943	43009
	J AM CHEM SOC	68	2460	1946	46001
	J AM CHEM SOC	69	883	1947	47002
	J AM CHEM SOC	69	2095	1947	47003
	J AM CHEM SOC	70	977	1948	48014
	J AM CHEM SOC	70	980	1948	48019
	J AM CHEM SOC	70	2918	1948	48020
	J AM CHEM SOC	70	983	1948	48021
	J PHYS COLLOID CHEM	52	1494	1948	48023
	J AM CHEM SOC	70	436	1948	48027
	J AM CHEM SOC	71	2145	1949	49008
	J AM CHEM SOC	71	672	1949	49009
	J AM CHEM SOC	71	671	1949	49013
RAY A	J PHYS CHEM	67	190	1963	63032
	J PHYS CHEM	70	2138	1966	66004
	J PHYS CHEM	70	2144	1966	66005
	J PHYS CHEM	70	2150	1966	66006
REED RM	J AM CHEM SOC	57	570	1935	35006
	J AM CHEM SOC	58	322	1936	36006
REEVES RL	J PHYS CHEM	69	2357	1965	65030
REITMIER RE	J AM CHEM SOC	62	2375	1940	40001
REYNOLDS CA	J AM CHEM SOC	76	4300	1954	54014
RHODES CT	TRANS FARADAY SOC	61	1043	1965	65003
RICCIERI FM	J PHARM SCI	52	1011	1963	63024
	J PHARM SCI	54	919	1965	65010
ROBINS DC	J PHARM PHARMACOL	15	522	1963	63005
	J PHARM PHARMACOL	15	157	1963	63031
ROBINSON RU	J PHYS CHEM	56	701	1952	52013
ROBSON P	TRANS FARADAY SOC	62	987	1966	66015
ROE CP	J AM CHEM SOC	76	4703	1954	54013
ROSE GRF	CAN J CHEM ENGR	F28	213	1950	50010
ROSENBLUM C	J PHYS CHEM	46	662	1942	42003
ROSS J	IND ENG CHEM	36	610	1944	44001
ROSS S	J COLLOID SCI	8	385	1953	53007
	J COLLOID SCI	12	523	1957	57010
	J PHYS CHEM	61	1261	1957	57031
	J PHYS CHEM	63	1671	1959	59020
RUSH RM	J PHYS CHEM	68	81	1964	64017
SAITO S	BULL CHEM SOC JAPAN	30	186	1957	57023
SAITO T	J CHEM SOC JAPAN, PURE CHEM SEC	82	1512	1961	61010
SAKAI T	J CHEM SOC JAPAN, PURE CHEM SEC	82	1512	1961	61010
SAMIS CS	TRANS FARADAY SOC	32	795	1936	36001
	TRANS FARADAY SOC	34	1288	1938	38005
SASAKI H	BULL CHEM SOC JAPAN	30	186	1957	57023
	BULL CHEM SOC JAPAN	30	326	1957	57030
SATA N	BULL CHEM SOC JAPAN	26	177	1953	53006
SATAKE I	BULL CHEM SOC JAPAN	35	1050	1962	62033
	BULL CHEM SOC JAPAN	36	204	1963	63034
	BULL CHEM SOC JAPAN	36	813	1963	63035
SATANEK J	J AM OIL CHEMISTS SOC	38	169	1961	61006
SATO M	REV PHYS CHEM JAPAN	35	32	1965	65036
	J CHEM SOC JAPAN, PURE CHEM SEC	87	329	1966	66033
SAWYER WM	J PHYS CHEM	62	159	1958	58029
SCHICK MJ	J PHYS CHEM	61	1062	1957	57014
	J COLLOID SCI	17	801	1962	62019
	J PHYS CHEM	66	1326	1962	62020
	J PHYS CHEM	67	1796	1963	63026
	J AM OIL CHEMISTS SOC	40	680	1963	63027
	J COLLOID SCI	18	378	1963	63028
	J PHYS CHEM	68	3585	1964	64020
	J COLLOID SCI	20	464	1965	65011
	J AM OIL CHEMISTS SOC	43	133	1966	66001
	J AM OIL CHEMISTS SOC	43	681	1966	66025
SCHMID G	Z ELEKTROCHEM	44	651	1938	38004
SCHOLBERG HM	J PHYS CHEM	57	923	1953	53014
SCHOTT H	J PHYS CHEM	68	3612	1964	64004
	J PHYS CHEM	70	2966	1966	66036

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SCOTT AB	J AM CHEM SOC	65	692	1943	43003
	J AM CHEM SOC	65	698	1943	43004
SCOTT R	NATURE	167	195	1951	51012
SEARLES J	J PHYS CHEM	40	493	1936	36005
SEBBA F	ANAL CHEM	38	1926	1966	66024
SHAFFER PM	Z PHYSIK CHEM (FRANKFURT)	31	397	1962	62016
SHANE N	J PHYS CHEM	69	968	1965	65025
SHEAFFER VE	J PHYS CHEM	68	2818	1964	64037
SHEDLOVSKY L	IND ENG CHEM	36	610	1944	44001
	J PHYS CHEM	48	57	1944	44002
	J PHYS CHEM	67	2075	1963	63001
SHIGEHARA K	BULL CHEM SOC JAPAN	38	1700	1965	650
	BULL CHEM SOC JAPAN	39	2643	1966	66009
	BULL CHEM SOC JAPAN	39	2332	1966	66010
SHIGEHIRO F	KOGYO KAGAKU ZASSHI	68	1090	1965	65017
SHINODA K	J PHYS CHEM	58	1136	1954	54003
	J PHYS CHEM	58	541	1954	54005
	BULL FAC ENG. YOKOHAMA NAT UNIV	4	77	1955	55004
	J PHYS CHEM	59	432	1955	55007
	J PHYS CHEM	60	1439	1956	56003
	J PHYS CHEM	63	648	1959	59013
	BULL CHEM SOC JAPAN	34	237	1961	61008
	J COLLOID SCI	18	705	1963	63036
	J PHYS CHEM	68	1568	1964	64011
	J PHYS CHEM	70	3502	1966	66026
SHIRAHAMA K	BULL CHEM SOC JAPAN	38	373	1965	65020
SHIRAI M	BULL CHEM SOC JAPAN	28	545	1955	55010
	BULL CHEM SOC JAPAN	29	733	1956	56009
	BULL CHEM SOC JAPAN	30	411	1957	57015
	BULL CHEM SOC JAPAN	30	542	1957	57016
	BULL CHEM SOC JAPAN	31	467	1958	58007
SHISHIDO S	BULL CHEM SOC JAPAN	24	41	1951	51017
SHOLTES EH	J COLLOID SCI	1	385	1946	46020
SHUCK GR	J AM CHEM SOC	71	1325	1949	49004
SHUTE HL	TRANS FARADAY SOC	34	758	1938	38007
SIMON E	J CHEM PHYS	15	496	1947	47012
SINGER K	ANN REP PROGR CH M (HI S. LONDON)	45	51	1948	48003
SINGLETERRY CR	J AM CHEM SOC	70	3965	1948	48017
	J COLLOID SCI	4	537	1949	49012
	J AM CHEM SOC	73	4574	1951	51007
	J PHYS CHEM	58	201	1954	54009
	J PHYS CHEM	60	1108	1956	56012
	J PHYS CHEM	60	1114	1956	56013
	J COLLOID SCI	12	465	1957	57026
	J COLLOID SCI	13	569	1958	58024
	J PHYS CHEM	62	1257	1958	58025
	WORLD PETROL CONGR, PROC	V 6	18	1959	59021
	J PHYS CHEM	68	3453	1964	64042
	J PHYS CH M	68	2709	1964	64044
SIROIS EH	J PHYS CHEM	56	701	1952	52013
SIVERTZ V	J AM CHEM SOC	61	549	1939	39007
	J AM CHEM SOC	62	2375	1940	40001
SMEDS K	ACTA CHEM SCAND	6	441	1952	52006
SMITH FD	J AM OIL CHEMISTS SOC	40	538	1963	63013
SNELL FD	PROC INTERN CONGR SURFACE ACTIVITY	2ND		1957	57024
SOLINAS M	BIOCHEM BIOPHYS ACTA	88	415	1964	64031
	J PHYS CHEM	68	3624	1964	64032
SOMASUNDARAN P	J PHYS CHEM	68	3562	1964	64035
	TRANS SME AIME		321	1964	64038
SPARKS B	TRANS FARADAY SOC	62	3244	1966	66038
SPINGOLA F	J COLLOID SCI	20	732	1965	65012
STANLEY JS	PROC INTERN CONGR SURFACE ACTIVITY	3RD	246	1960	60021
STAUFF J	Z PHYSIK CHEM (LEIPZIG)	183	55	1939	39006
STEARN RS	J CHEM PHYS	14	215	1946	46011
	J CHEM PHYS	14	214	1946	46017
	J CHEM PHYS	15	496	1947	47012
STEIGMAN J	J COLLOID SCI	20	732	1965	65012
	J PHYS CHEM	69	968	1965	65025
STERNBERG RJ	J POLYMER SCI	5	191	1950	50005
STEWART A	TRANS FARADAY SOC	31	208	1935	35003
STEWART JC	RES CORRESPONDENCE	7	1	1955	55018
STIRTON AJ	J PHYS CHEM	60	899	1956	56008
	J PHYS CHEM	62	1083	1958	58003
	J AM OIL CHEMISTS SOC	36	241	1959	59004
	J AM OIL CHEMISTS SOC	37	679	1960	60008
	J AM OIL CHEMISTS SOC	39	55	1962	62008
	J AM OIL CHEMISTS SOC	40	538	1963	63013
	J AM OIL CHEMISTS SOC	41	205	1964	64002
	J AM OIL CHEMISTS SOC	43	157	1966	66003
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SWARBICK J	NATURE	208	780	1965	65028
SZEGLOWSKI Z	TRANS FARADAY SOC	61	1043	1965	65003
TACHIBANA T	ZESZTY NAUK UNIW JAGIEL	211	199	1966	66034
TAMAKI K	ZESZTY NAUK UNIW JAGIEL	211	209	1966	66035
TAMAMUSHI B	J PHYS CHEM	66	363	1962	62024
TARTAR HV	BULL CHEM SOC JAPAN	31	467	1958	58007
TATE JR	REP INST SCI TECH UNIV TOKYO	7	401	1953	53001
TAUBMAN AB	BULL CHEM SOC JAPAN	28	545	1955	55010
TAYLOR FH	BULL CH M S J) 1-	9	733	1956	56009
TAYLOR H	BULL CHEM SOC JAPAN	30	411	1957	57015
THAKKAR AL	BULL CHEM SOC JAPAN	30	542	1957	57016
THIBAULT HG	BULL CHEM SOC JAPAN	31	467	1958	58007
THOMAS IL	J AM CHEM SOC	57	570	1935	35006
TOFIAS A	J AM CHEM SOC	58	322	1936	36006
TOKIWA F	J AM CHEM SOC	60	544	1938	38008
TONG LKJ	J AM CHEM SOC	61	539	1939	39002
TORI K	J AM CHEM SOC	61	549	1939	39007
TRAP HJL	J AM CHEM SOC	61	544	1939	39010
TREBBI GF	J AM CHEM SOC	62	2375	1940	40001
TSIKURINA NN	J AM CHEM SOC	65	686	1943	43001
TUDDENHAM RF	J AM CHEM SOC	65	692	1943	43003
TSIKURINA NN	J AM CHEM SOC	65	698	1943	43004
TUDDENHAM RF	J AM CHEM SOC	68	1490	1946	46003
TSIKURINA NN	J AM CHEM SOC	70	1989	1948	48007
TUDDENHAM RF	J AM CHEM SOC	70	1992	1948	48012
TSIKURINA NN	J AM CHEM SOC	73	5411	1951	51003
TUDDENHAM RF	J PHYS CHEM	59	1190	1955	55006
TSIKURINA NN	J PHYS CHEM	59	1193	1955	55012
TUDDENHAM RF	J PHYS CHEM	59	1195	1955	55013
TSIKURINA NN	J PHYS CHEM	59	1185	1955	55021
TUDDENHAM RF	J COLLOID SCI	14	115	1959	59010
TSIKURINA NN	J COLLOID SCI	17	243	1962	62014
TUDDENHAM RF	PROC ROY SOC	273	84	1963	63014
TSIKURINA NN	TRANS FARADAY SOC	60	986	1964	64012
TUDDENHAM RF	TRANS FARADAY SOC	60	996	1964	64027
TSIKURINA NN	TRANS FARADAY SOC	62	994	1966	66014
TUDDENHAM RF	TRANS FARADAY SOC	62	987	1966	66015
TSIKURINA NN	TRANS FARADAY SOC	62	979	1966	66016
TSIKURINA NN	KHIM TEKHNOL	5	61	1960	60033
TUDDENHAM RF	PHIL TRANS ROY SOC LONDON, SER A	252	225	1960	60011
TSIKURINA NN	TRANS FARADAY SOC	58	1233	1962	62004
TSIKURINA NN	J COLLOID SCI	19	495	1964	64050
TSIKURINA NN	J PHARM SCI	54	1529	1965	65027
TSIKURINA NN	J PHYS CHEM	56	701	1952	52013
TSIKURINA NN	J PHARM PHARMACOL	15	522	1963	63005
TSIKURINA NN	J PHARM PHARMACOL	15	157	1963	63031
TSIKURINA NN	J PHYS CHEM	56	701	1952	52013
TSIKURINA NN	J CHEM SOC JAPAN, PURE CHEM SEC	82	1657	1961	61009
TSIKURINA NN	J CHEM SOC JAPAN, PURE CHEM SEC	82	1512	1961	61010
TSIKURINA NN	J CHEM SOC JAPAN, PURE CHEM SEC	82	1660	1961	61011
TSIKURINA NN	BULL CHEM SOC JAPAN	34	1236	1961	61012
TSIKURINA NN	J CHEM SOC JAPAN, PURE CHEM SEC	83	732	1962	62007
TSIKURINA NN	BULL CHEM SOC JAPAN	35	1737	1962	62012
TSIKURINA NN	BULL CHEM SOC JAPAN	35	240	1962	62022
TSIKURINA NN	BULL CHEM SOC JAPAN	36	222	1963	63003
TSIKURINA NN	BULL CHEM SOC JAPAN	36	1589	1963	63004
TSIKURINA NN	BULL CHEM SOC JAPAN	36	281	1963	63007
TSIKURINA NN	BULL CHEM SOC JAPAN	36	1585	1963	63023
TSIKURINA NN	J CHEM EDUC	40	472	1963	63029
TSIKURINA NN	BULL CHEM SOC JAPAN	37	1837	1964	64024
TSIKURINA NN	J PHYS CHEM	68	2818	1964	64037
TSIKURINA NN	BULL CHEM SOC JAPAN	38	751	1965	65014
TSIKURINA NN	J PHYS CHEM	70	3437	1966	66027
TSIKURINA NN	J PHYS CHEM	69	2357	1965	65030
TUDDENHAM RF	J CHEM SOC JAPAN, PURE CHEM SEC	78	1573	1957	57020
TUDDENHAM RF	KOLLOID-Z	168	132	1960	60009
TUDDENHAM RF	KOLLOID-Z	189	50	1963	63008
TUDDENHAM RF	KOLLOID-Z	188	47	1963	63009
TUDDENHAM RF	KOLLOID-Z	187	44	1963	63010
TUDDENHAM RF	KOLLOID-Z	191	48	1963	63012
TUDDENHAM RF	KOLLOID-Z	194	143	1964	64015
TUDDENHAM RF	KONINKI NED AKAD WETEN. PROC SER B	58	97	1955	55009
TUDDENHAM RF	J PHYS CHEM	67	1987	1963	63017
TUDDENHAM RF	J AM OIL CHEMISTS SOC	41	231	1964	64013
TUDDENHAM RF	J PHYS CHEM	68	3592	1964	64014
TUDDENHAM RF	J COLLOID SCI	20	191	1965	65001
TUDDENHAM RF	KOLLOID ZH	26	76	1964	64051
TUDDENHAM RF	J PHYS CHEM	66	1839	1962	62035

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TYUZO K	BULL CHEM SOC JAPAN	26	177	1953	53006
	BULL CHEM SOC JAPAN	27	259	1954	54008
	BULL CHEM SOC JAPAN	31	117	1958	58026
	KOLLOID-Z	175	40	1961	61025
UBBELOHDE AR	J COLLOID SCI	8	424	1953	53008
ULEVITCH IN	J COLLOID SCI	9	382	1954	54006
V DEN TEMPEL M	PROC INTERN CONGR SURFACE ACTIVITY	3RD	248	1960	60022
V VOORST VA.F	PROC INTERN CONGR SURFACE ACTIVITY	3RD	276	1960	60013
	TRANS FARADAY SOC	56	1078	1960	60014
	PROC INTERN CONGR SURFACE ACTIVITY	3RD	248	1960	60022
	TRANS FARADAY SOC	56	1067	1960	60025
	TRANS FARADAY SOC	57	110	1961	61026
VASSILIADES T	J PHYS CHEM	65	1781	1961	61027
	J PHYS CHEM	65	1774	1961	61028
VEIS A	J COLLOID SCI	15	427	1960	60001
VENABLE RL	J PHYS CHEM	68	3498	1964	64001
VERGNOLLE J	MEM SERV CHIM ETAT (PARIS)	41	363	1956	56010
	PROC INTERN CONGR SURFACE ACTIVITY	2ND	395	1957	57017
VERMA SP	INDIAN J CHEM	3	441	1965	65021
VETTER RJ	J PHYS COLLOID CHEM	51	263	1947	47011
VINOGRAD JR	J AM CHEM SOC	63	670	1941	41005
VOEKS JF	J PHYS CHEM	59	1190	1955	55006
WACHS W	KOLLOID-Z	181	139	1962	62023
WAGNER E	PROC INTERN CONGR SURFACE ACTIVITY	3RD	27	1960	60017
WALKER T	TRANS FARADAY SOC	61	589	1965	65004
WALTON HF	J COLLOID SCI	1	385	1946	46020
WAN LSC	J PHARM SCI	55	1395	1966	66018
WARD AFH	J CHEM SOC		522	1939	39003
	PROC ROY SOC	176	412	1940	40004
WASIK SP	J RES NAT BUR STD A	68	359	1964	64043
WATARI Y	J SCI HIROSHIMA UNIV, SER A-II	28	41	1964	64036
WEATHERBURN AS	CAN J CHEM ENGR	F28	213	1950	50010
WEIL JK	J PHYS CHEM	60	899	1956	56008
	J PHYS CHEM	62	1083	1958	58003
	J AM OIL CHEMISTS SOC	36	241	1959	59004
	J AM OIL CHEMISTS SOC	37	679	1960	60008
	J AM OIL CHEMISTS SOC	39	55	1962	62008
	J AM OIL CHEMISTS SOC	40	538	1963	63013
	J AM OIL CHEMISTS SOC	41	205	1964	64002
	J AM OIL CHEMISTS SOC	43	157	1966	66003
WEINER ND	J PHARM SCI	54	436	1965	65026
WESTWELL AE	J PHYS CHEM	63	1022	1959	59011
	J PHYS CHEM	68	3490	1964	64018
WHEELER OL	J AM CHEM SOC	68	1490	1946	46003
	J AM CHEM SOC	70	1989	1948	48007
WHITE P	J COLLOID SCI	13	584	1958	58005
	TRANS FARADAY SOC	54	1638	1958	58006
	CAN J CHEM	37	2086	1959	59008
	TRANS FARADAY SOC	55	1025	1959	59012
	J PHYS CHEM	64	599	1960	60002
WILDER AG	J PHYS COLLOID CHEM	52	12	1948	48026
WILLIAMS DE	J POLYMER SCI	5	201	1950	50002
WILLIAMS EF	J COLLOID SCI	12	452	1957	57009
WILLIAMS G	ANN REP PROGR CHEM (CH. S. LONDON)	45	51	1948	48003
WILLIAMS RJ	TRANS FARADAY SOC	51	728	1955	55005
WINSLOW L	J PHYS CHEM	58	1124	1954	54012
	Z PHYSIK CHEM (FRANKFURT)	11	165	1957	57021
WINSOR PA	TRANS FARADAY SOC	44	463	1948	48008
	J PHYS CHEM	56	391	1952	52009
WOODBERRY N	J COLLOID SCI	12	452	1957	57009
WOODWARD RJ	J PHARM PHARMACOL	15	422	1963	63015
WRIGHT KA	J AM CHEM SOC	61	539	1939	39002
	J AM CHEM SOC	61	549	1939	39007
	J AM CHEM SOC	61	544	1939	39010
YAMAGUCHI T	BULL CHEM SOC JAPAN	34	237	1961	61008
YAMANAKA T	J PHYS CHEM	63	648	1959	59013
YANG JT	J PHYS CHEM	57	628	1953	53015
YODA O	J CHEM SOC JAPAN, PURE CHEM SEC	77	1236	1956	56020
	BULL CHEM SOC JAPAN	30	760	1957	57033
YOUNG HS	J AM CHEM SOC	71	309	1949	49017
YURZHENKO AI	COLLOID J (USSR)	14	243	1952	52018
	COLLOID J (USSR)	14	311	1952	52019
ZAKHAROVA NN	UKR KHIM ZH	28	611	1962	62038
ZOELLNER M	J PHYS CHEM	65	1804	1961	61001
ZOGRAFI G	J PHARM SCI	54	436	1965	65026
	J PHARM SCI	55	1345	1966	66017
ZUTRAUEN HA	J CHIM PHYS	53	62	1956	56017

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27001	EKWALL P		ACTA ACAD ABOENSIS, MATH PHYS	4	1	1927	YIELDED 5 ENTRIES
27002	JONES E	BURY CR	PHIL MAG	4	841	1927	YIELDED 2 ENTRIES
28001	EKWALL P		KOLLOID-Z	45	291	1928	NO ENTRIES
29001	GRINDLEY J	BURY CR	J CHEM SOC		679	1929	YIELDED 5 ENTRIES
30001	DAVIES DG	BURY CR	J CHEM SOC		2263	1930	YIELDED 1 ENTRIES
32001	EKWALL P		KOLLOID-Z	161	195	1932	YIELDED 5 ENTRIES
33001	EKWALL P		ACTA ACAD ABOENSIS, MATH PHYS	7	3	1933	NO ENTRIES
33002	EKWALL P		ACTA ACAD ABOENSIS, MATH PHYS	7	3	1933	NO ENTRIES
33003	LOTTERMOSER A	PUSCHEL F	KOLLOID-Z		63	175	1933 NO ENTRIES
33004	LOTTERMOSER A	STOLL F	KOLLOID-Z		63	49	1933 NO ENTRIES
34001	MALSCH J	HARTLEY GS	Z PHYSIK CHEM (LEIPZIG)	170	321	1934	YIELDED 1 ENTRIES
35001	MURRAY RC	HARTLEY GS	TRANS FARADAY SOC	31	183	1935	YIELDED 1 ENTRIES
35002	LOTTERMOSER A	GIESE E	KOLLOID-Z	73	276	1935	NO ENTRIES
35003	STEWART A	BUNBURY HM	TRANS FARADAY SOC	31	208	1935	NO ENTRIES
35004	LOTTERMOSER A		TRANS FARADAY SOC	31	200	1935	NO ENTRIES
35005	POWNEY J		TRANS FARADAY SOC	31	1510	1935	NO ENTRIES
35006	REED RM	TARTAR HV	J AM CHEM SOC	57	570	1935	NO ENTRIES
35007	MCBAIN JW	BETZ MD	J AM CHEM SOC	57	1905	1935	NO ENTRIES
35008	BURY CR	PARRY GA	J CHEM SOC		626	1935	YIELDED 2 ENTRIES
36001	HARTLEY GS	COLLIE B	SAMIS CS	TRANS FARADAY SOC	32	795	1936 YIELDED 5 ENTRIES
36002	HARTLEY GS		J AM CHEM SOC	58	2347	1936	YIELDED 8 ENTRIES
36003	EKWALL P		KOLLOID-Z	77	320	1936	NO ENTRIES
36004	ADAM NK		TRANS FARADAY SOC	32	653	1936	NO ENTRIES
36005	MCBAIN JW	SEARLES J	J PHYS CHEM	40	493	1936	NO ENTRIES
36006	REED RM	TARTAR HV	J AM CHEM SOC	58	322	1936	NO ENTRIES
37001	EKWALL P		KOLLOID-Z	80	77	1937	NO ENTRIES
37003	POWNEY J	ADDISON CC	TRANS FARADAY SOC	33	851	1937	NO ENTRIES
37004	LOTTERMOSER A	FROTSCHER H	KOLLOID-BEIH.	45	303	1937	NO ENTRIES
37005	LONG FA	NUTTING GC	HARKINS WD	J AM CHEM SOC	59	2197	1937 NO ENTRIES
38001	HARTLEY GS		J CHEM SOC		1968	1938	YIELDED 4 ENTRIES
38002	EKWALL P		KOLLOID-Z	84	284	1938	NO ENTRIES
38003	HARTLEY GS		NATURE	142	161	1938	NO ENTRIES
38004	SCHMID G	LARSEN EC	Z ELEKTROCHEM	44	651	1938	NO ENTRIES
38005	SAMIS CS	HARTLEY GS	TRANS FARADAY SOC	34	1288	1938	NO ENTRIES
38006	POWNEY J	ADDISON CC	TRANS FARADAY SOC	34	372	1938	YIELDED 7 ENTRIES
38007	ADAM NK	SHUTE HL	TRANS FARADAY SOC	34	758	1938	NO ENTRIES
38008	HOULTON HG	TARTAR HV	J AM CHEM SOC	60	544	1938	NO ENTRIES
39001	HARTLEY GS		TRANS FARADAY SOC	35	1109	1939	NO ENTRIES
39002	TARTAR HV	WRIGHT KA	J AM CHEM SOC	61	539	1939	YIELDED 5 ENTRIES
39003	WARD AFH		J CHEM SOC		522	1939	NO ENTRIES
39004	EKWALL P		FINSKA KEMISTSAMFUNDETS MEDD	1	8	1939	NO ENTRIES
39005	HARTLEY GS		KOLLOID-Z	88	22	1939	NO ENTRIES
39006	STAUFF J		Z PHYSIK CHEM (LEIPZIG)	183	55	1939	YIELDED 5 ENTRIES
39007	WRIGHT KA	ABBOTT AD	SIVERTZ V	J AM CHEM SOC	61	549	1939 YIELDED 12 ENTRIES
39008	HARTLEY GS		TARTAR HV				
39009	HESS K	PHILIPPOFF W	KIESSIG H	J CHEM SOC		1828	1939 NO ENTRIES
39010	WRIGHT KA	TARTAR HV		KOLLOID-Z	88	40	1939 YIELDED 9 ENTRIES
39011	MCBAIN MEL	DYE WB	JOHNSTON SA	J AM CHEM SOC	61	544	1939 NO ENTRIES
40001	TARTAR HV		REITMIER RE	J AM CHEM SOC	61	3210	1939 YIELDED 7 ENTRIES
40002	EKWALL P			KOLLOID-Z	62	2375	1940 NO ENTRIES
40003	EKWALL P			TEK FOREN FINLAND FORH	92	141	1940 NO ENTRIES
40004	WARD AFH			PROC ROY SOC	10	1	1940 YIELDED 4 ENTRIES
40005	NUTTING GC	LONG FA	HARKINS WD	J AM CHEM SOC	176	412	1940 YIELDED 8 ENTRIES
41001	HARTLEY GS			TRANS FARADAY SOC	62	1496	1940 NO ENTRIES
41002	EKWALL P	LINDSTROM RE		MEDD.ABO AKAD.FYS.KEM.IN.	37	130	1941 NO ENTRIES
41003	EKWALL P	LINDSTROM RE		SP NO	97	3	1941 NO ENTRIES
41004	EKWALL P	LINDSTROM RE		KOLLOID-Z	94	71	1941 YIELDED 2 ENTRIES
41005	MCBAIN JW	MERRILL RC JR	VINOGRAD JR	KOLLOID-Z	63	42	1941 YIELDED 1 ENTRIES
41006	NUTTING GC	LONG FA		J AM CHEM SOC	63	670	1941 NO ENTRIES
42001	HOERR CW	RALSTON AW		J AM CHEM SOC	63	84	1941 NO ENTRIES
42002	RALSTON AW	HOERR CW		J AM CHEM SOC	64	2824	1942 NO ENTRIES
42003	HAFFNER FD	PICCIONE GA	ROSENBLUM C	J PHYS CHEM	64	772	1942 YIELDED 7 ENTRIES
42004	EKWALL P			KOLLOID-Z	46	662	1942 YIELDED 8 ENTRIES
42005	RALSTON AW	HOERR CW	HOFFMAN EJ	J AM CHEM SOC	101	135	1942 YIELDED 10 ENTRIES
42006	MERRILL RC JR	MCBAIN JW		J PHYS CHEM	64	97	1942 NO ENTRIES
42007	MCBAIN JW	MERRILL RC JR		IND ENG CHEM	46	10	1942 NO ENTRIES
42008	HOFFMAN EJ	BOYD GE	RALSTON AW	J AM CHEM SOC	34	915	1942 NO ENTRIES
42009	HOFFMAN EJ	BOYD GE	RALSTON AW	J AM CHEM SOC	64	2067	1942 NO ENTRIES
43001	PAQUETTE RG	LINGAFELTER EC	TARTAR HV	J AM CHEM SOC	64	498	1942 NO ENTRIES
43002	HOERR CW	RALSTON AW		J AM CHEM SOC	65	686	1943 YIELDED 8 ENTRIES
43003	SCOTT AB	TARTAR HV		J AM CHEM SOC	65	976	1943 NO ENTRIES
43004	SCOTT AB	TARTAR HV	LINGAFELTER EC	J AM CHEM SOC	65	692	1943 YIELDED 10 ENTRIES
43005	EKWALL P	HARVA O		FINSKA KEMISTSAMFUNDETS MEDD	65	698	1943 YIELDED 2 ENTRIES
43006	MCBAIN JW	BRADY AP		J AM CHEM SOC	25	257	1943 NO ENTRIES
					65	2072	1943 YIELDED 2 ENTRIES

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43007	MCBAIN ML		J PHYS CHEM	47	196	1943	NO ENTRIES	
43008	MCBAIN JW	BOLDUAN OEA	J PHYS CHEM	47	94	1943	NO ENTRIES	
43009	HOERR CW	MCCORKLE MR	RALSTON AW	65	328	1943	NO ENTRIES	
44001	DREGER EE	KEIM GI	MILES GD	36	610	1944	NO ENTRIES	
	SHEDLOVSKY L	ROSS J						
44002	MILES GD	SHEDLOVSKY L	J PHYS CHEM	48	57	1944	NO ENTRIES	
44003	MCBAIN JW	JOHNSON KE	J AM CHEM SOC	66	9	1944	NO ENTRIES	
45001	GONICK E		J AM CHEM SOC	67	1191	1945	NO ENTRIES	
45002	MILES GD		J PHYS CHEM	49	71	1945	NO ENTRIES	
46001	RALSTON AW	HOERR CW	J AM CHEM SOC	68	2460	1946	YIELDED 6 ENTRIES	
46002	CORRIN ML	KLEVENS HB	HARKINS WD	14	216	1946	YIELDED 3 ENTRIES	
46003	LINGAFELTER EC	WHEELER OL	TARTAR HV	68	1490	1946	NO ENTRIES	
46004	CORRIN ML	HARKINS WD	J CHEM PHYS	14	640	1946	YIELDED 11 ENTRIES	
46005	CORRIN ML	HARKINS WD	J CHEM PHYS	14	641	1946	YIELDED 5 ENTRIES	
46006	KOLTHOFF IM	JOHNSON WF	J PHYS CHEM	50	440	1946	YIELDED 5 ENTRIES	
46007	KLEVENS HB		J CHEM PHYS	14	742	1946	YIELDED 8 ENTRIES	
46008	GONICK E	MCBAIN JW	J COLLOID SCI	1	127	1946	YIELDED 1 ENTRIES	
46009	GONICK E		J COLLOID SCI	1	393	1946	NO ENTRIES	
46010	CORRIN ML	KLEVENS HB	HARKINS WD	14	480	1946	YIELDED 7 ENTRIES	
46011	HARKINS WD	STEARN RS	J CHEM PHYS	14	215	1946	NO ENTRIES	
46012	KLEVENS HB		J CHEM PHYS	14	567	1946	YIELDED 2 ENTRIES	
46013	HARKINS WD	MATTOON RW	CORRIN ML	68	220	1946	NO ENTRIES	
46014	HARKINS WD	MATTOON RW	CORRIN ML	1	105	1946	NO ENTRIES	
46015	CORRIN ML	HARKINS WD	J COLLOID SCI	1	469	1946	YIELDED 37 ENTRIES	
46016	GONICK E	MCBAIN JW	REC TRAV CHIM	65	601	1946	YIELDED 1 ENTRIES	
46017	STEARNS RS	HARKINS WD	J CHEM PHYS	14	214	1946	NO ENTRIES	
46018	EVERS EC	GRIEGER PF	KRAUS CA	68	1137	1946	NO ENTRIES	
46019	GONICK E		J AM CHEM SOC	68	177	1946	YIELDED 2 ENTRIES	
46020	WALTON HF	HIEBERT EN	SHOLTES EH	1	385	1946	NO ENTRIES	
46021	ADAM NK	PANKHURST KGA	TRANS FARADAY SOC	42	523	1946	NO ENTRIES	
46022	MCBAIN JW	GREEN AA	J AM CHEM SOC	68	731	1946	NO ENTRIES	
47001	BROWN GL	GRIEGER PF	EVERS EC	J AM CHEM SOC	69	1835	1947	NO ENTRIES
	KRAUS CA							
47002	RALSTON AW	HOERR CW	J AM CHEM SOC	69	883	1947	NO ENTRIES	
47003	RALSTON AW	EGGENBERGER DN	HARWOOD HJ	J AM CHEM SOC	69	2095	1947	YIELDED 7 ENTRIES
	DU BROW PL							
47004	KLEVENS HB		J PHYS COLLOID CHEM	51	114	1947	YIELDED 27 ENTRIES	
47005	KLEVENS HB		J COLLOID SCI	2	301	1947	YIELDED 19 ENTRIES	
47006	HARKINS WD	CORRIN ML	J AM CHEM SOC	69	679	1947	YIELDED 14 ENTRIES	
47007	GONICK E	MCBAIN JW	J AM CHEM SOC	69	334	1947	YIELDED 2 ENTRIES	
47008	HARKINS WD	MATTOON RW	MITTELMANN R	15	763	1947	NO ENTRIES	
47009	HARKINS WD		J CHEM PHYS	69	1428	1947	NO ENTRIES	
47010	CORRIN ML	HARKINS WD	J AM CHEM SOC	69	683	1947	YIELDED 106 ENTRIES	
47011	VETTER RJ		J PHYS COLLOID CHEM	51	263	1947	YIELDED 2 ENTRIES	
47012	STEARNS RS	OPPENHEIMER H	SIMON E	J CHEM PHYS	15	496	1947	NO ENTRIES
	HARKINS WD							
47013	CARR CW	JOHNSON WF	KOLTHOFF IM	J PHYS COLLOID CHEM	51	636	1947	YIELDED 6 ENTRIES
48001	DAINTON FS		ANN REP PROGR CHEM (CH. S. LONDON)	45	5	1948	NO ENTRIES	
48002	HARTLEY GS		ANN REP PROGR CHEM (CH. S. LONDON)	45	33	1948	NO ENTRIES	
48003	WILLIAMS G	SINGER K	ANN REP PROGR CHEM (CH. S. LONDON)	45	51	1948	NO ENTRIES	
48004	BRADY AP	HUFF H	J COLLOID SCI	3	511	1948	YIELDED 10 ENTRIES	
48005	KLEVENS HB		J PHYS COLLOID CHEM	52	130	1948	YIELDED 68 ENTRIES	
48006	DEBYE P		J COLLOID SCI	3	407	1948	NO ENTRIES	
48007	NEFF LL	WHEELER OL	TARTAR HV	J AM CHEM SOC	70	1989	1948	YIELDED 1 ENTRIES
	LINGAFELTER EC							
48008	WINSOR PA		TRANS FARADAY SOC	44	463	1948	YIELDED 23 ENTRIES	
48009	CUSHMAN A	BRADY AP	MCBAIN JW	J COLLOID SCI	3	425	1948	YIELDED 5 ENTRIES
48010	GRIEGER PF	KRAUS CA	J AM CHEM SOC	70	3803	1948	YIELDED 7 ENTRIES	
48011	FINEMAN MN	MCBAIN JW	J PHYS COLLOID CHEM	52	881	1948	YIELDED 10 ENTRIES	
48012	GREGORY NW	TARTAR HV	J AM CHEM SOC	70	1992	1948	YIELDED 2 ENTRIES	
48013	CORRIN ML		J COLLOID SCI	3	333	1948	NO ENTRIES	
48014	RALSTON AW	EGGENBERGER DN	DU BROW PL	J AM CHEM SOC	70	977	1948	YIELDED 1 ENTRIES
48015	MCBAIN JW	MCHAN H	J AM CHEM SOC	70	3838	1948	NO ENTRIES	
48016	KOLTHOFF IM	STRICKS W	J PHYS COLLOID CHEM	52	915	1948	YIELDED 19 ENTRIES	
48017	ARKIN L	SINGLETERRY CR	J AM CHEM SOC	70	3965	1948	NO ENTRIES	
48018	EXNER ML		NATURWISSENSCHAFTEN	35	344	1948	YIELDED 1 ENTRIES	
48019	RALSTON AW	EGGENBERGER DN	J AM CHEM SOC	70	980	1948	NO ENTRIES	
48020	RALSTON AW	EGGENBERGER DN	J AM CHEM SOC	70	2918	1948	YIELDED 9 ENTRIES	
48021	RALSTON AW	EGGENBERGER DN	J AM CHEM SOC	70	983	1948	YIELDED 4 ENTRIES	
48022	BRADY AP		J PHYS COLLOID CHEM	3	57	1948	YIELDED 2 ENTRIES	
48023	RALSTON AW	EGGENBERGER DN	J PHYS COLLOID CHEM	52	1494	1948	YIELDED 9 ENTRIES	
48024	MERRILL RC	GETTY R	J PHYS COLLOID CHEM	52	774	1948	YIELDED 76 ENTRIES	
48025	KOLTHOFF IM	JOHNSON WF	J PHYS COLLOID CHEM	52	22	1948	YIELDED 6 ENTRIES	
48026	MCBAIN JW	WILDER AG	MERRILL RC JR	J PHYS COLLOID CHEM	52	12	1948	NO ENTRIES
48027	RALSTON AW	EGGENBERGER DN	J AM CHEM SOC	70	436	1948	YIELDED 6 ENTRIES	
48028	EVERS EC	KRAUS CA	J AM CHEM SOC	70	3049	1948	YIELDED 17 ENTRIES	
49001	DEBYE P		ANN N Y ACAD SCI	51	575	1949	YIELDED 8 ENTRIES	
49002	DEBYE P		J PHYS COLLOID CHEM	53	1	1949	NO ENTRIES	
49003	KLEVENS HB		J AM OIL CHEMISTS SOC	26	456	1949	NO ENTRIES	
49004	SHUCK GR	LINGAFELTER EC	J AM CHEM SOC	71	1325	1949	YIELDED 3 ENTRIES	

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Reference No.	Authors		Journal	Vol.	Pg.	Year	No. of Entries from Reference	
49005	KOLTHOFF IM	STRICKS W	J PHYS COLLOID CHEM	53	424	1949	YIELDED 49 ENTRIES	
49006	HARKINS WD	MITTELMANN R CORRIN ML	J PHYS COLLOID CHEM	53	1350	1949	YIELDED 41 ENTRIES	
49007	HARKINS WD	MITTELMANN R	J COLLOID SCI	4	367	1949	NO ENTRIES	
49008	RALSTON AW	EGGENBERGER DN BROOME FK	J AM CHEM SOC	71	2145	1949	YIELDED 45 ENTRIES	
49009	RALSTON AW	EGGENBERGER DN HARWOOD HJ	J AM CHEM SOC	71	672	1949	YIELDED 1 ENTRIES	
49010	HARTLEY GS		NATURE	163	767	1949	NO ENTRIES	
49011	HARKINS WD	OPPENHEIMER H	J AM CHEM SOC	71	808	1949	NO ENTRIES	
49012	ARKIN L	SINGLETERRY CR	J COLLOID SCI	4	537	1949	NO ENTRIES	
49013	RALSTON AW	BROOME FK	HARWOOD HJ	J AM CHEM SOC	71	671	1949	YIELDED 4 ENTRIES
49014	BROWN GL	GRIEGER PF	KRAUS CA	J AM CHEM SOC	71	95	1949	YIELDED 22 ENTRIES
49015	ADCOCK WA	COLE RH		J AM CHEM SOC	71	2835	1949	NO ENTRIES
49017	YOUNG HS	GRIEGER PF	KRAUS CA	J AM CHEM SOC	71	309	1949	YIELDED 15 ENTRIES
49018	GRIEGER PF	KRAUS CA	J AM CHEM SOC	71	1455	1949	YIELDED 2 ENTRIES	
50001	DEBYE P		AN. R. SOC ESPAN FIS QUIM (MADRID)	46	335	1950	NO ENTRIES	
50002	CARR CW	KOLTHOFF IM	MEEHAN EJ	J POLYMER SCI	5	201	1950	NO ENTRIES
	WILLIAMS DE							
50003	KLEVENS HB		J PHYS COLLOID CHEM	54	1012	1950	YIELDED 30 ENTRIES	
50004	KLEVENS HB		J AM CHEM SOC	72	3780	1950	NO ENTRIES	
50005	CARR CW	KOLTHOFF IM	MEEHAN EJ	J POLYMER SCI	5	191	1950	NO ENTRIES
	STERNBERG RJ							
50006	KLEVENS HB		ANAL CHEM	22	1141	1950	NO ENTRIES	
50007	KLEVENS HB		CHEM REV	47	1	1950	NO ENTRIES	
50008	HERZFIELD SH	CORRIN ML	HARKINS WD	J PHYS COLLOID CHEM	54	271	1950	YIELDED 98 ENTRIES
50009	GOTTE E			KOLLOID-Z	117	42	1950	NO ENTRIES
50010	WEATHERBURN AS	ROSE GRF		CAN J CHEM ENGR	F28	213	1950	NO ENTRIES
50011	HARKINS WD			SCI MONTHLY	70	220	1950	NO ENTRIES
50012	COLICHMAN EL			J AM CHEM SOC	72	4036	1950	YIELDED 38 ENTRIES
50013	KUHN DW	KRAUS CA		J AM CHEM SOC	72	3676	1950	NO ENTRIES
51001	DEBYE P	ANACKER EW		J PHYS COLLOID CHEM	55	644	1951	YIELDED 1 ENTRIES
51002	KOLTHOFF IM	JOHNSON WF		J AM CHEM SOC	73	4563	1951	NO ENTRIES
51003	LELONG ALM	TARTAR HV	LINGAFELTER EC	J AM CHEM SOC	73	5411	1951	YIELDED 10 ENTRIES
51004	HUFF H	MCBAIN JW	BRADY AP	J PHYS COLLOID CHEM	55	311	1951	YIELDED 29 ENTRIES
51005	LANGE H			KOLLOID-Z	121	66	1951	YIELDED 94 ENTRIES
51006	EGGENBERGER DN	HARWOOD HJ		J AM CHEM SOC	73	3353	1951	YIELDED 8 ENTRIES
51007	SINGLETERRY CR	ARKIN L		J AM CHEM SOC	73	4574	1951	NO ENTRIES
51008	COLICHMAN EL			J AM CHEM SOC	73	3385	1951	YIELDED 15 ENTRIES
51009	MCDOWELL MJ	KRAUS CA		J AM CHEM SOC	73	2173	1951	YIELDED 5 ENTRIES
51010	HARKINS WD	KRIZEK H	CORRIN ML	J COLLOID SCI	6	576	1951	YIELDED 6 ENTRIES
51011	HENNE AL	FOX CJ		J AM CHEM SOC	73	2323	1951	NO ENTRIES
51012	SCOTT R	BOLAM TR		NATURE	167	195	1951	NO ENTRIES
51013	DAGGETT HM JR	BAIR EJ	KRAUS CA	J AM CHEM SOC	73	799	1951	NO ENTRIES
51014	BAIR EJ	KRAUS CA		J AM CHEM SOC	73	1129	1951	NO ENTRIES
51016	MCDOWELL MJ	KRAUS CA		J AM CHEM SOC	73	2170	1951	NO ENTRIES
51017	SHISHIDO S			BULL CHEM SOC JAPAN	24	41	1951	NO ENTRIES
51018	NAKAGAKI M			J CHEM SOC JAPAN, PURE CHEM SEC	72	113	1951	NO ENTRIES
52001	CELLA JA	EGGENBERGER DN	NOEL DR	J AM CHEM SOC	74	2061	1952	YIELDED 14 ENTRIES
	HARRIMAN LA	HARWOOD HJ						
52002	KOLTHOFF IM	JOHNSON WF		J AM CHEM SOC	74	20	1952	NO ENTRIES
52003	JOHNSON WF	KOLTHOFF IM		J AM CHEM SOC	74	22	1952	NO ENTRIES
52004	KLEVENS HB			MEM SERV CHIM ETAT (PARIS)	37	13	1952	NO ENTRIES
52005	KLEVENS HB			J AM CHEM SOC	74	4624	1952	NO ENTRIES
52006	EKWALL P	SMEDS K		ACTA CHEM SCAND	6	441	1952	NO ENTRIES
52007	EKWALL P	HASAN A	DANIELSSON I	ACTA CHEM SCAND	6	440	1952	NO ENTRIES
52008	KLEVENS HB			KOLLOID-Z	128	61	1952	NO ENTRIES
52009	WINSOR PA			J PHYS CHEM	56	391	1952	NO ENTRIES
52011	BURY CR	BROWNING J		TRANS FARADAY SOC	48	209	1952	YIELDED 3 ENTRIES
52013	BROWN AS	ROBINSON RU	SIROIS EH	J PHYS CHEM	56	701	1952	YIELDED 3 ENTRIES
52014	MEADER AL	THIBAULT HG	MCNEILL W	FRIES BA				
				IND ENG CHEM	44	1636	1952	NO ENTRIES
52015	HERZFIELD SH			J PHYS CHEM	56	953	1952	YIELDED 20 ENTRIES
52016	RAISON M			COMPT REND	235	1129	1952	YIELDED 10 ENTRIES
52017	HERZFIELD SH			J PHYS CHEM	56	959	1952	YIELDED 108 ENTRIES
52018	YURZHENKO AI	KUCHER RV		COLLOID J (USSR)	14	243	1952	YIELDED 3 ENTRIES
52019	YURZHENKO AI	KUCHER RV		COLLOID J (USSR)	14	311	1952	NO ENTRIES
53001	TAMAMUSHI B	NAKADATE S		REP INST SCI TECH UNIV TOKYO	7	401	1953	YIELDED 2 ENTRIES
53002	ANACKER EW			J COLLOID SCI	8	402	1953	YIELDED 5 ENTRIES
53003	ARRINGTON CH	PATTERSON GD		J PHYS CHEM	57	247	1953	YIELDED 13 ENTRIES
53004	FLOCKHART BD	GRAHAM H		J COLLOID SCI	8	105	1953	YIELDED 5 ENTRIES
53005	LANGE H			KOLLOID-Z	131	96	1953	YIELDED 21 ENTRIES
53006	SATA N	TYUZO K		BULL CHEM SOC JAPAN	26	177	1953	YIELDED 11 ENTRIES
53007	ROSS S	KWARTLER CE	BAILEY JH	J COLLOID SCI	8	385	1953	YIELDED 19 ENTRIES
53008	FLOCKHART BD	UBBELOHDE AR		J COLLOID SCI	8	424	1953	YIELDED 29 ENTRIES
53009	KUSHNER LM	HUBBARD WD		J PHYS CHEM	57	808	1953	NO ENTRIES
53010	KLEVENS HB			J AM OIL CHEMISTS SOC	30	74	1953	YIELDED 31 ENTRIES
53011	KRAUS CA			PROC NAT ACAD SCI U S	39	1213	1953	NO ENTRIES
53012	GODDARD ED	HARVA O	JONES TG	TRANS FARADAY SOC	49	980	1953	YIELDED 61 ENTRIES
53013	LAL H			J COLLOID SCI	8	414	1953	NO ENTRIES
53014	SCHOLBERG HM	GUENTHNER RA	COON RI	J PHYS CHEM	57	923	1953	NO ENTRIES
53015	YANG JT	FOSTER JF		J PHYS CHEM	57	628	1953	YIELDED 2 ENTRIES

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Reference No.	Authors	Journal	Vol.	Pg.	Year	No. of Entries from Reference	
54001	PILPEL N	J COLLOID SCI	9	285	1954	NO ENTRIES	
54002	KUSHNER LM	HUBBARD WD	J PHYS CHEM	58	1163	1954 NO ENTRIES	
54003	SHINODA K	J PHYS CHEM	58	1136	1954 YIELDED 57 ENTRIES		
54004	KLEVENS HB	RAISON M	PROC INTERN CONGR SURFACE ACTIVITY	1ST	1	1954 YIELDED 25 ENTRIES	
54005	SHINODA K	J PHYS CHEM	58	541	1954 YIELDED 139 ENTRIES		
54006	MARON SH	ELDER ME	J COLLOID SCI	9	382	1954 YIELDED 10 ENTRIES	
54007	GOTTE E	ULEVITCH IN	FETTE, SEIFEN, ANSTRICHMI	56	583	1954 NO ENTRIES	
54008	OKUYAMA H	TYUZYO K	BULL CHEM SOC JAPAN	27	259	1954 YIELDED 3 ENTRIES	
54009	HONIG JG	SINGLETERRY CR	J PHYS CHEM	58	201	1954 NO ENTRIES	
54010	KLEVENS HB	RAISON M	J CHIM PHYS	51	1	1954 YIELDED 25 ENTRIES	
54011	COHEN I	HISKEY CF	OSTER G	J COLLOID SCI	9	243	1954 NO ENTRIES
54012	HUTCHINSON E	MANCHESTER KE	WINSLOW L	J PHYS CHEM	58	1124	1954 NO ENTRIES
54013	ROE CP	BRASS PD	J AM CHEM SOC	76	4703	1954 YIELDED 4 ENTRIES	
54014	HUBBARD HM	REYNOLDS CA	J AM CHEM SOC	76	4300	1954 NO ENTRIES	
54015	HIGUCHI T	LACH JL	J AM PHARM ASSOC	43	465	1954 NO ENTRIES	
55001	HERMANS JJ		KONINKI NED AKAD WETEN. PROC SER B	58	91	1955 NO ENTRIES	
55002	PRINS W	HERMANS JJ	J PHYS CHEM	59	576	1955 NO ENTRIES	
55003	KUSHNER LM	HUBBARD WD	J COLLOID SCI	10	428	1955 YIELDED 6 ENTRIES	
55004	SHINODA K		BULL FAC ENG, YOKOHAMA NAT UNIV	4	77	1955 YIELDED 167 ENTRIES	
55005	WILLIAMS RJ	PHILLIPS JN	TRANS FARADAY SOC	51	728	1955 YIELDED 17 ENTRIES	
55006	VOEKS JF	TARTAR HV	J PHYS CHEM	59	1190	1955 YIELDED 2 ENTRIES	
55007	SHINODA K		J PHYS CHEM	59	432	1955 YIELDED 39 ENTRIES	
55008	LIN W		BULL CHEM SOC JAPAN	28	227	1955 YIELDED 20 ENTRIES	
55009	TRAP HJL	HERMANS JJ	KONINKI NED AKAD WETEN. PROC SER B	58	97	1955 YIELDED 34 ENTRIES	
55010	SHIRAI M	TAMAMUSHI B	BULL CHEM SOC JAPAN	28	545	1955 NO ENTRIES	
55011	GODDARD ED	JONES TG	RES CORRESPONDENCE	8	1	1955 NO ENTRIES	
55012	ABBOTT AD	TARTAR HV	J PHYS CHEM	59	1193	1955 NO ENTRIES	
55013	TARTAR HV		J PHYS CHEM	59	1195	1955 NO ENTRIES	
55014	PHILLIPS JN	MYSELS KJ	J PHYS CHEM	59	325	1955 NO ENTRIES	
55015	MUKERJEE P	MYSELS KJ	J AM CHEM SOC	77	2937	1955 YIELDED 8 ENTRIES	
55016	MYSELS KJ		J COLLOID SCI	10	507	1955 NO ENTRIES	
55017	KLEVENS HB		NATURE	176	879	1955 NO ENTRIES	
55018	GODDARD ED	HIGHAM EH	STEWART JC	RES CORRESPONDENCE	7	1	1955 YIELDED 2 ENTRIES
55019	KITAHARA A		BULL CHEM SOC JAPAN	28	234	1955 NO ENTRIES	
55020	HSIAO L	DUNNING HN	J PHYS CHEM	59	362	1955 NO ENTRIES	
55021	TARTAR HV	LELONG ALM	J PHYS CHEM	59	1185	1955 YIELDED 11 ENTRIES	
55022	FULLER GW		J COLLOID SCI	10	403	1955 NO ENTRIES	
55023	MCBAIN MEL		J COLLOID SCI	10	223	1955 NO ENTRIES	
55024	PHILLIPS JN		TRANS FARADAY SOC	51	561	1955 NO ENTRIES	
55025	ERIKSON JA	LINGAFELTER EC	J COLLOID SCI	10	71	1955 NO ENTRIES	
55026	GRIESS W		FETTE, SEIFEN, ANSTRICHMI	57	236	1955 NO ENTRIES	
55027	GRIESS W		FETTE, SEIFEN, ANSTRICHMI	57	168	1955 NO ENTRIES	
55028	GRIESS W		FETTE, SEIFEN, ANSTRICHMI	57	24	1955 YIELDED 15 ENTRIES	
56001	KLEVENS HB	CARR CW	J PHYS CHEM	60	1245	1956 YIELDED 25 ENTRIES	
56002	DORST W	PRINS W	HERMANS JJ	KONINKI NED AKAD WETEN. PROC SER B	59	190	1956 YIELDED 20 ENTRIES
56003	SHINODA K		J PHYS CHEM	60	1439	1956 YIELDED 5 ENTRIES	
56004	PILPEL N		J PHYS CHEM	60	779	1956 NO ENTRIES	
56005	LUDLUM DB		J PHYS CHEM	60	1240	1956 YIELDED 6 ENTRIES	
56006	EVANS HC		J CHEM SOC	579		1956 YIELDED 32 ENTRIES	
56007	PRINS W	HERMANS JJ	KONINKI NED AKAD WETEN. PROC SER B	59	162	1956 NO ENTRIES	
56008	WEIL JK	STIRTON AJ	J PHYS CHEM	60	899	1956 YIELDED 31 ENTRIES	
56009	SHIRAI M	TAMAMUSHI B	BULL CHEM SOC JAPAN	29	733	1956 NO ENTRIES	
56010	KLEVENS HB	VERGNOLLE J	MEM SERV CHIM ETAT (PARIS)	41	363	1956 YIELDED 5 ENTRIES	
56011	PRINS W	HERMANS JJ	KONINKI NED AKAD WETEN. PROC SER B	59	298	1956 YIELDED 13 ENTRIES	
56012	HONIG JG	SINGLETERRY CR	J PHYS CHEM	60	1108	1956 NO ENTRIES	
56013	HONIG JG	SINGLETERRY CR	J PHYS CHEM	60	1114	1956 NO ENTRIES	
56014	HSIAO L	DUNNING HN	LORENZ PB	J PHYS CHEM	60	657	1956 YIELDED 25 ENTRIES
56015	KITAHARA A		BULL CHEM SOC JAPAN	29	15	1956 NO ENTRIES	
56016	FAVA A	EYRING H	J PHYS CHEM	60	890	1956 YIELDED 3 ENTRIES	
56017	ZUTRAUEN HA		J CHIM PHYS	53	62	1956 NO ENTRIES	
56018	HARVA O		REC TRAV CHIM	75	112	1956 YIELDED 43 ENTRIES	
56019	ADDISON CC	FURMIDGE CGL	J CHEM SOC	3229		1956 YIELDED 13 ENTRIES	
56020	MEGURO K	KONDO T	YODA O	J CHEM SOC JAPAN, PURE CHEM SEC	77	1236	1956 YIELDED 10 ENTRIES
57001	MATIJEVIC E	PETHICA BA	CROAT CHEM ACTA	29	431	1957 NO ENTRIES	
57002	GODDARD ED	HOEVE CAJ	J PHYS CHEM	61	593	1957 NO ENTRIES	
57003	KUSHNER LM	HUBBARD WD	DOAN AS	J PHYS CHEM	61	371	1957 NO ENTRIES
57004	HOYER HW	GREENFIELD A	J PHYS CHEM	61	818	1957 YIELDED 5 ENTRIES	
57005	LIN W		J CHINESE CHEM SOC	4	28	1957 YIELDED 2 ENTRIES	
57006	KUSHNER LM	HUBBARD WD	PARKER RA	J RES NAT BUR STD A	59	113	1957 YIELDED 19 ENTRIES
57007	CARRINGTON RAG	EVANS HC	J CHEM SOC		1701	1957 NO ENTRIES	
57008	LIN W		J CHINESE CHEM SOC	4	21	1957 NO ENTRIES	
57009	WILLIAMS EF	WOODBERRY N	DIXON JK	J COLLOID SCI	12	452	1957 YIELDED 6 ENTRIES
57010	ROSS S	HUDSON JB		J COLLOID SCI	12	523	1957 YIELDED 1 ENTRIES
57011	GODDARD ED	BENSON GC	CAN J CHEM	35	986	1957 YIELDED 30 ENTRIES	
57012	GERSHMAN JW		J PHYS CHEM	61	581	1957 YIELDED 17 ENTRIES	
57013	FLOCKHART BD		J COLLOID SCI	12	557	1957 YIELDED 33 ENTRIES	
57014	SCHICK MJ	FOWKES FM	J PHYS CHEM	61	1062	1957 YIELDED 57 ENTRIES	
57015	SHIRAI M	TAMAMUSHI B	BULL CHEM SOC JAPAN	30	411	1957 NO ENTRIES	
57016	SHIRAI M	TAMAMUSHI B	BULL CHEM SOC JAPAN	30	542	1957 YIELDED 1 ENTRIES	
57017	KLEVENS HB	VERGNOLLE J	PROC INTERN CONGR SURFACE ACTIVITY	2ND	395	1957 YIELDED 14 ENTRIES	

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Reference No.	Authors		Journal	Vol.	Pg.	Year	No. of Entries from Reference	
57018	NAKAGAWA T	KURIYAMA K	J CHEM SOC JAPAN, PURE CHEM SEC	78	1568	1957	NO ENTRIES	
57019	NAKAGAWA T	INOUE H	J CHEM SOC JAPAN, PURE CHEM SEC	78	636	1957	NO ENTRIES	
57020	NAKAGAWA T	KURIYAMA K	TORI K	J CHEM SOC JAPAN, PURE CHEM SEC	78	1573	1957	YIELDED 22 ENTRIES
57021	HUTCHINSON E	WINSLOW L	Z PHYSIK CHEM (FRANKFURT)	11	165	1957	YIELDED 8 ENTRIES	
57022	LANGE H	KLING W	PROC INTERN CONGR SURFACE ACTIVITY	2ND	295	1957	YIELDED 15 ENTRIES	
57023	SASAKI H	SAITO S	OKUYAMA H	BULL CHEM SOC JAPAN	30	186	1957	NO ENTRIES
57024	OSIPOW L	SNELL FD	HICKSON J	PROC INTERN CONGR SURFACE ACTIVITY	2ND		1957	YIELDED 2 ENTRIES
57025	MIURA M	MATSUMOTO T		J SCI HIROSHIMA UNIV, SER A-II	21	51	1957	YIELDED 5 ENTRIES
57026	KAUFMAN S	SINGLETERRY CR	J COLLOID SCI	12	465	1957	NO ENTRIES	
57027	KITAHARA A		J COLLOID SCI	12	342	1957	NO ENTRIES	
57028	KITAHARA A		BULL CHEM SOC JAPAN	30	586	1957	NO ENTRIES	
57029	KITAHARA A		BULL CHEM SOC JAPAN	31	288	1957	NO ENTRIES	
57030	SASAKI H		BULL CHEM SOC JAPAN	30	326	1957	NO ENTRIES	
57031	ROSS S	BRAMFITT TH	J PHYS CHEM	61	1261	1957	YIELDED 18 ENTRIES	
57032	MEGURO K	KONDO T	OHBA N	BULL CHEM SOC JAPAN	30	905	1957	NO ENTRIES
57033	MEGURO K	KONDO T	OHBA N	BULL CHEM SOC JAPAN	30	760	1957	NO ENTRIES
58001	MILLER ML	DIXON JK	J COLLOID SCI	13	411	1958	YIELDED 7 ENTRIES	
58002	HARRIS JC		SOAP CHEM SPECIALTIES	1958		1958	NO ENTRIES	
58003	WEIL JK	BISTLINE RG	STIRTON AJ	J PHYS CHEM	62	1083	1958	YIELDED 20 ENTRIES
58004	HARRIS JC		J AM OIL CHEMISTS SOC	35	670	1958	YIELDED 1 ENTRIES	
58005	WHITE P	BENSON GC	J COLLOID SCI	13	584	1958	NO ENTRIES	
58006	WHITE P	MOULE D	BENSON GC	TRANS FARADAY SOC	54	1638	1958	NO ENTRIES
58007	TAMAMUSHI B	SHIRAI M	TAMAKI K	BULL CHEM SOC JAPAN	31	467	1958	YIELDED 10 ENTRIES
58008	GINN ME	HARRIS JC	J PHYS CHEM	62	1554	1958	YIELDED 18 ENTRIES	
58009	ANACKER EW		J PHYS CHEM	62	41	1958	YIELDED 7 ENTRIES	
58010	HARRIS JC		J AM OIL CHEMISTS SOC	35	428	1958	NO ENTRIES	
58011	KLEVENS HB		KOLLOID-Z	158	53	1958,	YIELDED 20 ENTRIES	
58012	HAYDON DA	PHILLIPS JN	TRANS FARADAY SOC	54	698	1958	YIELDED 2 ENTRIES	
58013	MUKERJEE P	MYSELS KJ	DULIN CI	J PHYS CHEM	62	1390	1958	NO ENTRIES
58014	MUKERJEE P		J PHYS CHEM	62	1397	1958	NO ENTRIES	
58015	MUKERJEE P	MYSELS KJ	J PHYS CHEM	62	1400	1958	NO ENTRIES	
58016	MUKERJEE P		J PHYS CHEM	62	1404	1958	NO ENTRIES	
58017	NAKAGAWA T		SHINOGI KENKYUSHO NEMPO	8	805	1958	YIELDED 9 ENTRIES	
58018	NAKAGAWA T	INOUE H	KURIYAMA K	J CHEM SOC JAPAN, PURE CHEM SEC	79	348	1958	NO ENTRIES
58019	NAKAGAWA T	INOUE H	KURIYAMA K	J CHEM SOC JAPAN, PURE CHEM SEC	79	345	1958	NO ENTRIES
58020	NAKAGAWA T	YOYAMA T		TRANS FARADAY SOC	54	587	1958	YIELDED 9 ENTRIES
58021	KASHIWAGI KM	PETHICA BA	J COLLOID SCI	13	618	1958	YIELDED 2 ENTRIES	
58022	KITAHARA A		BULL CHEM SOC JAPAN	31	653	1958	NO ENTRIES	
58023	MIURA M	ARISHI S	J SCI HIROSHIMA UNIV, SER A-II	22	57	1958	YIELDED 9 ENTRIES	
58024	BASCOM WD	SINGLETERRY CR	J COLLOID SCI	13	569	1958	NO ENTRIES	
58025	KAUFMAN S	SINGLETERRY CR	J PHYS CHEM	62	1257	1958	NO ENTRIES	
58026	TYUZO K		BULL CHEM SOC JAPAN	31	117	1958	NO ENTRIES	
58027	NASH T		CHEM IND (LONDON)		590	1958	NO ENTRIES	
58028	NAKAGAWA T	INOUE H	KAZUO T	J CHEM SOC JAPAN, PURE CHEM SEC	79	48	1958	YIELDED 8 ENTRIES
58029	SAWYER WM	FOWKES FM	J PHYS CHEM	62	159	1958	NO ENTRIES	
58030	MARKAN AL	KRIUENTSOV WI	ZAVODSKAYA LAB	24	158	1958	NO ENTRIES	
58031	FEW A	GILBY A	OTTEWILL RH	J CHEM SOC	1712	1958	YIELDED 1 ENTRIES	
59001	HARROLDE SP		J PHYS CHEM	63	317	1959	YIELDED 3 ENTRIES	
59002	MYSELS KJ	PRINCEN LH	J PHYS CHEM	63	1696	1959	YIELDED 4 ENTRIES	
59003	PRINCEN LH	MYSELS KJ	J PHYS CHEM	63	1781	1959	NO ENTRIES	
59004	WEIL JK	STIRTON AJ	BISTLINE RG	J AM OIL CHEMISTS SOC	36	241	1959	YIELDED 19 ENTRIES
59005	MAURER EW		J COLLOID SCI	14	519	1959	YIELDED 1 ENTRIES	
59006	BECHER P	CLIFTON NK	J PHYS CHEM	63	1675	1959	YIELDED 22 ENTRIES	
59007	MOULE D	BENSON GC	CAN J CHEM	37	2083	1959	YIELDED 4 ENTRIES	
59008	MOULE D	WHITE P	BENSON GC	CAN J CHEM	37	2086	1959	YIELDED 2 ENTRIES
59009	GINN ME	KINNEY FB	HARRIS JC	J AM OIL CHEMISTS SOC	36	332	1959	YIELDED 23 ENTRIES
59010	TARTAR HV		J COLLOID SCI	14	115	1959	YIELDED 2 ENTRIES	
59011	WESTWELL AE	ANACKER EW	J PHYS CHEM	63	1022	1959	NO ENTRIES	
59012	WHITE P	BENSON GC	TRANS FARADAY SOC	55	1025	1959	YIELDED 9 ENTRIES	
59013	SHINODA K	YAMANAKA T	KINOSHITA K	J PHYS CHEM	63	648	1959	YIELDED 3 ENTRIES
59014	ASCOLI F	BOTRE C	CRESCENZI V	NATURE	184	1482	1959	NO ENTRIES
59015	MELE A	CRESCENZI VL	LIQUORI AM	TRANS FARADAY SOC	55	1975	1959	YIELDED 2 ENTRIES
59016	BOTRE C	CRESCENZI VL	MELE A	J PHYS CHEM	63	650	1959	YIELDED 6 ENTRIES
59017	KASHIWAGI M	EZAKI H		BULL CHEM SOC JAPAN	32	624	1959	YIELDED 4 ENTRIES
59018	HUTCHINSON E		Z PHYSIK CHEM (FRANKFURT)	21	38	1959	YIELDED 6 ENTRIES	
59019	FEURSTENAU DW	MODI HJ	J ELECTROCHEM SOC	106	336	1959	NO ENTRIES	
59020	ROSS S	OLIVIER JP	J PHYS CHEM	63	1671	1959	YIELDED 7 ENTRIES	
59021	BASCOM WD	KAUFMAN S	SINGLETERRY CR	WORLD PETROL CONGR, PROC	V 6	18	1959	NO ENTRIES
59022	KOLBEL H	KUHN P		ANGEW CHEM	71	211	1959	NO ENTRIES
59023	HOI LAHAN JR	CADY GH		J PHYS CHEM	63	757	1959	YIELDED 1 ENTRIES
59024	MEGURO K	KONDO T		J CHEM SOC JAPAN, PURE CHEM SEC	80	818	1959	YIELDED 11 ENTRIES
59025	MEGURO K	KONDO T		J CHEM SOC JAPAN, PURE CHEM SEC	80	821	1959	NO ENTRIES
59026	MEGURO K	KONDO T		J CHEM SOC JAPAN, PURE CHEM SEC	80	823	1959	YIELDED 9 ENTRIES
60001	VEIS A	HOERR CW		J COLLOID SCI	15	427	1960	YIELDED 2 ENTRIES

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Reference No.	Authors		Journal	Vol.	Pg.	Year	No. of Entries from Reference	
60002	WHITE P	BENSON GC	J PHYS CHEM	64	599	1960	YIELDED 4 ENTRIES	
60003	BECHER P		J PHYS CHEM	64	1221	1960	NO ENTRIES	
60004	HARROLD SP		J COLLOID SCI	15	280	1960	YIELDED 2 ENTRIES	
60005	CARTAN F	ANACKER EW	J CHEM EDUC	37	36	1960	YIELDED 2 ENTRIES	
60006	NAKAGAWA T	KURIYAMA K	INOUE H	J COLLOID SCI	15	268	1960	YIELDED 20 ENTRIES
60007	NAKAGAWA T		PROC INTERN CONGR SURFACE ACTIVITY	3RD	636	1960	NO ENTRIES	
60008	WEIL JK	STIRTON AJ	BISTLINE RG	J AM OIL CHEMISTS SOC	37	679	1960	YIELDED 9 ENTRIES
	AULT WC							
60009	NAKAGAWA T	TORI K	KOLLOID-Z	168	132	1960	NO ENTRIES	
60010	GINN ME	KINNEY FB	HARRIS JC	J AM OIL CHEMISTS SOC	37	183	1960	YIELDED 78 ENTRIES
60011	HAYDON DA	TAYLOR FH	PHIL TRANS ROY SOC LONDON, SER A	252	225	1960	YIELDED 8 ENTRIES	
60012	LANGE H		PROC INTERN CONGR SURFACE ACTIVITY	3RD	279	1960	YIELDED 7 ENTRIES	
60013	V VOORST VA.F		PROC INTERN CONGR SURFACE ACTIVITY	3RD	276	1960	NO ENTRIES	
60014	V VOORST VA.F		TRANS FARADAY SOC	56	1078	1960	NO ENTRIES	
60015	PARREIRA HC		ANALIS ACAD BRASIL CIENC	32	207	1960	YIELDED 4 ENTRIES	
60016	KOLBEL H	KLAMANN D	KURZENDORFER P	PROC INTERN CONGR SURFACE ACTIVITY	3RD	1	1960	NO ENTRIES
60017	KOLBEL H	KLAMANN D	WAGNER E	PROC INTERN CONGR SURFACE ACTIVITY	3RD	27	1960	YIELDED 3 ENTRIES
60018	GOTTE E		PROC INTERN CONGR SURFACE ACTIVITY	3RD	45	1960	YIELDED 20 ENTRIES	
60019	DERVICHIAN DG		PROC INTERN CONGR SURFACE ACTIVITY	3RD	182	1960	NO ENTRIES	
60020	JAMES JW	PETHICA BA	PROC INTERN CONGR SURFACE ACTIVITY	3RD	227	1960	YIELDED 1 ENTRIES	
60021	STANLEY JS	RADLEY JA	PROC INTERN CONGR SURFACE ACTIVITY	3RD	246	1960	YIELDED 3 ENTRIES	
60022	V VOORST VA.F	V DEN TEMPEL M	PROC INTERN CONGR SURFACE ACTIVITY	3RD	248	1960	NO ENTRIES	
60023	LUCK W		PROC INTERN CONGR SURFACE ACTIVITY	3RD	264	1960	NO ENTRIES	
60024	BOTRE C	CRESCENZI V	LIQUORI AM	PROC INTERN CONGR SURFACE ACTIVITY	3RD	302	1960	YIELDED 3 ENTRIES
60025	V VOORST VA.F		TRANS FARADAY SOC	56	1067	1960	YIELDED 9 ENTRIES	
60026	HUGO WB	NEWTON JM	J PHARM PHARMACOL	12	447	1960	YIELDED 2 ENTRIES	
60027	ELWORTHY PH		J PHARM PHARMACOL	012	293	1960	YIELDED 9 ENTRIES	
60028	BISWAS AK	MUKHERJI BK	J PHYS CHEM	64	1	1960	YIELDED 11 ENTRIES	
60029	MIYAMOTO S		BULL CHEM SOC JAPAN	33	375	1960	YIELDED 25 ENTRIES	
60030	BESPYATOV MP	LESHCHENKO ZY	MASLOB ZHIR PROM	26	24	1960	NO ENTRIES	
60031	BESPYATOV MP	OLEINIKOVA ZV	MASLOB ZHIR PROM	26	26	1960	NO ENTRIES	
60032	DEMCHENKO PA	DUMANSKII AV	DOKLADY AKAD NAUK SSSR	131	120	1960	YIELDED 2 ENTRIES	
60033	TAUBMAN AB	KONSTANTIN. VV KRYUKOVA AS	KHIM TEKHNOL	5	61	1960	YIELDED 28 ENTRIES	
60034	DEMCHENKO PA		MASLOB ZHIR PROM	26	26	1960	YIELDED 4 ENTRIES	
60035	MIYAMOTO S		BULL CHEM SOC JAPAN	33	371	1960	NO ENTRIES	
61001	HOYER HW	MARMO A	ZOELLNER M	J PHYS CHEM	65	1804	1961	YIELDED 2 ENTRIES
61002	HOYER HW	MARMO A		J PHYS CHEM	65	1807	1961	YIELDED 8 ENTRIES
61003	BECHER P		J COLLOID SCI	16	49	1961	YIELDED 7 ENTRIES	
61004	CORKILL JM	GOODMAN JF	OTTEWILL RH	TRANS FARADAY SOC	57	1627	1961	YIELDED 6 ENTRIES
61005	MYSELS KJ	KAPAUAN P		J COLLOID SCI	16	481	1961	YIELDED 5 ENTRIES
61006	HARRIS JC	SATANEK J	J AM OIL CHEMISTS SOC	38	169	1961	NO ENTRIES	
61007	FLOCKHART BD		J COLLOID SCI	16	484	1961	YIELDED 83 ENTRIES	
61008	SHINODA K	YAMAGUCHI T	HORI R	BULL CHEM SOC JAPAN	34	237	1961	YIELDED 11 ENTRIES
61009	MARUTA I	TOKIWA F		J CHEM SOC JAPAN, PURE CHEM SEC	82	1657	1961	NO ENTRIES
61010	MARUTA I	SAKAI T	TOKIWA F	J CHEM SOC JAPAN, PURE CHEM SEC	82	1512	1961	NO ENTRIES
	SAITO T							
61011	MARUTA I	TOKIWA F		J CHEM SOC JAPAN, PURE CHEM SEC	82	1660	1961	NO ENTRIES
61012	ISEMURA T	IKEDA S	TOKIWA F	BULL CHEM SOC JAPAN	34	1236	1961	NO ENTRIES
	NOGUCHI J							
61013	PILPEL N			TRANS FARADAY SOC	57	1426	1961	NO ENTRIES
61014	GINN ME	HARRIS JC		J AM OIL CHEMISTS SOC	38	605	1961	YIELDED 16 ENTRIES
61015	GINN ME	KINNEY FB	HARRIS JC	J AM OIL CHEMISTS SOC	38	138	1961	YIELDED 21 ENTRIES
61016	BRUNING W	HOLTZER A		J AM CHEM SOC	83	4865	1961	YIELDED 4 ENTRIES
61017	MYSELS KJ	OTTER RJ		J COLLOID SCI	16	462	1961	YIELDED 13 ENTRIES
61018	MYSELS KJ	OTTER RJ		J COLLOID SCI	16	474	1961	NO ENTRIES
61019	GINN ME	BROWN EL	HARRIS JC	J AM OIL CHEMISTS SOC	38	361	1961	NO ENTRIES
61020	NAKAGAWA T			CHEM IND (LONDON)	14	1135	1961	NO ENTRIES
61023	ASCOLI F	BOTRE C	LIQUORI AM	J MOL BIOL	3	202	1961	NO ENTRIES
61024	ASCOLI F	BOTRE C	LIQUORI AM	J PHYS CHEM	65	1991	1961	NO ENTRIES
61025	TYUZO K			KOLLOID-Z	175	40	1961	YIELDED 2 ENTRIES
61026	V VOORST VA.F			TRANS FARADAY SOC	57	110	1961	YIELDED 5 ENTRIES
61027	COHEN I	VASSILIADES T		J PHYS CHEM	65	1781	1961	YIELDED 2 ENTRIES
61028	COHEN I	VASSILIADES T		J PHYS CHEM	65	1774	1961	NO ENTRIES
61029	ANGELESCU E.	NICOLESCU A	BARBULESCU EM	REV CHIM AC.REP POP.ROUM.	6	309	1961	YIELDED 6 ENTRIES
61030	DEMCHENKO PA			KOLLOID ZH	23	528	1961	YIELDED 4 ENTRIES
61031	DEMCHENKO PA			DOPOV. AKAD NAUK UKR RSR		928	1961	YIELDED 19 ENTRIES
61032	DEMCHENKO PA			UKR KHIM ZH	27	322	1961	NO ENTRIES
61033	DEMCHENKO PA			MASLOB ZHIR PROM	27	19	1961	NO ENTRIES
62001	BECHER P			J PHYS CHEM	66	374	1962	YIELDED 6 ENTRIES
62002	BECHER P			J COLLOID SCI	17	325	1962	YIELDED 27 ENTRIES
62003	DEBYE P	COLL H		J COLLOID SCI	17	220	1962	NO ENTRIES
62004	HAYDON DA	TAYLOR FH		TRANS FARADAY SOC	58	1233	1962	YIELDED 16 ENTRIES
62005	HERRMANN KW			J PHYS CHEM	66	295	1962	YIELDED 12 ENTRIES
62006	CORKILL JM	GOODMAN JF		TRANS FARADAY SOC	58	206	1962	YIELDED 12 ENTRIES
62007	MARUTA I	TOKIWA F		J CHEM SOC JAPAN, PURE CHEM SEC	83	732	1962	NO ENTRIES
62008	STIRTON AJ	BISTLINE RG	WEIL JK	J AM OIL CHEMISTS SOC	39	55	1962	YIELDED 13 ENTRIES
	AULT WC							
62009	KURIYAMA K			KOLLOID-Z	180	55	1962	YIELDED 49 ENTRIES
62010	KURIYAMA K	INOUE H	NAKAGAWA T	KOLLOID-Z	183	68	1962	YIELDED 22 ENTRIES
62011	KURIYAMA K			KOLLOID-Z	181	144	1962	YIELDED 20 ENTRIES
62012	TOKIWA F	ISEMURA T		BULL CHEM SOC JAPAN	35	1737	1962	NO ENTRIES

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62013	BALMBRA RR	CLUNIE JS	CORKILL JM	TRANS FARADAY SOC	58	1661	1962	NO ENTRIES
	GOODMAN JF			J COLLOID SCI	17	243	1962	NO ENTRIES
62014	TARTAR HV			J COLLOID SCI	17	523	1962	YIELDED 4 ENTRIES
62015	MULLEY BA	METCALF AD		Z PHYSIK CHEM (FRANKFURT)	31	397	1962	NO ENTRIES
62016	HUTCHINSON E	SHAFFER PM		ANN CHIM (ROME)	52	1199	1962	NO ENTRIES
62017	BOTRE C	SCIBONA G		FETTE, SEIFEN, ANSTRICHMI	64	457	1962	NO ENTRIES
62018	LANGE H			J COLLOID SCI	17	801	1962	YIELDED 71 ENTRIES
62019	SCHICK MJ			J PHYS CHEM	66	1326	1962	YIELDED 14 ENTRIES
62020	SCHICK MJ	ATLAS SM	EIRICH FR	PONTIF ACAD SCIENTIAR. SCRIPTA VAR.	22	/12/1	1962	NO ENTRIES
62021	LIQUORI AM	ASCOLI F	BOTRE C	BULL CHEM SOC JAPAN	35	240	1962	NO ENTRIES
62022	ISEMURA T	TOKIWA F	IKEDA S	KOLLOID-Z	181	139	1962	YIELDED 6 ENTRIES
62023	WACHS W	HAYANO S		J PHYS CHEM	66	363	1962	NO ENTRIES
62024	KITAHARA A	KOBAYASHI T	TACHIBANA T	J COLLOID SCI	17	231	1962	NO ENTRIES
62025	KAUFMAN S			J CHEM SOC JAPAN, PURE CHEM SEC	83	786	1962	NO ENTRIES
62026	MARUTA I			J PHARM PHARMACOL	14	100	1962	YIELDED 6 ENTRIES
62027	ELWORTHY PH	MACFARLANE CB		J CHEM SOC JAPAN, PURE CHEM SEC	83	395	1962	NO ENTRIES
62028	MARUTA I			J CHEM SOC JAPAN, PURE CHEM SEC	83	782	1962	NO ENTRIES
62029	MARUTA I			J CHEM SOC JAPAN, PURE CHEM SEC	83	788	1962	NO ENTRIES
62030	MARUTA I			J CHEM SOC JAPAN, PURE CHEM SEC	83	858	1962	NO ENTRIES
62031	MARUTA I			J CHEM SOC JAPAN, PURE CHEM SEC	83	861	1962	NO ENTRIES
62032	MARUTA I			BULL CHEM SOC JAPAN	35	1050	1962	NO ENTRIES
62033	MATUURA R	SATAKE I	IWAMATSU I	LAKSHMINAR. GR CAN J CHEM	40	839	1962	NO ENTRIES
	HOSOKAWA S			J PHYS CHEM	66	1839	1962	YIELDED 15 ENTRIES
62034	CAMPBELL AN	KARTZMARK EM		J PHYS CHEM	66	1359	1962	YIELDED 5 ENTRIES
62035	TUDDENHAM RF	ALEXANDER AE		UKR KHIM ZH	28	46	1962	YIELDED 6 ENTRIES
62036	HAMANN SD			MASLOB ZHIR PROM	28	611	1962	YIELDED 5 ENTRIES
62037	DEMCHENKO PA			J PHYS CHEM	66	2239	1962	YIELDED 4 ENTRIES
62038	DEMCHENKO PA	ZAKHAROVA NN	DEMCHENKO LG	J PHYS CHEM	67	2075	1963	YIELDED 12 ENTRIES
62039	BESPYATOV MP	LESHCHENKO ZY		CHEM REV	63	221	1963	NO ENTRIES
62040	KURZ J L			BULL CHEM SOC JAPAN	36	222	1963	NO ENTRIES
63001	SHEDLOVSKY S	JAKOB CW	EPSTEIN MB	BULL CHEM SOC JAPAN	36	1589	1963	NO ENTRIES
63002	PILPEL N			J PHARM PHARMACOL	15	522	1963	NO ENTRIES
63003	TOKIWA F			BULL CHEM SOC JAPAN	36	1250	1963	NO ENTRIES
63004	TOKIWA F			BULL CHEM SOC JAPAN	36	281	1963	NO ENTRIES
63005	ROBINS DC	THOMAS IL		KOLLOID-Z	189	50	1963	YIELDED 49 ENTRIES
63006	KAKIUCHI K	HATTORI K	ISEMURA T	KOLLOID-Z	188	47	1963	YIELDED 6 ENTRIES
63007	TOKIWA F			KOLLOID-Z	187	44	1963	YIELDED 3 ENTRIES
63008	TORI K	NAKAGAWA T		KOLLOID-Z	191	48	1963	YIELDED 4 ENTRIES
63009	TORI K	NAKAGAWA T		KOLLOID-Z	40	538	1963	YIELDED 43 ENTRIES
63010	TORI K	NAKAGAWA T		J AM OIL CHEMISTS SOC				
63011	TORI K	KURIYAMA K	NAKAGAWA T					
63012	WEIL JK	SMITH FD	STIRTON AJ					
63013	BISTLINE RG							
63014	CORKILL JM	GOODMAN JF	OGDEN CP	PROC ROY SOC	273	84	1963	YIELDED 2 ENTRIES
	TATE JR							
63015	BECKETT AH	WOODWARD RJ		J PHARM PHARMACOL	15	422	1963	YIELDED 25 ENTRIES
63016	ANACKER EW	GHOSE HM		J PHYS CHEM	67	1713	1963	YIELDED 7 ENTRIES
63017	CROOK EH	FORDYCE DB	TREBBI GF	J PHYS CHEM	67	1987	1963	YIELDED 54 ENTRIES
63018	CORKILL JM	HERRMANN KW		J PHYS CHEM	67	935	1963	NO ENTRIES
63019	GRABENSTETT RJ	CORKILL JM		J COLLOID SCI	18	401	1963	NO ENTRIES
63020	BECHER P			J COLLOID SCI	18	196	1963	YIELDED 5 ENTRIES
63021	DONBROW M	JAN ZA		J PHARM PHARMACOL	15	825	1963	YIELDED 6 ENTRIES
63022	BECHER P			J COLLOID SCI	18	665	1963	NO ENTRIES
63023	TOKIWA F			BULL CHEM SOC JAPAN	36	1585	1963	NO ENTRIES
63024	BOTRE C	RICCieri FM		J PHARM SCI	52	1011	1963	NO ENTRIES
63025	ASCOLI F	BOTRE C		BIOPOLYMERS	1	353	1963	NO ENTRIES
63026	SCHICK MJ			J PHYS CHEM	67	1796	1963	YIELDED 59 ENTRIES
63027	SCHICK MJ			J AM OIL CHEMISTS SOC	40	680	1963	NO ENTRIES
63028	SCHICK MJ			J COLLOID SCI	18	378	1963	NO ENTRIES
63029	HUTCHINSON E	TOKIWA F		J CHEM EDUC	40	472	1963	NO ENTRIES
63030	PACKTER A	DONBROW M		J PHARM PHARMACOL	15	317	1963	YIELDED 12 ENTRIES
63031	ROBINS DC	THOMAS IL		J PHARM PHARMACOL	15	157	1963	NO ENTRIES
63032	MUKERJEE P	RAY A		J PHYS CHEM	67	190	1963	YIELDED 18 ENTRIES
63033	EKWALL P	EIKREM H	MANDELL L	ACTA CHEM SCAND	17	111	1963	NO ENTRIES
63034	SATAKE I	IWAMATSU I	HOSOKAWA S	BULL CHEM SOC JAPAN	36	204	1963	YIELDED 14 ENTRIES
	MATUURA R							
63035	SATAKE I	MATUURA R		BULL CHEM SOC JAPAN	36	813	1963	NO ENTRIES
63036	SHINODA K	NAKAYAMA H		J COLLOID SCI	18	705	1963	NO ENTRIES
63037	KATO Y			CHEM PHARM BULL (TOKYO)	11	1202	1963	YIELDED 3 ENTRIES
63038	BESPYATOV MP	LESHCHENKO ZY		MASLOB ZHIR PROM	29	19	1963	NO ENTRIES
63039	LESYUIS AA	KARNAUKH AM		MASLOB ZHIR PROM	29	22	1963	NO ENTRIES
64001	VENABLE RL	NAUMAN RV		J PHYS CHEM	68	3498	1964	YIELDED 9 ENTRIES
64002	MAURER EW	STIRTON AJ	AULT WC	J AM OIL CHEMISTS SOC	41	205	1964	YIELDED 30 ENTRIES
	WEIL JK							
64003	CORKILL JM	GOODMAN JF	HARROLDS SP	TRANS FARADAY SOC	60	202	1964	YIELDED 25 ENTRIES
64004	SCHOTT H			J PHYS CHEM	68	3612	1964	YIELDED 2 ENTRIES
64005	BECHER P			PROC INTERN CONGR SURFACE ACTIVITY	4TH		1964	NO ENTRIES
64006	HERRMANN KW			J PHYS CHEM	68	1540	1964	YIELDED 4 ENTRIES
64007	HOYER HW	DOERR IL		J PHYS CHEM	68	3494	1964	YIELDED 1 ENTRIES
64008	BECHER P	DEL VECCHIO AJ		J PHYS CHEM	68	3511	1964	NO ENTRIES
64009	CARLESS JE	CHALLIS RA	MULLEY BA	J COLLOID SCI	19	201	1964	YIELDED 16 ENTRIES

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Reference No.	Authors		Journal	Vol.	Pg.	Year	No. of Entries from Reference	
64010	MANKOWICH AM		J AM OIL CHEMISTS SOC	41	449	1964	YIELDED 6 ENTRIES	
64011	SHINODA K	KATSURA K	J PHYS CHEM	68	1568	1964	YIELDED 24 ENTRIES	
64012	CORKILL JM	GOODMAN JF	TRANS FARADAY SOC	60	986	1964	YIELDED 1 ENTRIES	
64013	CROOK EH	FORDYCE DB	J AM OIL CHEMISTS SOC	41	231	1964	NO ENTRIES	
64014	CROOK EH	TREBBI GF	J PHYS CHEM	68	3592	1964	YIELDED 80 ENTRIES	
64015	NAKAGAWA T	TORI K	KOLLOID-Z	194	143	1964	NO ENTRIES	
64016	BENJAMIN L		J PHYS CHEM	68	3575	1964	YIELDED 5 ENTRIES	
64017	ANACKER EW	RUSH RM	J PHYS CHEM	68	81	1964	YIELDED 4 ENTRIES	
64018	ANACKER EW	WESTWELL AE	J PHYS CHEM	68	3490	1964	NO ENTRIES	
64019	INOUE H		KOLLOID-Z	196	1	1964	NO ENTRIES	
64020	SCHICK MJ		J PHYS CHEM	68	3585	1964	YIELDED 108 ENTRIES	
64021	NAKAGAWA T	INOUE H	KOLLOID-Z	195	93	1964	NO ENTRIES	
64022	PILPEL N		NATURE	204	378	1964	NO ENTRIES	
64023	BALMBRA RR	CLUNIE JS	CORKILL JM	TRANS FARADAY SOC	60	979	1964	YIELDED 8 ENTRIES
	GOODMAN JF							
64024	TOKIWA F		BULL CHEM SOC JAPAN	37	1837	1964	YIELDED 12 ENTRIES	
64025	NAKAGAKI M	NINOMIYA Y	BULL CHEM SOC JAPAN	37	817	1964	YIELDED 1 ENTRIES	
64026	CAMPBELL AN	GIESKES JMTM	CAN J CHEM	43	1004	1964	NO ENTRIES	
64027	CORKILL JM	GOODMAN JF	TATE JR	TRANS FARADAY SOC	60	996	1964	YIELDED 2 ENTRIES
64028	BERRY RWH	BROCKLEHURST P	J CHEM SOC		2264	1964	NO ENTRIES	
64030	HARTLEY GS		CHEM IND (LONDON)	24	1012	1964	NO ENTRIES	
64031	BOTRE C	SOLINAS M	BIOCHEM BIOPHYS ACTA	88	415	1964	NO ENTRIES	
64032	BOTRE C	DE MARTIIS F	SOLINAS M	J PHYS CHEM	68	3624	1964	YIELDED 1 ENTRIES
64033	LIQUORI AM	BOTRE C	RIC SCI	6	71	1964	NO ENTRIES	
64034	NAKAGAKI M	KAWAMURA S	YAKUGAKU ZASSHI	84	246	1964	YIELDED 5 ENTRIES	
64035	SOMASUNDARAN P	HEALY TW	FUERSTENAU DW	J PHYS CHEM	68	3562	1964	YIELDED 5 ENTRIES
64036	MIURA M	FUJITA H	WATARI Y	J SCI HIROSHIMA UNIV. SER A-II	28	41	1964	NO ENTRIES
64037	HUTCHINSON E	SHEAFFER VE	TOKIWA F	J PHYS CHEM	68	2818	1964	YIELDED 7 ENTRIES
64038	FUERSTENAU DW	HEALY TW	SOMASUNDARAN P	TRANS SME AIME		321	1964	NO ENTRIES
64039	LANGE H		PROC INTERN CONGR SURFACE ACTIVITY	4TH	1	1964	NO ENTRIES	
64040	NAKAGAWA T	INOUE H	PROC INTERN CONGR SURFACE ACTIVITY	4TH	N	1964	NO ENTRIES	
64041	KAUFMAN S		J PHYS CHEM	68	2814	1964	NO ENTRIES	
64042	LITTLE RC	SINGLETERRY CR	J PHYS CHEM	68	3453	1964	NO ENTRIES	
64043	WASIK SP	HUBBARD WD	J RES NAT BUR STD A	68	359	1964	YIELDED 13 ENTRIES	
64044	LITTLE RC	SINGLETERRY CR	J PHYS CHEM	68	2709	1964	NO ENTRIES	
64045	HUISMAN HF		KONINKI NED AKAD WETEN. PROC SER B	67	367	1964	NO ENTRIES	
64046	HUISMAN HF		KONINKI NED AKAD WETEN. PROC SER B	67	376	1964	NO ENTRIES	
64047	HUISMAN HF		KONINKI NED AKAD WETEN. PROC SER B	67	388	1964	YIELDED 59 ENTRIES	
64048	HUISMAN HF		KONINKI NED AKAD WETEN. PROC SER B	67	407	1964	NO ENTRIES	
64049	ELWORTHY PH	FLORENCE AT	KOLLOID-Z	195	23	1964	YIELDED 20 ENTRIES	
64050	ADDERSON JE	TAYLOR H	J COLLOID SCI	19	495	1964	YIELDED 14 ENTRIES	
64051	MARKINA ZN	TSIKURINA NN	KOSTOVA NZ	KOLLOID ZH	26	76	1964	YIELDED 31 ENTRIES
65001	CROOK EH	FORDYCE DB	TREBBI GF	J COLLOID SCI	20	191	1965	NO ENTRIES
65002	KONNO K	KITAHARA A	KOGYO KAGAKU ZASSHI	68	2058	1965	NO ENTRIES	
65003	MOLYNEUX P	RHODES CT	SWARBICK J	TRANS FARADAY SOC	61	1043	1965	YIELDED 13 ENTRIES
65004	CORKILL JM	GOODMAN JF	WALKER T	TRANS FARADAY SOC	61	589	1965	NO ENTRIES
65005	CORKILL JM	GOODMAN JF	OGDEN CP	TRANS FARADAY SOC	61	583	1965	YIELDED 3 ENTRIES
65006	EMERSON MF	HOLTZER A	J PHYS CHEM	69	3718	1965	NO ENTRIES	
65007	BUJAKE JE	GODDARD ED	TRANS FARADAY SOC	61	190	1965	YIELDED 2 ENTRIES	
65010	BOTRE C	BRUFANI M	RICCIERI FM	J PHARM SCI	54	919	1965	NO ENTRIES
65011	SCHICK MJ	GILBERT AH	J COLLOID SCI	20	464	1965	YIELDED 24 ENTRIES	
65012	STEIGMAN J	COHEN I	SPINGOLA F	J COLLOID SCI	20	732	1965	YIELDED 9 ENTRIES
65013	BECHER P		J COLLOID SCI	20	728	1965	YIELDED 9 ENTRIES	
65014	TOKIWA F		BULL CHEM SOC JAPAN	38	751	1965	NO ENTRIES	
65015	V VOORST VA.F							
65016	ELWORTHY PH	MACFARLANE CB	J PHARM PHARMACOL	17	65	1965	NO ENTRIES	
65017	ARAI H	SHIGEHIRO F	MARUTA I	KOGYO KAGAKU ZASSHI	68	1090	1965	NO ENTRIES
65018	MYSELS EK	MYSELS KJ	J COLLOID SCI	20	315	1965	YIELDED 6 ENTRIES	
65019	ABU-HAMDIYY M	MYSELS KJ	J PHYS CHEM	69	1466	1965	YIELDED 6 ENTRIES	
65020	SHIRAHAMA K	MATUURA R	BULL CHEM SOC JAPAN	38	373	1965	YIELDED 34 ENTRIES	
65021	MALIK WU	VERMA SP	INDIAN J CHEM	3	441	1965	NO ENTRIES	
65022	SHIGEHARA K		BULL CHEM SOC JAPAN	38	1700	1965	YIELDED 3 ENTRIES	
65023	EKWALL P	HOLMBERG P	ACTA CHEM SCAND	19	573	1965	NO ENTRIES	
65024	CAMPBELL AN	LAKSHMINAR. GR	CAN J CHEM	43	1729	1965	YIELDED 16 ENTRIES	
65025	STEIGMAN J	SHANE N	J PHYS CHEM	69	968	1965	YIELDED 13 ENTRIES	
65026	WEINER ND	ZOGRAFI G	J PHARM SCI	54	436	1965	YIELDED 6 ENTRIES	
65027	BJAASTAD SG	HALL NA	THAKKAR AL	J PHARM SCI	54	1529	1965	NO ENTRIES
65028	KUPPUSAMI J	SURYANARAY.CV	NATURE	208	780	1965	YIELDED 4 ENTRIES	
65029	MIJNLIEFF PF	DITMARSCH R	NATURE	208	889	1965	NO ENTRIES	
65030	TONG LKJ	REEVES RL	J PHYS CHEM	69	2357	1965	YIELDED 3 ENTRIES	
65031	CZERNIAWSKI M		ROCNZ CHEM	39	1059	1965	YIELDED 4 ENTRIES	
65032	CZERNIAWSKI M		ROCNZ CHEM	39	1469	1965	NO ENTRIES	
65033	INOUE H	NAKAGAWA T	J PHYS CHEM	70	1108	1965	NO ENTRIES	
65034	LELONG ALM	CONSOLE L	ANALES ASOC QUIM ARGENT	53	39	1965	NO ENTRIES	
65035	LELONG ALM	NATALE I	ANALES ASOC QUIM ARGENT	53	11	1965	NO ENTRIES	
65036	OSUGI J	SATO M	IFUKU N	REV PHYS CHEM JAPAN	35	32	1965	YIELDED 19 ENTRIES
65037	CZERNIAWSKI M		ROCNZ CHEM	39	1275	1965	YIELDED 4 ENTRIES	
66001	SCHICK MJ	MANNING DJ	J AM OIL CHEMISTS SOC	43	133	1966	YIELDED 67 ENTRIES	
66002	MUKERJEE P	KAPAUAN P	MEYER HG	J PHYS CHEM	70	783	1966	YIELDED 4 ENTRIES
66003	WEIL JK	STIRTON AJ	BARR EA	J AM OIL CHEMISTS SOC	43	157	1966	YIELDED 5 ENTRIES

List of References—Continued

Reference No.	Authors		Journal	Vol.	Pg.	Year	No. of Entries from Reference
66004	MUKERJEE P	RAY A	J PHYS CHEM	70	2138	1966	NO ENTRIES
66005	MUKERJEE P	RAY A	J PHYS CHEM	70	2144	1966	NO ENTRIES
66006	MUKERJEE P	RAY A	J PHYS CHEM	70	2150	1966	YIELDED 10 ENTRIES
66007	ELWORTHY PH	MYSELS KJ	J COLLOID SCI	21	331	1966	YIELDED 3 ENTRIES
66008	ELWORTHY PH	FLORENCE AT	KOLLOID-Z Z POLYMERE	208	157	1966	NO ENTRIES
66009	SHIGEHARA K		BULL CHEM SOC JAPAN	39	2643	1966	NO ENTRIES
66010	SHIGEHARA K		BULL CHEM SOC JAPAN	39	2332	1966	YIELDED 3 ENTRIES
66011	CARDWELL PH		J COLLOID INTERFACE SCI	22	430	1966	YIELDED 6 ENTRIES
66012	BENJAMIN L		J COLLOID INTERFACE SCI	22	386	1966	YIELDED 19 ENTRIES
66013	HERRMANN KW		J COLLOID INTERFACE SCI	22	352	1966	YIELDED 12 ENTRIES
66014	CORKILL JM	GOODMAN JF	HARROLD SP	62	994	1966	YIELDED 10 ENTRIES
	TATE JR						
66015	CORKILL JM	GOODMAN JF	ROBSON P	62	987	1966	YIELDED 12 ENTRIES
	TATE JR						
66016	CORKILL JM	GOODMAN JF	TATE JR	62	979	1966	NO ENTRIES
66017	PATEL RM	ZOGRAFI G		55	1345	1966	NO ENTRIES
66018	WAN LSC			55	1395	1966	YIELDED 5 ENTRIES
66019	DONBROW M	JACOBS J		18	92S	1966	YIELDED 2 ENTRIES
66020	FLORENCE AT			18	384	1966	YIELDED 1 ENTRIES
66021	MANKOWICH AM			43	615	1966	YIELDED 11 ENTRIES
66022	KOMOR JA	BEISWANGE. JPG		43	435	1966	YIELDED 12 ENTRIES
66023	MALIK WU	CHAND P		43	446	1966	YIELDED 16 ENTRIES
66024	LOVELL VM	SEBBA F		38	1926	1966	NO ENTRIES
66025	SCHICK MJ			43	681	1966	YIELDED 45 ENTRIES
66026	NAKAYAMA H	SHINODA K	HUTCHINSON E	70	3502	1966	NO ENTRIES
66027	TOKIWA F	OHKI K		70	3437	1966	YIELDED 6 ENTRIES
66028	FORD WPJ	OTTEWILL RH	PARREIRA HC	21	522	1966	YIELDED 21 ENTRIES
66029	CZERNIAWSKI M			40	1265	1966	NO ENTRIES
66030	CZERNIAWSKI M			40	1935	1966	YIELDED 4 ENTRIES
66031	MATHAI K	OTTEWILL RH		62	759	1966	NO ENTRIES
66032	IDA O	MEGURO K	KONDO A	77	905	1966	NO ENTRIES
66033	OSUGI J	SATO M	IFUKU N	87	329	1966	NO ENTRIES
66034	PYTASZ G	SZEGLOWSKI Z		211	199	1966	NO ENTRIES
66035	PYTASZ G	SZEGLOWSKI Z		211	209	1966	NO ENTRIES
66036	SCHOTT H			70	2966	1966	YIELDED 8 ENTRIES
66037	MATHAI K	OTTEWILL R		62	750	1966	YIELDED 1 ENTRIES
66038	BENTON D	SPARKS B		62	3244	1966	YIELDED 6 ENTRIES
66039	HERRMANN WK	BRUSHMILLER J	COURCHENE W	70	2909	1966	YIELDED 7 ENTRIES
66040	BENJAMIN L			70	3790	1966	YIELDED 11 ENTRIES

Abbreviations: Units of Measure

Code	Description
A	MOLES/100 MOLES OF SOLVENT (INCLUDING ADDITIVES)
B	VOLUME % OF SOLVENT
C	MOLES/100 MOLES OF SURFACTANT MIXTURE
D	W/V % (GRAMS/100 MILLILITERS OF SOLUTION)
E	% SATURATION OF SOLUTION BY ADDITIVE
H	GRAMS/100 GRAMS OF SOLVENT
I	MOLES/100 MOLES OF SURFACTANT
K	TOTAL NORMALITY OF COUNTERIONS
M	MOLAR (MOLES/LITER OF SOLUTION)
N	NORMAL (EQUIVALENTS/LITER OF SOLUTION)
P	W/W % (GRAMS/100 GRAMS OF SOLUTION)
Q	GRAMS/100 GRAMS OF SURFACTANT
R	VALUE VARIED DURING EXPERIMENT
S	MOLES/KILOGRAM OF SOLUTION
T	GRAMS/100 GRAMS OF SURFACTANT MIXTURE
U	MOLES/LITER OR KILOGRAM OF SOLUTION (UNSPECIFIED)
W	MOLAL (MOLES/KILOGRAM OF SOLVENT)
Y	PRESSURE IN ATMOSPHERES

Abbreviations: Words and Names

Abbreviations	Meaning	Abbreviations	Meaning
AZBZ	AZOBENZENE	N-3SOA*	N-(3SULFOLANYL) OLEYL AMIDE
BRPB	BROMPHENOL BLUE (DYE)	OROT	ORANGE OT (DYE)
BZL*	BENZYL	PDMAB	P-DIMETHYLAMINOAZOBENZENE
BZP4	BENZOPURPURINE 4B (DYE)	PLT	PLOT (GRAPH)
COND	CONDUCTANCE (ELECTRIC)	PMS*	POLYMETHYLSILOXANE
DCFL	DICHLOROFLUORESCINE (DYE)	PNCN	PINACYANOL
DMYL	DIMETHYL YELLOW (DYE)	RHD6	RHODAMINE 6G (DYE)
EOSN	EOSINE (DYE)	SDN4	SUDAN 4 (DYE)
EQN	EQUATIONS	SKYB	SKY BLUE SF (DYE)
ERTS	ERYTHROSIN (DYE)	SP	SPECIFIC
FL	FLUORESCINE (DYE)	SPCTR	SPECTRAL
FLUOR	FLUORESCENCE	TMCHCGLET*	TRIMETHYL CYCLOHEXYL CARBONYL
FOTMTR	PHOTOMETRIC	TNSN	GLYCEROL ETHER
GLET*	GLYCERINE ETHER	UNSPEC	TENSION
HXMTP*	HEXAMETAPHOSPHATE	V.BR*	NOT SPECIFIED BY REFERENCE
INPX	INDOPHENOL (DYE)	YLOB	VERY BRANCHED
I2	IODINE	2NPA	YELLOW OB (DYE)
LOGMIC	LOGARITHMIC GRAPH		2-NITRODIPHENYLAMINE
LOH	LAURYL (DODECYL) ALCOHOL		

Source Symbols

Source Code	Description
A	MISPRINT CORRECTED IN FIGURE QUOTED IN REFERENCE
C	UNITS CONVERTED FROM TABULATION IN REFERENCE
E	FROM EQUATIONS IN REFERENCE OF METHOD RESULTS
G	FROM GRAPH OF CMC VALUES IN REFERENCE
K	FROM GRAPH OF METHOD RESULTS IN REFERENCE
L	ALSO PERSONAL COMMUNICATIONS FROM AUTHOR
M	UNITS CONVERTED FROM PRECEDING
P	FROM OUR PLOT OF DATA IN REFERENCE
S	AVERAGE OF TABULATED VALUES
T	TABULATED OR FIGURE QUOTED IN REFERENCE

The code E in the numerical value of the concentration in the "Additive" column of the CMC tables stands for "times ten to the power".

Symbols for the quality of materials and methods are discussed on page 6 et seq.
Symbols for methods are discussed on page 8 et seq.

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