

ANNIVERSARY REVIEW

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MHC ligands and peptide motifs: first listing

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Introduction

The purpose of this article is to provide a compendium of major histocompatibility complex (MHC) peptide motifs and MHC ligands known to date, together with a discussion of the methods used to gain this information. A problem here is the exponential growth of information in this field over the recent years. The number of known MHC ligands was zero in 1989 and three in 1990. This article, written in 1994, lists a couple of hundred such ligands, plus a large number of likely ligands. By the time this work is published, we expect a lot more ligands to be known. On the other hand, the peptide motifs of many of the more important MHC class I molecules are known already, so that this article will still be useful as a source of information. For class II, the situation is a bit different. Only a few allele-specific motifs have been reported, and the data from different authors are partially conflicting. The principles of allele-specific ligand motifs, however, have emerged recently by the combination of information on MHC class II structure, ligand sequencing, and peptide binding assays. Thus, these principles can be applied to further ligands to be identified.

Overview of MHC function

MHC molecules are peptide receptors. Their function is to collect peptides inside the cell and to transport them to the cell surface, where the complex of peptide and MHC molecule may be recognized by the T-cell receptor (TCR) for antigen of T lymphocytes (Rammensee et al. 1993). In normal cells, MHC-associated peptides are derived from normal, that is, self proteins. During their differentiation,

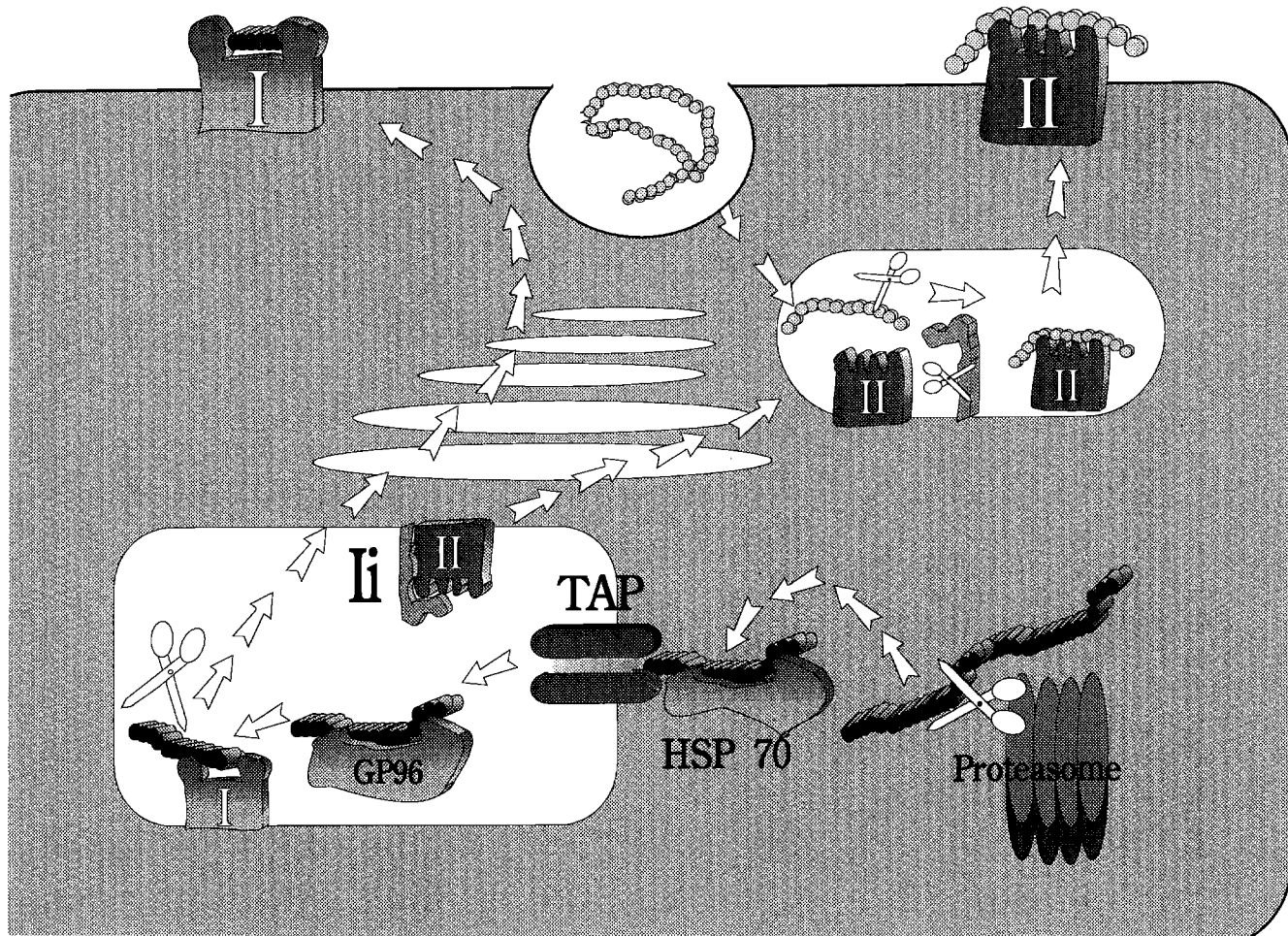
T cells become tolerant to complexes of self peptides and self MHC molecules (Von Boehmer 1992). Thus, if any new peptides, e.g., derived from an infectious agent, occur later, they can be recognized by T cells. Since the specific immune system is regulated by T cells, the trimolecular complex of TCR, MHC molecule, and peptide can be considered a control switch for the immune system. Thus, a study of the molecular interactions between the three parts is essential for our understanding of the immune system.

Two classes of MHC molecules are distinguished, class I and class II. Class I molecules consist of a membrane-inserted heavy chain of about 45 000 M_r , and a non-covalently attached light chain of 12 000 M_r (Klein 1986). The latter is also known as β_2 -microglobulin (β_2m). The structure of class I molecules has been resolved by X-ray crystallography (Stern and Wiley 1994). It has some resemblance to a moose's head, whereby the antlers would form a groove that is recognized as a peptide-binding device. HLA-A, B, and C are the "classical" class I molecules of humans, and H-2K, H-2D, and H-2L those of the mouse. Class II molecules are heterodimers consisting of two chains α and β , of similar size (about 30 000 M_r), both of which are membrane inserted. HLA-DR, DQ, and DP are the human class II molecules, H-2A and E those of the mouse. Their structure is surprisingly similar to that of class I molecules (Stern and Wiley 1994; Stern et al. 1994; Brown et al. 1993).

All HLA molecules, including the numerous "non-classical", are encoded on chromosome 6, with the exception of β_2m which is on chromosome 15. $H2$ genes are on chromosome 17 of the mouse, and the mouse β_2m gene is on chromosome 2.

A peculiarity of MHC genes is their extensive polymorphism, characterized by the presence of dozens or hundreds of alleles in a species. $H2$ alleles are designated $H2K^b$, $H2K^d$, $H2K^k$ and so on for class I, and $H2Aa^b$, $H2Aa^k$, $H2Ab^k$, $H2Eb^d$ and so on for class II, whereby different alleles may differ in as many as 40 amino acids (Klein 1986). The present nomenclature (Bodmer et al. 1994) of HLA genes and products (which has been changed several times) is outlined as follows: class I heavy chain

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loci: *HLA-A*, *B*, and *C*; class II α chain loci: e.g., *HLA-DRA*, *DQA1*, *DPA1*, class II β chain loci: e.g., *HLA-DRB1*, *DRB3*, *DQB1*, *DPB1*. Alleles are designated, for example, *HLA-A*0201*, or *HLA-DRB1*0101*. This nomenclature can only be applied if the respective sequences are known. Since this is not the case in many situations, the old designations, e.g., *HLA-A2* or *HLA-DR3*, based on serology, are still being used, and describe collections of alleles with shared serologic determinants (e.g., *HLA-A2* for *A*0201* through *A*02012*). Both class I light chains and *HLA-DR\alpha* chains are not very polymorphic (Klein 1986). The high (*HLA-B*) or at least moderate polymorphism of the other genes results in the expression of a large number of combinations of alleles at the different loci per chromosome (haplotype), and in a high degree of heterozygosity. Thus each individual has his or her particular combination of HLA molecules, namely up to six different class I and about six different class II molecules (if the non-classical HLA molecules, whose function is not known, are not considered), making it unlikely to find two unrelated individuals with exactly the same combination of HLA genes.

A simplified outline of MHC function is given in the diagram in Figure 1. Class I molecules, both heavy and light chains, are synthesized into the ER (reviewed in Jackson and Peterson 1993). The peptides to be loaded on class I molecules are, in many cases, derived from cytosolic

Fig. 1 A simplified and partially hypothetical overview of antigen processing. For explanation see text

proteins. The details of peptide generation are not known definitely. A widely held view, however, is that cytosolic proteins are partially degraded by an endopeptidase activity of the proteasome, a multiunit structure with several activities located in the cytosol (Rock et al. 1994). It is not clear, however, how the products of such endopeptidase activity are related to the final class I ligands (Dick et al. 1994). One possibility is that the proteasomes directly produce the correct ligands. Alternatively, proteasomes could cut out larger peptides requiring further processing. The endopeptidase specificity of the proteasome is such that a protein is cut after hydrophobic or charged residues, in principle. The fine specificity of endopeptidase activity is influenced by two proteasome subunits, LMP2 and LMP7, which are encoded in the MHC region and regulated by IFN. However, the exact kind of LMP influence on specificity is controversial (Howard and Seelig 1993). In any case, such peptides must be transported into the ER lumen by the TAP molecule [(transporter associated with processing) (Neefjes and Momburg 1993)]. According to one hypothesis, these peptides are bound and protected from complete degradation by a chaperone, HSP70, before reaching TAP (Srivastava et al. 1994). Peptide transport by TAP molecules has

been directly demonstrated recently (reviewed in Momburg et al. 1994). Transport has specificity especially regarding the C-termini of peptides, and selectivity for peptide lengths. Peptides of 7 to 23 amino acids have been shown to be transported, whereby optima of 10 to 15 amino acids are seen. Human TAPs do not discriminate much between the C-termini of peptides. In contrast, the mouse TAP has a preference for peptides with hydrophobic C-termini and dislikes peptides with charged termini. This pattern of specificities fits well with the peptide specificities of human and mouse MHC class I molecules, since all mouse class I molecules require peptides with hydrophobic C-termini, whereas some human class I molecules require peptides with basic C-termini. On the other hand, mouse cells transfected with the *HLA-A3* gene, requiring peptide ligands with basic C-termini, can be loaded with the fitting peptides (Maier et al. 1994). This indicates that MHC peptide specificity need not be strictly dependent on TAP specificity. That TAP specificity indeed can influence MHC peptide loading is evident from two different TAP forms in the rat, TAP^a and TAP^u. Dependent on co-expressions of the respective TAP, the peptide spectrum of rat MHC class I molecules, RT1^u, is different, as indicated by different HPLC behavior of RT1^a-associated peptides. When measured in a peptide transporter assay, TAP^a has the same specificity as human TAP, that is, it does not discriminate between hydrophobic and basic C-termini, whereas TAP^u is more like the mouse transporter, with a preference for peptides with hydrophobic C-termini.

Once they are inside the ER lumen, the further fate of transported peptides is not exactly known. The recently reported physical association of TAP molecules and class I molecules suggested that peptides are directly loaded onto class I molecules immediately after discharge from the transporter (Ortmann et al. 1994; Suh et al. 1994). However, this would require that either the incoming peptides are already of the right size for loading to class I molecules, or that they bind as longer peptides (Falk et al. 1990) and are trimmed while somehow bound to MHC. An alternative hypothesis is that peptides are first bound by a chaperone, gp96, which shuttles the peptides to class I molecules, perhaps with some trimming of peptides underway. The main reason for assuming that gp96 plays a role in antigen processing stems from an impressive series of experiments by Srivastava and co-workers (1994), showing that gp96 molecules are associated with a large array of peptides and are able to immunize mice against antigens presented by MHC class I molecules.

In any event, the peptide somehow reaches the class I molecule and binds into the groove, perhaps after a final trimming step while already in touch with MHC. Unusually long peptides found associated with MHC class I molecules might have escaped such a final trimming (Urban et al. 1994). The assembly sequence of class I heavy chain, β_2m and peptide is not quite clear. A recent report indicates that another chaperone, calnexin, is bound to assembled complexes of heavy chain and β_2m , and thus retains empty class I molecules in the ER (Jackson et al. 1994). It is only upon peptide loading that the fully assembled heavy chain/

β_2m /peptide complex is released by calnexin for transportation to the cell surface.

Class II molecules also start their existence in the ER. The two chains, α and β , assemble and are bound by a chaperone-like molecule, the invariant chain [(Ii) (Cresswell 1994)]. This molecule has two functions; one is to direct the α,β -heterodimer to the class II loading compartment, which appears to be a specialized vesicle characterized by the presence of class II molecules. The second function of Ii is the prevention of premature peptide loading to class II molecules. The molecular interactions between Ii and the α,β -heterodimer preventing peptide binding are not completely sorted out; one possibility is an allosteric effect of Ii on the dimer such that the peptide binding groove is closed due to conformational change. The other possibility is that a particular stretch of the invariant chain binds into the groove and thereby competitively prevents the binding of peptides. This latter view is derived from the observation that Ii peptides, called CLIPs (class II-associated invariant chain peptides) are frequently found associated with immunoprecipitated class II molecules, and that CLIPs are especially abundant in cells with a defect in antigen processing. In any case, Ii is removed from the α,β -heterodimer in the class II loading compartment, or shortly before. The peptides loaded onto class II molecules can be derived not only from endocytosed protein but also from protein endogenous to the cells, especially membrane-bound proteins which have a chance to co-localize in the class II loading compartment. Finally, the peptide-loaded α,β -heterodimers are translocated to the cell surface.

The simplified view shown in Figure 1 suggests a strict separation of the processing pathways for class I and class II, respectively. There is strong evidence, however, for considerable cross-talk between the two pathways. Peptides derived from cytosolic proteins, for example, can be loaded onto class II molecules (Pinet et al. 1994). On the other hand, peptides derived from phagocytosed proteins can be loaded onto class I molecules, especially if the phagocytosed protein is aggregated (Pfeifer et al. 1993; Rock et al. 1993). Such side-lines of processing pathways deserve interest because they could be exploited for new strategies of immune intervention.

Methods of characterizing MHC/peptide interactions

The most seminal approach to gain information on the function of MHC molecules as peptide receptors is the X-ray analysis of MHC crystals (Stern and Wiley 1994). The two other principle methods are: 1) Biochemical isolation and study of naturally MHC-associated peptides, and 2) Binding studies with synthetic peptides. The latter two approaches are discussed below:

1) Analysis of natural MHC ligands

The diagram in Figure 2 gives an overview on the approaches used for isolation and analysis of MHC-associated peptides.

The major technical challenge is the small copy number of individual peptides. It is estimated that a cell presents well over 1000 different peptides on its 100 000 or so copies of a given MHC allelic product. A few of these peptides are present in high copy number, that is, up to 10 000 or more. By far the most ligands, however, occur in a much lower copy number, maybe even down to as low as one copy per cell.

The most sensitive means of detecting isolated peptides is the T-cell assay, which is able to detect peptides in the sub-picomolar range, at least as far as cytotoxic T cells are concerned (Rötzschke et al. 1990). Typically, a peptide-containing sample (e.g., a few μ l of an HPLC fraction) is incubated in a total volume of 100 μ l together with MHC-expressing, ^{51}Cr -labeled target cells. After some incubation time, e.g., 90 min, CTL are added, the supernatant is harvested 4 to 6 h later, and the relative radioactivity measured indicates the degree of target cell lysis. If the 100 μ l volume used for target cell incubation has a concentration of 1 pM, the absolute amount of peptide is 100 attomol, a sensitivity not reached by any other method. The use of the CTL assay, of course, is limited to the detection of T-cell epitopes for which T cells are on hand: Viral antigens, minor H antigens, tumor-associated antigens, transfected model antigens, or antigens recognized by alloreactive T cells. On the other hand, peptide detection assays for class-II-restricted T cells appear to be less sensitive than for class I-restricted T cells.

The major shortcoming of the T-cell assay for peptide detection is that it does not give sequence information. However, the location of a T-cell epitope among HPLC-separated MHC ligands of an infected cell can allow identification of the peptide in combination with biochemical analysis such as Edman degradation or mass spectrometry. The first naturally processed viral T-cell epitopes indeed were identified by the combination of T-cell assay with mass spectrometry, comparison of the HPLC behavior of synthetic and natural peptides, or partially direct sequencing, using radiolabeled amino acids incorporated by virus-infected cells (Rötzschke et al. 1990; van Bleek and Nathenson 1990). A combination of these methods for identification of T-cell epitopes is only possible if the proteins of origin are known. Direct sequencing of HPLC fractions containing a T-cell epitope is rarely successful, namely, only in cases where the T-cell epitope happens to be a peptide highly abundant in that fraction. A marked improvement of sensitivity was brought about by an ingenious combination of HPLC, CTL assay, and mass spectrometry by Cox and co-workers (1994).

By far the most ligands known to date are not T-cell epitopes and these ligands were determined by direct sequencing, either by Edman degradation, or by mass spectrometry, or by a combination of the two methods. Detection limit of Edman degradation is about 1 pmol, that

Source of MHC-expressing cells

(tumor cells, transformed cells, cells transfected to express a specific MHC molecule, or fresh or frozen tissue).

↓

Detergent extract

↓

Precipitation of MHC molecules with solid-phase bound antibodies

↓

Dissociation of peptides from MHC molecules with acid (0.1 % TFA or 10 % acetic acid)

↓

Ultrafiltration

↓

Separation of peptides by reversed phase HPLC

◀ ▶ ↘

T cell assay

Edman degradation

Mass spectrometry

Fig. 2 Methods for analysis of MHC ligands

is, the equivalent of 6×10^9 cells for a peptide occurring in 100 copies per cell. Sequencing by tandem mass spectroscopy has been reported to be more sensitive – down to 30 fmol or less. It is, however, challenging to achieve this degree of sensitivity, so that, apart from the pioneering group of Hunt and co-workers (1992), not many other laboratories have come up with similar results.

A special application of Edman degradation is pool sequencing, that is, altogether-sequencing of the complex mixture of peptides eluted from a given MHC species (Falk et al. 1991 b). Although disliked by purists, this approach allows one to determine the common characteristics of all peptides associated with a given MHC molecule, with relatively little effort. Pool sequencing of MHC class I ligands led to the discovery of the principle of allele-specific motifs, and allowed a large number of such motifs to be determined. The clear information that can be obtained from pool sequencing of class I ligands is made possible by their uniform length, e.g., 9 amino acids. But even for class II ligands, which can range in length from about 12 to 25 amino acids, pool sequencing is a valuable tool for gaining detailed information on motifs (Falk et al. 1994 b).

It appears that the number of amino acids between the N-terminus of class II ligands and the first anchor varies by about three amino acids for the majority of ligands. For DR1, for example, the distance from the N-terminus to the first anchor of the majority of ligands is 5 ± 1 (Falk et al. 1994 b). Thus, pool sequencing indicates a cluster at position 4, 5, and 6 for the anchor residues used, aromatic and aliphatic. Again for DR1, the next cluster stretches over

positions 7, 8, and 9, indicating the next anchor for aliphatic residues. The rough motif obtained by such interpretations – absolute position 5 set as relative position 1 to mark the first anchor – can then be complemented and worked out in depth by applying 1) alignment of natural ligands, 2) consideration of the pockets, as revealed recently by crystallography of a monopeptidic DR1 molecule (Stern et al. 1994), and 3) considerations of peptide binding assays. If all four sources of information are considered, a detailed picture of the degenerate (as compared with class I) peptide specificities of class II molecules can be obtained that should be useful for epitope predictions (Friede and co-workers, submitted).

2) Peptide binding assays

MHC/peptide binding assays have a history of leading to obsolete results. On the other hand, with our increasing knowledge of MHC structure and MHC/peptide interaction and specificity, it is possible to design straightforward peptide binding experiments to answer specific questions. A number of approaches can be used to measure peptide binding to MHC. The oldest method is as follows (Buus et al. 1987): MHC molecules are purified and incubated with radioactively labeled peptides. Then the mixture is subjected to a gel filtration column. MHC molecules with the radioactive peptide bound will elute in the exclusion volume, whereas free peptides will elute later. Thus, the amount of radioactivity in the exclusion volume is a measure for peptides bound to MHC. The binding behavior of other, unlabeled peptides can be tested via their capacity to inhibit binding of the radioactive peptide. A number of variations of this method have been used. For example, the radioactive label can be replaced by a fluorescent marker. Furthermore, MHC/peptide complexes can be separated from free peptides by gel electrophoresis, or upon binding of the MHC/peptide complex to solid phase with the help of antibodies. In the latter case, however, two different antibodies reactive with different sites of the MHC molecule are required, one for purification of the MHC molecule, the other for capturing the MHC/peptide complex from the reaction mixture.

Depending on the conditions, these kinds of peptide binding assays can be made very sensitive to detect even low-affinity peptide binding. This may result in problems of interpretations, since low-affinity binding might not be relevant for physiological MHC/peptide interactions.

A second type of binding assay depends on the stabilization of MHC class I molecules by bound peptides. Cells with a defect in antigen processing, for example, TAP-defective RMA-S cells, express only a low density of antibody-detectable MHC class I molecules on their surface, if cultured under normal conditions (37 °C). If such cells are incubated with peptides binding to the expressed class I molecules with high affinity, the latter are stabilized, and their surface density increases (Townsend et al. 1989). Since determination of class I surface density can be easily done by FACS analysis, this approach has been widely

used. Since only few cell lines with transporter defects are known, the assay can only be used for MHC molecules expressed by such cells, e.g., H-2K^b or D^b for RMA-S cells. To study peptide binding for additional MHC-molecules, the desired MHC molecule can be expressed in RMA-S or other TAP-defective cells upon gene transfection. The advantage of this MHC-stabilization assay is that it is rather insensitive and thus detects only peptides binding with high affinity that are likely to be physiologically relevant. Stabilization of MHC molecules by peptides can also be measured with purified MHC molecules.

For class II molecules, the binding of high-affinity peptides leads to a compact form of the MHC/peptide complex, as seen by SDS gel electrophoresis, whereas a peptide of lower affinity leads to a "floppy" form of class II molecules.

A very elegant approach for studying the peptide specificity of class II molecules has been developed by Hammer and co-workers (Sinigaglia and Hammer 1994). A peptide library is expressed by bacteriophages. From the peptide-expressing phages only those are selected which are able to bind to a given class II molecule. The peptide sequences expressed by the selected phages are then determined. With this approach, a peptide binding motif of HLA-DR1 has been established that is well reflected among the natural ligands, and can be well explained by the crystal structure of HLA-DR1.

MHC class I ligands and motifs

The main purposes for which this information will be useful are the prediction of T-cell epitopes within proteins of known sequences and the detailed analysis of peptide/MHC interaction. For epitope prediction it is important not to consider just the basic motif of a given MHC molecule, since the non-anchor positions of peptides could also contribute considerably to the interaction with MHC. This is evident from the preferences seen for certain residues at non-anchor-positions in pool sequencing data, from the interaction of such residues with MHC sites as seen in crystals (Madden et al. 1993; Zhang et al. 1992; Fremont et al. 1992), and from detailed binding studies showing that certain residues at a given peptide position can be detrimental for binding (Ruppert et al. 1993; Kast et al. 1994; Parker et al. 1994).

The basic approach to search a protein sequence for an epitope fitting to a given class I molecule is as follows. First, the sequence is screened for stretches fitting to the basic anchor motif (2 anchors in most cases), whereby allowance should be made for some variation in peptide length as well as in anchor occupancy. If a motif, for example, calls for 9mers with I or L at the end, 10mers with a fitting C-terminus should be considered as well, and other aliphatic residues at the C-terminus, like Val or Met, should also be considered. In this way, a list of candidates will be obtained. These are now inspected for having as many non-anchor residues as possible in common with

ligands already known, or with the residues listed among the "preferred residues" or "others" on top of each motif Table. If possible, a binding assay can be performed at this stage to exclude weak binders which occur frequently among peptides conforming to a basic motif. If a detailed study on peptide binding requirements is available, the candidates can also be screened for non-anchor residues detrimental or optimal for binding (Ruppert et al. 1993; Kast et al. 1994; Romero et al. 1991; Ebert et al. 1993). One should keep in mind, however, that pure peptide binding motifs can be misleading in the search for natural ligands, since other constraints, such as enzyme specificity during antigen processing and specificity of transporters or chaperones, are likely to contribute to ligand identity in addition to the MHC binding specificity.

A careful consideration of the pocket structure of the MHC molecule of interest can also be useful for epitope prediction (Falk and Rötzschke 1993). For the P1 residue, for example, preferences can be explained by the residues contributing to the P1 contact site (Falk et al. 1995 a,c). Since the MHC residues contributing to the different contact sites can differ among MHC molecules, such considerations should be held with caution, however (Guo et al. 1993). Computer modeling of the MHC molecule in question can be of help here.

The use of allele-specific peptide motifs is limited to a certain extent by exceptional ligands not fitting to a motif, e.g., Frumento and co-workers (1993) and Mandelboim and co-workers (1994). Such ligands will be missed by epitope predictions based on allele-specific motifs. It is not clear as yet how frequently this happens. In most cases, natural ligands will fit to the motifs, whereby substitutions of anchor residues with residues of similar chemistry (e.g., one aliphatic residue for another) and length variations are not infrequent and should be considered. A special case is the mouse H-2M3 molecule. A complete motif is not known, except that this molecule is specialized to present N-formylated peptides of bacterial or mitochondrial origin (Fischer-Lindahl 1991; Shawar et al. 1991).

MHC class II ligands and motifs

The long-awaited X-ray analysis of class II molecules has given us invaluable insight into peptide/class II interactions (Brown et al. 1993; Stern et al. 1994). Especially the detailed information on the 5 DR1-pockets accommodating anchoring side chains of one particular ligand, influenza haemagglutinin 306-318, provided a structural basis for the previously worked out peptide ligand motif of DR1 molecules (Rötzschke and Falk 1994; Sinigaglia and Hammer 1994). Moreover, pocket spacing and structure, as found for this one particular DR1/peptide complex, can be used to deduce the putative interaction for other DR1-peptide complexes and even for some other class II molecules. We found it particularly useful to evaluate pool sequencing data under the aspect of the expected pocket structure (Friede and co-workers, submitted; Schild and co-workers,

submitted). Combined with the alignment of individual class II ligands, this approach is a powerful tool to determine allele-specific class II peptide motifs, as we have exercised recently for several closely related DR4 subtypes (Friede and co-workers, submitted).

The general picture for allele-specific class II motifs emerging is as follows. A stretch of nine amino acids, on average starting at absolute positions 3 to 5 of natural ligands, is determined by the respective allele-specific motif, corresponding to the peptide portion embedded in the MHC groove. The first position of this nonamer stretch, P1, represents a hydrophobic anchor for all class II ligand motifs known so far. Anchoring of the hydrophobic P1 side chain in the respective class II pocket appears to be particularly intensive, as impressively illustrated by the deep pocket seen in the monopeptidic DR1 crystal. The importance of P1 side chains is also indicated by the striking influence of P1 on peptide binding, and by the significant clustering of hydrophobic residues at cycles 3 to 5 of self-peptide pools. In addition to P1, several other anchors follow up to P9. For DR1, these are at P4, P6, P7, and P9, as indicated by structural data, whereby the specificity of P7 is somewhat degenerate and escapes detection in binding assays or natural ligand analysis. For several other class II molecules, the same anchor spacing – P1, P4, P6, P7, P9 – is compatible with ligand motif data. DR2, DR3, and DR4 motifs as well as H-2E motifs fall into this category. Other molecules, like DR5, DPw4, and DQ7 appear to have slightly different anchor spacing, e.g., the second anchor at P3, or an anchor at P5. Allele-specific differences can occur at each of the anchor positions, although differences of P1 specificity in HLA-DR molecules are limited by the β 86Gly/Val polymorphism. More pronounced allele-specific differences are found for P4, P6, and P9, respectively. Charge differences are particularly evident; P4 of DR17, for example, requires Asp, whereas P4 of DR4Dw10 does not tolerate Asp or Glu but prefers basic or hydrophobic residues. P9, on the other hand, prefers hydrophobic residues for DR1 but negative charges for DR4Dw15 and positive charges for H-2E^k. Interestingly, charge differences in polymorphic stretches of class II molecules (probably reflecting counter charges for charged anchors) have been found to be associated with autoimmune diseases (Gregersen et al. 1987; Khalil et al. 1990; Todd et al. 1987).

Epitope prediction of class II ligands within a protein is not as easy as with class I, because the anchors, or interaction sites, are more degenerate in their specificity. The first step should be to pick out the most allele-specific anchor beyond P1, for example, P4 of DR17, P6 of DR1, or P9 of H-2E^k or DR4Dw15. The selection of nonamer sequences fitting to P1 and the other anchor of the respective motif is then further examined for adherence to the additional anchors. The resulting collection of nonamer stretches might then be inspected for adherence to the putative processing motif XPXX in the flanking regions (Rötzschke and Falk 1994). A quantitative ranking of the contribution of each amino acid residue at almost every position has been determined in an elegant approach by

Hammer and co-workers (1994) for DR4, which led to highly accurate predictions of good DR4 binders.

Technical notes

We have tried to put together all the motifs and natural ligands we were aware of. Due to the flood of data emerging in the past years, however, we anticipate that some information has been overlooked. We apologize in advance to those authors whose work was inadvertently not adequately covered.

In case of those class II ligands occurring as nested sets, we included only one or a few members of the set in many cases.

An X in peptide sequences stands for an undetermined amino acid. However, if the peptide sequence has been determined by mass spectrometry, as is the case for the peptides reported by Hunt and co-workers (1992a, b), X stands for either Leu or Ile (which have the same mass). Lowercase letters in peptide sequences indicate residue determination of lower confidence.

As far as T-cell epitopes are concerned, only those have been selected which are likely to be naturally processed;

criteria for judgement are to be found in Stevanović and Rammensee (1995). From the numerous class II motifs that have been published, we selected the more convincing ones, that is, those compatible with the class II structure. Due to the variable number of amino acids between the N-terminus and the first anchor of peptides, alignment of ligands or T-cell epitopes to class II motifs is always arbitrary, unless a structural analysis has been performed. For the class II molecules without reasonable motifs, a list of the published ligands is provided, without any attempt at alignment.

If you wish to have your motifs or ligands included in forthcoming listings, please send us reprints (no preprints) of the work describing them. We would also appreciate any comments on errors and omissions, as well as suggestions for improvements.

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Table 1 Mouse class I motifs
A H-2K^d

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues		Y				I					a
		F				L					
			V								
Preferred residues	N	P	M	K	T						
	I			F	N						
		L									
Others	K	A	A	V	H	P	H				
	A	H	E	N	I	H	E				
	R	V	S	D	M	D	K				
	S	R	D	I	Y	E	V				
	V	S	H	L	V	Q	F				
	T	F	N	S	R	S	R				
			E		T	L					
			Q		G						
			K								
			M								
			T								
Examples for ligands	T	Y	Q	R	T	R	A	L	V*	Influenza A NP 147–154	b, c
	S	Y	F	P	E	I	T	H	I	Tyrosine kinase JAK1 355–363	a, d
	K	Y	Q	A	V	T	T	T	L*	Tum-P198 14–22	a, e, f
	G	Y	K	D	G	N	E	Y	I*	Lysteriolysin O 91–99	g
	K	Y	G	V	S	V	Q	D	I*	L. monocytogenes p60 217–225	h
	G	Y	L	G	Q	V	T	X	I	Unknown	u
	S	F	V	D	T	R	T	L	L	Collagen 1 α 2 4–12	u
T-cell epitopes	L	Y	Q	N	V	G	T	Y	V	Influenza JAP HA 204–212	i
	T	Y	V	S	V	G	T	S	T	Influenza JAP HA 210–219	i
	V	Y	Q	I	L	A	I	Y	A	Influenza JAP HA 523–531	i
	I	Y	A	T	V	A	G	S	L	Influenza JAP HA 529–537	i
	T	Y	V	S	V	S	T	S	T	Influenza A HA 210–219	k, l
	I	Y	S	T	V	A	S	S	L	Influenza A HA 518–526	k
	R	Y	L	E	N	G	K	E	T	HLA-A24 170–179	a, m
	R	Y	L	K	N	G	K	E	T	HLA-Cw3 170–179	a, m
	S	Y	I	P	S	A	E	K	I	P. berghei CSP 252–260	n
	S	Y	V	P	S	A	E	Q	I	P. yoelii CSP 281–289	o
	S	Y	I	G	S	I	N	N	I	RSV M2 82–90	p
	D	Y	A	T	L	G	V	G	V	HSV-1 ICP27 448–456	q
	L	Y	R	T	F	A	G	N	P	HSV-1 ICP27 322–332	q
	T	Y	K	D	T	V	Q	L		Polio VP1 111–118	r
	F	Y	D	G	F	S	K	V	P	Polio VP1 208–217	r
	A	Y	I	S	S	G	S	S	T	Human Ig VH 49–58	l
	N	Y	D	N	A	G	T	N	L	P. falciparum CSP 39–47	s
	K	Y	L	K	K	I	K	N	S	P. falciparum CSP 333–342	s
	K	Y	L	K	I	K	H	L	L	APC frameshift	t

* Also a T-cell epitope

References:

- a: Falk et al. 1991 b; b: Rötzschke et al. 1990; c: Falk et al. 1991 a; d: Harpur et al. 1993; e: Sibille et al. 1990; f: Wallny et al. 1992; g: Palmer et al. 1991; h: Palmer 1994; i: Braciale et al. 1987; k: Kuwano et al. 1988; l: Cao et al. 1994; m: Maryanski et al. 1986; n: Romero et al. 1989; o: Weiss et al. 1990; p: Kulkarni et al. 1993; q: Banks et al. 1993; r: Kutubuddin et al. 1992; s: Blum-Tirouvanziam et al. 1994; t: Townsend et al. 1994; u: Reich et al. 1994

Table 1 (Continued)**B H-2D^d**

	Position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or auxiliary anchor residues	G	P		R		I					a, b	
				K		L						
					F							
Other preferred residues	D		N	D								
	E		I	E								
	Q		L									
Examples for ligands	K	G	P	I	T	V	Q	I		Unknown	b	
	V	G	P	Q	<u>K</u>	N	E	N	L	Unknown	b	
	S	G	P	R	<u>K</u>	X	I	X	L	Homol. mRNA CD40 mouse	b	
	A	G	P	D	<u>R</u>	T	E	K	X	L	Unknown	b
	K	G	P	D	<u>K</u>	G	N	E	F	Homol. metalloproteinase 2 inhibitor	b	
	I	G	P	E	<u>R</u>	G	H	N	L	Homol. hypoxanthine phosphoribosyl-transferase	b	
	D	G	P	V	<u>R</u>	E	H	N	L	Homol. urease canavalia ensiformis	b	
	K	G	P	E	<u>R</u>	X	N	G	L	Unknown	b	
	S	G	P	E	<u>R</u>	G	E	K	L	Homol. proliferating cell nucleolar antigen P40	b	
	D	G	P	V	<u>R</u>	G	I	S	I	Homol. ribosomal protein S17 rat	b	
	N	G	P	Q	<u>R</u>	I	Y	N	L	Unknown	b	
	S	G	P	V	A	L	V	N	F	Unknown	b	
	I	G	P	N	<u>R</u>	A	F	N	F	Unknown	b	
	S	G	P	E	<u>R</u>	L	L	S	X	Y	Homol. heterog. nucl.,RNP complex K	b
	V	G	P	S	<u>G</u>	K	Y	F	I	L	Unknown	b
	F	G	P	Y	<u>K</u>	L	N	R	L	Homol. feline leukemia virus envelope polyprotein	b	
	F	G	P	L	<u>K</u>	F	N	V	L	T	Unknown	b
	A	G	P	D	<u>R</u>	F	I	X	X	M	Unknown	b
	F	G	P	Y	<u>R</u>	F	Y	V	L	T	Unknown	b
	S	E	Q	D	L	N	F				Unknown	b
	S	X	H	K	E	Q	P	A	T		Homol. transforming protein spi-1 human	b
	S	X	P	K	T	D	X	Q	T	L	Homol. insulin receptor precursor	b
T-cell epitopes	G	P	P	H	S	N	N	F	G	Y	Tum-P35B 4-13	c
	R	G	P	G	<u>R</u>	A	F	V	T	I	HIV gp160 318-327	d, f
	L	M	G	Y	I	P	L	V	G	A	HCV core 133-142	e

References:

- a: Falk and co-workers, unpublished; b: Corr et al. 1993; c: Szikora et al. 1993; d: Takahashi et al. 1988; e: Shirai et al. 1994;
f: Bergmann et al. 1993b

Table 1 (Continued)C H-2L^d

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues		P					F				a, b, c
		S					L				
							M				
Other preferred residues		G	T	T	I	F	Q				
		Q			K		N				
		M			F						
		L									
Examples for ligands	Y	P	H	F	M	P	T	N	L*	MCMV pp 89 168–176	d
V A I T R I E Q	L	S	P	F	P	F	D	L*		OGDH 105–112	e
	L	S	P	F	P	F	D	L*		OGDH 97–112	e
	X	P	L	E	A	N	Y	Q	X F	Unknown	c
	A	P	Q	P	G	M	E	N	F	Unknown	c
	Q	P	Q	R	G	R	E	N	F	Unknown	c
	X	P	Q	P	G	R	E	Q		Unknown	c
	X	P	Q	P	N	L	Y	Q	L	Unknown	c
	X	P	A	X	A	Y	P	Y		Unknown	c
	Y	P	N	V	N	I	H	N	F	Unknown	c
	X	P	Q	K	A	G	G	F	L M	Phosphoglycerate kinase 180–189	c
T-cell epitopes	R	P	Q	A	S	G	V	Y	M	LCMV NP 118–126	f, g
	I	S	T	Q	N	H	R	A	L	Tumor antigen P91A 12–20	h
	L	P	Y	L	G	W	L	V	F	Tumor antigen P815 35–43	i
	A	P	T	A	G	A	F	F	F	JHMV Nucleocapsid 318–326	k
	Y	P	A	L	G	L	H	E	F	Measles NP 281–289	l
	T	P	H	P	A	R	I	G	L	E. coli β -gal. 876–884	m
	D	P	V	I	D	R	L	Y	L	Measles HA 343–351	n
	S	P	G	R	S	F	S	Y	F	Measles HA 544–552	n

* Also a T-cell epitope

References:

- a: Falk et al. 1991 b; b: Falk and co-workers, unpublished; c: Corr et al. 1992; d: Reddehase et al. 1989; e: Udaka et al. 1992; Udaka et al. 1993; f: Whitton et al. 1989; g: Schulz et al. 1991; h: Lurquin et al. 1989; i: Lethé et al. 1992; k: Bergmann et al. 1993 a; l: Beauverger et al. 1993; m: Gavin et al. 1994; n: Beauverger et al. 1994

Table 1 (Continued)**D H-2K^b**

	Position								Source	Ref.
	1	2	<u>3</u>	4	5	6	7	8		
Anchor or auxiliary anchor residues			Y		F		L			a
				Y		M				
					I					
					V					
Other preferred residues	R	N	P	R		T	N			
	I			D		I	Q			
	L			E		E	K			
	S			K		S				
	A			T						
Examples for ligands	R	G	<u>Y</u>	V	<u>Y</u>	Q	G	L*	VSV NP 52–59	b
	S	I	<u>I</u>	N	F	E	K	L*	Ovalbumin 258–276	a, c, d
	H	I	<u>Y</u>	E	F	P	Q	L	Unknown	n
T-cell epitopes	I	I	<u>Y</u>	R	F	L	L	I	Rotavirus VP7 33–40	e
	S	S	<u>I</u>	E	F	A	R	L	HSV glycoprotein B 498–505	f
	F	A	P	G	N	<u>Y</u>	P	A	Sendai virus NP 324–332	g, h
		K	S	P	W	F	T	T	MuLV p15E 574–581	i, k
		V	G	P	V	F	P	P	Rotavirus VP6 376–384	l
		Y	S	G	<u>Y</u>	I	F	R	Rotavirus VP3 585–593	l
			F	E	Q	N	T	A	MUT 2 tumor antigen	m
			F	E	Q	N	T	A	MUT 1 tumor antigen	m
						Q	P ⁺			

* Also a T-cell epitope

+ One of these peptides was found in a total cell extract of K^b-expressing tumor cells

References:

- a: Falk et al. 1991 b; b: van Bleek and Nathenson 1990; c: Rötzschke et al. 1991; d: Carbone et al. 1988; e: Franco et al. 1993; f: Bonneau et al. 1993; g: Kast et al. 1991; h: Schumacher et al. 1991; i: Sijts et al. 1994; k: White et al. 1994; l: Franco et al. 1994; m: Mandelboim et al. 1994; n: Wallny 1992

Table 1 (Continued)

E H-2Db

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues	N M I										a
Preferred residues	M	I	K		L						
		L	E		F						
		P	Q								
		V	V								
Others	A	A	G	D	A	D	F				
	N	Q		T	Y	E	H				
	I	D			T	Q	K				
	F				V	V	S				
	P				M	T	Y				
	S				E	Y					
	T				Q						
	V				H						
					I						
					K						
					P						
					S						
Examples for ligands	A	S	N	E	N	M	E	T	M*	Influenza A34 NP 366–374	a, b, c
	I	Q	V	G	N	T	R	T	I*	Yersinia YOP 51 249–257	n
T-cell epitopes	A	S	N	E	N	M	D	A	M	Influenza A68 NP 366–374	d
	S	A	I	N	N	Y	A	Q	K	SV 40 T 206–215	e, o
	C	K	G	V	N	K	E	Y	L	SV 40 T 223–231	e, o
	Q	G	I	N	N	L	D	N	L	SV 40 T 489–497	e, o
	S	G	P	S	N	T	P	P	E	Adenovirus 5 E1A 234–243	f
	F	Q	P	Q	N	G	Q	F	I	LCMV NP 396–404	g
	S	G	V	E	N	P	G	G	Y	LCMV GP 276–286	h
	K	A	V	Y	N	F	A	T	C	LCMV GP 33–42	i, k
	R	A	H	Y	N	I	V	T	F	HPV16 E7 49–57	l
	N	N	L	D	N	L	R	D	Y (L)	SV 40 T 492–500 (501)	m

* Also a T-cell epitope

References:

- a: Falk et al. 1991b; b: Rötzschke et al. 1990; c: Townsend et al. 1986; d: Cerundolo et al. 1991; e: Deckhut et al. 1992; f: Kast et al. 1989; g: Yanagi et al. 1992; h: Oldstone et al. 1988; i: Oldstone et al. 1993; k: Klavinskis et al. 1990; l: Feltkamp et al. 1993; m: Alsheikly 1994; n: Starnbach and Bevan 1994; o: Tevethia et al. 1990

Table 1 (Continued)**F H-2K^k**

	Position									Comments	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor residues		E			I	I				C-terminus at P8 or P9	a, b, c	
Preferred residues	V	D	K	L	A	N	T					
	F		N		G	K						
			Y		P	H						
			M		T							
			Q		V							
			I		F							
			L		S							
			F									
			P									
			H									
			T									
Source												
Examples for natural ligands	H	E	T	T	F	N	S	I		β Actin 275–282	k	
	D	D	H	R	A	G	K	I		S24 ribosomal protein 53–60	k	
	Y	E	D	T	G	K	T	I		Unknown	k	
	K	E	M	K	A	K	V	I		Homol. T cell transcript. factor 1	k	
	E	E	E	P	V	K	K	I		Hn RNP C protein 84–91	k	
	S	E	I	V	G	K	R	I		S7/S8 ribos. protein 137–144	k	
	S	E	G	G	S	H	T	I		H-2D ^k 112–119	k	
	D	E	R	T	V	R	K	I		Unknown	k	
	E	E	D	P	V	K	K	V		CArG bind. factor A 209–216	k	
	E	A	Y	L	G	K	K	V		BiP 158–165	k	
T-cell epitopes	F	E	A	N	G	N	L	I		Influenza A HA 259–266	c, i	
	I	E	G	G	W	T	G	M	I	Influenza A HA 10–18	c, i	
	S	D	Y	E	G	R	L	I		Influenza A NP 50–57	d, l	
	F	E	S	T	G	N	L	I		Influenza JAP HA 255–262	e	
	S	E	F	L	L	E	K	R	I	SV 40 T 560–568	f	
	Y	E	N	D	I	E	K	K	I	P. falciparum CSP 375–383	g	
	D	E	L	D	Y	E	N	D	I	P. falciparum CSP 371–379	g	
	T	E	M	E	K	E	G	K	I	HIV-1 RT 206–214	h	
	V	E	A	E	I	A	H	Q	I	Rabies NS 197–205	i	
	E	E	G	A	I	V	G	E	I	Influenza A NSI 152–160	a	

References:

a: Cossins et al. 1993; b: Norda et al. 1993; c: Gould et al. 1991; d: Bastin et al. 1987; e: Sweetser et al. 1989; f: Rawie et al. 1988; g: Kumar et al. 1988; h: Hosmalin et al. 1990; i: Larson et al. 1991; Gould et al. 1987; k: Brown et al. 1994; l: Gould et al. 1989

G H-2K^{km1}

	Position								Source	Ref.
	1	<u>2</u>	3	4	5	6	7	8		
Anchor or auxiliary anchor residues		E			I					a
Other preferred residues	Q	K	P	A		R				
	G	N		R		Y				
	P	Q		K						
			G							
			M							
			P							
			Y							

References:

a: Norda et al. 1993

Table 1 (Continued)

H Qa-2

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	M	N		V	K	H		L			a, b
	L	I		I	M			I			
	Q	L			I			F			
Other preferred residues	K	T	P	L	L	R	E				
	A	E	T	F			Q				
	E	A	E	N			N				
	Q	G	H	Y		D					
			K	M			K				
			S	F			S				
			D	Y		T					
						R					
Examples for ligands	K	Q	N	P	I	A	H	Q	L	Unknown	b
	A	L	A	E	L	P	H	E	L	Unknown	b
	K	Q	N	P	T	V	H	H	L	Unknown	b
	A	G	L	L	G	M	R	S	G	Unknown	b
	K	L	I	K	V	Y	H	S	L	Unknown	b
	V	Q	N	X	T	M	H	P	I	Unknown	b
	R	S	N	G	Q	V	H	M	L	Unknown	b
	K	L	T	G	I	K	H	E	L	Cofilin 127–135	b
	A	M	L	A	T	Y	H	K	L	Unknown	b
	G	Q	L	X	V	X	H	K	L	Unknown	b
	I	L	M	E	H	I	H	K	L	Ribosomal protein L19 137–145	b
	A	Q	N	P	V	L	Y	Q	I	Unknown	b
	S	L	I	K	T	V	R	E	M	Unknown	b
	K	M	L	P	V	E	H	N	L	Unknown	b
	K	Q	L	I	V	T	Y	H	L	Unknown	b
	S	V	L	D	D	L	A	L	L	Unknown	b
	Y	E	S	P	L	M	X	F	L	Unknown	b
	D	L	L	G	T	L	H	N	L	Unknown	b

References:

a: Rötzscheck et al. 1993; b: Joyce et al. 1994

I Selected other T-cell epitopes

MHC	Sequence											Comments	Ref.
H-2D ^k	R	R	K	G	K	Y	T	G	L			T cell epitope of LEC-A	a
H-2M3	fM	F	F	I	N	I	L	T	L	L	V	ND1 α 1–17	b
	fM	F	F	I	N	A	L	T	L	L	V	ND1 β 1–17	b

References:

a: de Bergeyck et al. 1994; b: Fischer Lindahl 1991

Table 2 HLA-A motifs
A HLA-A1

	Position									Source	Ref		
	1	2	3	4	5	6	7	8	9				
Anchor or auxiliary anchor residues	T	D	P		L		Y				a, b, c, f, i		
	S	E											
Other preferred residues	L		G	G	G								
		I	N	V									
		Y	I										
Examples for ligands	A	T	D	F	K	F	A	M	Y	Cyclin-like protein 135–143	a, i		
	I	A	D	M	G	H	L	K	Y	Proliferation cell nuclear antigen 241–249	a, b, i		
	M	I	E	P	R	T	L	Q	Y	Ribosomal protein S16 40–48	a, b		
	Y	T	S	D	Y	F	I	S	Y	Ets-1 154–162	a, i		
	L	T	D	P	G	V	L	D	Y	Unknown	a		
	V	S	D	I	V	G	P	D	G	Fibrillarin 177–188	a, b		
	Y	T	D	Y	G	G	L	I	F	Cytochrome C oxidase II	a, i		
	Q	S	E	D	G	S	H	T	I	HLA class I α chain 111–123	a		
	Y	L	D	D	P	D	L	K	Y	Cytosine methyl transferase 238–246	i		
	S	T	D	H	I	P	I	L	Y	Fructose-6-amino transferase 217–225	i		
	D	S	D	G	S	F	F	L	Y	IgG4 279–287	i		
	G	T	D	E	X	R	N	X	Y	Unknown	i		
	V	S	D	P	Y	N	X	K	Y	Unknown	d, i		
	V	A	D	K	V	H	X	M	Y	Unknown	i		
	Y	T	A	V	V	P	L	V	Y	J-chain 102–110	i		
	Y	T	N	P	Q	F	N	V	Y	Unknown	i		
	E	T	X	X	P	D	W	S	Y	Unknown	i		
	F	T	D	V	N	S	X	X	R	Unknown	i		
	S	S	E	Q	T	F	M	Y	Y	Ornithine decarboxylase 309–317	b		
	S	T	E	P	V	N	I	L	Y	Unknown	b		
	G	T	D	P	G	V	L	I	Y	Unknown	b		
	S	T	E	P	P	M	L	N	Y	Unknown	b		
	S	L	E	P	Q	R	T	Q	Y	Unknown	b		
	F	T	E	V	S	I	R	K	Y	Unknown	b		
	K	F	D	P	V	N	L	V	Y	Unknown	b		
	A	V	D	P	G	G	M	Y	S	Unknown	b		
	F	G	S	G	A	R	D	X	Y	Unknown	b		
	Y	X	E	P	Q	F	L	T	Y	Unknown	b		
	A	X	I	P	A	F	I	N	Y	Unknown	b		
	I	T	E	D	M	G	H	L	K	Y	Unknown	f	
	E	T	D	X	X	X	D	R	S	E	Y	Unknown	i
T-cell epitopes	E	A	D	P	T	G	H	S	Y	MAGE-1 161–169	e, k		
	V	S	D	G	G	P	N	L	Y	Influenza A PB1 591–599	b, f		
	C	T	E	L	K	L	S	D	Y	Influenza A NP 44–52	f		
	E	V	D	P	I	G	H	L	Y	MAGE-3	g, h		

References:

a: Falk et al. 1994 c; b: Di Brino et al. 1993 b; c: Sette et al. 1994; d: Engelhard 1994; e: Traversari et al. 1992; f: DiBrino et al. 1994; g: Gaugler et al. 1994; h: Celis et al. 1994; i: Kubo et al. 1994; k: Van der Bruggen et al. 1991

Table 2 (Continued)
B HLA-A*0201

* Class I ligands allocated to A2 by motif. + Also a T-cell epitope

References:

- a: Falk et al. 1991b; b: Hunt et al. 1992; c: Henderson et al. 1992; d: Wei and Cresswell 1992; e: Henderson et al. 1993; f: Wölfel et al. 1994; g: Robbins et al. 1994; h: Brichard et al. 1993; i: Engelhard et al. 1993; j: Walker et al. 1989; k: Gotch et al. 1988; l: Harris et al. 1993; m: Nayersina et al. 1993; n: Bertoletti et al. 1993, 1994; o: Utz et al. 1992; p: Lee et al. 1993; q: Robbins et al. 1989; r: Chisari and co-workers, personal comm.; s: Shirai et al. 1994; t: Tarpey et al. 1994; u: Cox et al. 1994; v: Kawakami et al. 1994b; w: Coulie et al. 1994; x: Kawakami et al. 1994a, c; y: Falk et al. 1994a; z: Bednarek et al. 1991

Table 2 (Continued)

C HLA-A*0205

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	V				I			L			a
	L				V						
	I				L						
	M				A						
Other preferred residues	Q	Y	G	Y	T	Q	K				
	P	E	V								
	F	D	L								
	I	K	I								
	N										

References:

a: Rötzschke et al. 1992

D HLA-A3

	Position										Source	Ref.
	1	2	3	4	5	6	7	8	9	10		
Anchor or auxiliary anchor residues	L	F			I	I		K	K			a, b, g
	V	Y			M	L		Y				
	M				F	M		F				
					V	F						
					L							
Other preferred residues	I			I	T		Q					
				P		S						
			V			T						
			K			K						
Examples for ligands	K	X	F	K	M	I	L	R	K		Unknown	a
	K	L	F	K	N	I	L	Y	K		Unknown	a
	Y	L	X	V	R	X	A	X	i	V	Unknown	a
	K	L	H	K	Q	R	A	K	S		Unknown	a
	S	L	F	K	Q	V	V	T	K		Unknown	a
	K	X	F	V	K	X	L	X	Y		Unknown	a
	S	L	F	N	T	H	L	X	K		Unknown	a
	T	L	A	N	D	X	V	V	P		Unknown	a
	G	I	F	A	X	X	X	V	K	A	Unknown	a
	T	X	F	V	K	X	L	X	Y		Unknown	a
	S	L	F	D	H	I	L	X	K	H	Unknown	a
	K	L	F	K	V	V	X	N	Y		Unknown	a
	K	L	Y	E	K	V	Y	T	Y	K	Unknown	a
	K	L	F	N	I	M	V	T	Y		Unknown	a
	K	L	F	E	K	V	Y	N	Y		Unknown	a
	K	L	F	K	V	T	F	S	Y		Unknown	a
	G	L	F	P	X	Q	F	A	Y		Unknown	a
	S	L	F	E	L	V	F	X	Y		Unknown	a
	S	L	X	E	K	T	F	D	Y		Unknown	a
	S	L	H	K	Y	X	f	e	Y		Unknown	a
	K	M	F	N	I	T	v	T	Y		Unknown	a
	K	L	F	V	K	V	y	N	Y		Unknown	a
	K	I	V	R	K	P	G	M	A		Unknown	a
T-cell epitopes	R	L	R	D	L	L	I	V	T	R	HIV-1 env gp41 768–778	c
	Q	V	P	L	R	P	M	T	Y	K	HIV-1 nef 73–82	d
	T	V	Y	Y	G	V	P	V	W	K	HIV-1 env gp120 36–45	e
	R	L	R	P	G	G	K	K	K		HIV-1 gag p17 20–29	e
	I	L	R	G	S	V	A	H	K		Influenza NP 265–273	f

References:

a: DiBrino et al. 1993 a; b: Maier et al. 1994; c: Takahashi et al. 1991; d: Koenig et al. 1990; e: Venet and Walker 1993; f: DiBrino et al. 1993 b; g: Kubo et al. 1994

Table 2 (Continued)**E HLA-A*1101**

	Position											Source	Ref.
	1	2	3	4	5	6	7	8	9	10	11		
Anchor or auxiliary anchor residues	V	M				L		K	K	K			a, b, c
	I	L				I							
	F	F				Y							
	Y	Y				V							
	I					F							
	A												
Other preferred residues	A	T	N	P	P	I		R	R	R	R		
			D	G	I	V		K	D				
			E	D	F	M		N					
			Q	E	V		E						
				K	M		Q						
Examples for ligands	A	V	M	K	P	E	A	E	K	R	K	Unknown	b
	A	V	I	L	P	P	L	S	P	Y	F	HSB 66 EST 18–29	b
	A	S	F	D	K	A	K	L	K	K		Thymosin β -10 11–20	b
	G	Q	Y	G	N	P	L	N	K			Cattle metalloproteinase 19–27	b
	G	V	M	P	S	H	F	S	R			Ribosomal protein S19 93–101	b
	Y	F	D	P	A	N	G	K	F	S	K	Elongation factor 2 265–275	b
	A	T	A	G	D	G	L	I	E	L	R	Prohibitin (rat) 229–240	b, c
	S	T	Y	Y	G	S	F	V	T	R		Unknown (also presented by A33)	a, b
	S	V	L	N	L	V	I	V	K			Ribosomal protein S6 107–115	c
	K	V	V	N	P	L	F	E	K			Ribosomal protein L7A 25–33	c
	R	T	Q	N	V	L	G	E	K			Ribosomal protein S3 54–62	c
	G	T	M	T	T	S	X	Y	K			Unknown	c
	A	S	F	D	K	A	K	L	K			Thymosin β -10 11–19	c
	A	A	M	X	D	T	V	V	F	K		Unknown	c
	R	V	E	Q	A	V	E	S	M	V	K	Unknown	c
T-cell epitope	I	V	T	D	F	S	V	I	K			EBNA 4 416–424	a, d

References:

a: Zhang et al. 1993; b: Falk et al. 1994; c: Kubo et al. 1994; d: Gavioli et al. 1993

F HLA-A24

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	Y			I	F		I				a
				V			L				
				F							
Other preferred residues	N	D			Q	E					
	E	P			N	K					
	L										
	M										
	P										
	G										
Examples for ligands	K	Y	P	E	N	F	F	L	L	Protein phosphatase 1 113–121	b
	Y	Y	E	E	Q	H	P	E	L	NK/T-cell activation protein 107–115	b
	A	Y	V	H	M	V	T	H	F	Unknown	b
	V	Y	X	K	H	P	V	S	X	Unknown	b
T-cell epitope	R	Y	L	K	D	Q	Q	L	L	HIV gp 41 583–591	c

References:

a: Maier et al. 1994; b: Kubo et al. 1994; c: Dai et al. 1992

Table 2 (Continued)**G HLA-A*3101**

	Position									Comments	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	L	F			L		R				a
	V	L			F						
	Y	Y			V						
	F	W			I						
Other preferred residues	K	T	K	P	P	T	N	L		P1 deduced from individual ligands	
	R	Q	N	D	I	N	V	R			
				E	V	D	R	N			
				G	F	E	F	Q			
				S	L	R	T				
				V	Y		H				
				T	W		L				
							Y				
Source											
Examples for ligands	L	Q	F	P	V	G	R	V	H	R	
	Q	Q	L	Y	W	S	H	P	R		a
	R	G	Y	R	P	R	F	R	R		a
	K	V	F	G	P	I	H	E	R		a
	K	I	M	K	W	N	Y	E	R		[GlcNac]-P-transferase 371–379
	R	Y	M	D	A	W	N	T	Y	S	Unknown
											Lamin B2
T-cell epitope	S	T	L	P	E	T	T	V	V	R	R
											Hepatitis B cAg 141–151
References:											
a: Falk et al. 1994 c; b: Missale et al. 1993											

H HLA-A*3302

	Position									Comments	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	A				R						a
	I										
	L										
	F										
	Y										
	V										
Preferred residues	D	T	L	P	P	I				P1 deduced from individual ligands	
	E		K		L						
				F							
Other possible residues	M	Q	R	R	R	H	Q				
		W	D	I	D	Y	N				
		E	E	F	H	V	E				
		N	G	P	Y	T	M				
		S	V		S						
		H	L								
		P	W								
Source											
Examples for ligands	D	M	A	A	Q	I	T	Q	R	HLA class I α -chain 161–169	
	E	S	G	P	S	I	V	H	R	Actin 364–372	a
	T	Y	Y	G	S	F	V	T	R	Unknown	a
	D	Y	I	H	I	R	I	Q	R	Human cDNA HSB15F 102 65–74	a
	E	I	M	K	W	N	R	E	R	Unknown	a
	T	I	M	P	K	D	I	Q	L	Histon 3.1/3.3 118–129	a
T-cell epitope	I	V	G	L	N	K	I	V	R	HIV p24 gag 267–275	b, c

References:

a: Falk et al. 1994 c; b: Buseyne et al. 1993; c: Buseyne and Riviere 1993

Table 2 (Continued)**I HLA-A68.1**

	Position									Source	Ref.		
	1	2	3	4	5	6	7	8	9				
Anchor residues	V R T K										a		
Examples for ligands	A	V	A	A	V	A	A	R	R	Unknown	a		
	E	V	A	P	P	E	Y	H	R	Unknown	a		
	E	V	A	P	P	E	Y	H	R	Unknown	a		
	D	V	F	R	D	P	A	L	K	Homologous ribosomal 60S	a		
	K	T	G	G	P	I	Y	K	R *	Influenza NP 91–99	a, b		
	E	V	I	L	I	D	P	F	H	K	Unknown	a	
	T	V	F	D	A	K	R	L	I	G	R	a	
	X	V	L	K	X	I	A	K	R *	Unknown	d		
	P	V	K	Q	V	V	Y	H	R *	Unknown	d		
	E	S	G	P	S	I	V	H	R	K *	β-Actin 364–373	d	
	T	T	X	T	T	N	A	R *		Unknown	d		
	D	T	T	P	T	X	X	R *		Unknown	d		
T-cell epitopes	S	T	L	P	E	T	T	V	V	R	R	Hepatitis B cAg 141–151	c

* Class I ligands allocated to A68.1 by motif +Also a T-cell epitope

References:

a: Guo et al. 1992; b: Silver et al. 1992; c: Missale et al. 1993; d: Harris et al. 1993

Table 3 HLA-B motifs**A HLA-B7**

	Position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or auxiliary anchor residues	P R L F										a, b	
Other preferred residues	D D F L G P T											
Also detected	A	D	E	I	R	V						
	H	E	H	V	L							
	S	Q	L		I							
		K	K									
		Y	S									
		F	T									
		M	P									
		N										
		A										
Examples for ligands	A	P	<u>R</u>	T	V	A	L	T	A	HLA-DP signal sequence 9–17	a	
	A	P	<u>R</u>	T	V	A	L	T	A	HLA-DP signal sequence 9–18	a	
	A	P	<u>R</u>	A	X	X	X	X	X	Unknown	a	
	A	P	<u>R</u>	X	P	X	T	G	X	Unknown	a	
	A	P	<u>R</u>	A	S	R	P	S	X	Unknown	a	
	A	P	<u>R</u>	T	L	V	L	L	L	HLA-A2.1 signal sequence 5–13	a	
	M	P	<u>R</u>	G	V	V	V	T	X	Unknown	a	
	S	P	<u>R</u>	Y	I	F	T	M	L	Topoisomerase II 801–809	a	
	A	P	A	P	T	V	A	V	X	Unknown	a	
	R	P	S	G	P	G	P	E	X	Unknown	a	
	L	V	M	A	P	R	T	V	L	HLA-B7 signal sequence 2–10	a	
	R	V	M	A	P	R	A	X	X	Unknown	a	
	A	P	<u>R</u>	A	F	X	P	X	P	Unknown	a	
	A	A	S	K	E	R	S	G	V	S	Histone H1 49–59	a
	A	P	<u>R</u>	S	N	G	M	V	X	Unknown	c	
	A	P	<u>R</u>	Q	P	G	X	M	A	Unknown	c	
	A	P	A	P	P	K	p	M	M	Ribosomal S26 protein 107–115	c	
	A	P	Y	G	G	P	X	A	X	Unknown	c	
T-cell epitope	T	P	G	P	G	V	R	Y	P	L	HIV-1 nef 128–137	d

References:

a: Huczko et al. 1993; b: Maier et al. 1994; c: Engelhard 1994; d: Culmann et al. 1991

Table 3 (Continued)**B HLA-B8**

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues			K		K			L			a, b, d
					R						
Other preferred residues	G		R	E		N	E	E	F		
	L		Q		Q	H	Q		M		
	I		D		H	M	H				
			H		I	N	S				
			L		L	D	L				
			S		Y	Q	V				
			T		V	S	D				
			R		E	T	T				
			G		M	Y					
			K		S						
					T						
					F						
Examples for ligands	H	P	K	Y	K	T	E	L		Tristetraproline 148–155	d
	I	L	K	Q	K	I	A	d	I	IL-6 precursor 161–169	d
	E	L	K	V	K	N	l	e	I	Restin 1273–1281	d
	E	P	K	Y	K	T	Q	L		Yeast PRAI-SCS 95–102	d
	V	P	K	L	K	V	X	A	L	Rat ribosomal prot. L18, 94–102	d
	F	A	K	P	R	V	G	G		Unknown	d
	S	P	K	L	K	Y	M	Q		Unknown	d
	E	L	K	K	K	T	N	I		Unknown	d
	S	P	K	E	K	I	X	Y		Unknown	d
	X	A	K	E	K	L	A	D		Unknown	d
	S	P	K	E	K	Y	E	X		Unknown	d
	E	L	K	E	K	T	Q	L		Unknown	d
	L	P	K	V	K	L	A	L		Unknown	d
	H	p	K	Y	K	T	E	L		Unknown	d
	V	L	D	L	K	I	V	A	F	Unknown	d
	G	P	K	E	K	X	A	M		Unknown	d
	G	L	K	V	K	G	N	E	F	Unknown	d
	F	L	K	P	K	F	V	A	L	Unknown	d
	E	L	K	I	K	V	Y	X	I	Unknown	d
	S	L	K	E	K	V	X	L		Unknown	d
	E	L	K	E	K	X	y	e	I	Unknown	d
	I	P	K	L	K	N	V	K	r	Unknown	d
	S	L	K	I	K	X	L			Unknown	d
	D	L	K	Q	K	N	E	L		Unknown	d
T-cell epitopes	E	L	R	S	R	Y	W	A	I	Influenza NP 380–388	b
	F	L	R	G	R	A	Y	G	L	EBNA 3 339–347	c
	E	I	Y	K	R	W	I	I	L	HIV gag p24, 262–270	d, e
	G	E	I	Y	K	R	W	I	I	HIV gag p24, 261–269	d, e
	E	I	K	D	T	K	E	A	L	HIV gag p17, 93–101	d, f

References:

a: Malcherek et al. 1993; b: Sutton et al. 1993; c: Burrows et al. 1990; d: DiBrino et al. 1994; e: Phillips et al. 1989; f: Achour et al. 1990

Table 3 (Continued)
C HLA-B*2702

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues		R				F					a
						Y					
						I					
						L					
						W					
Other preferred residues	K	F	G	I	I	Y	K				
		L	P	K	V	L	V				
	X	K	E	Y	V	D					
		D	V	R	T	E					
		E	M	D	F	R					
		Q	T	H							
		T		E							
		S		Q							
Examples for ligands	S	R	D	K	T	I	I	M	W	HGNBPβ-subunit 35–43	a
	G	R	L	T	K	H	T	K	F	Rat ribosomal protein L36 36–44	a
	R	R	F	V	N	V	V	P	T	Human fau protein 114–123	a
	K	R	Y	K	S	I	V	K	Y	HFPS 191–199	a
	K	R	K	K	A	Y	A	D	F	Cytochrome C oxidase 42–50	a
	K	R	G	I	L	T	L	K	Y	Actin 63–71	a
	G	R	F	G	V	G	N	R	Y	Unknown	a
	G	R	F	K	L	I	V	L	Y	Unknown	a

References:

a: Rötzschke et al. 1994

Table 3 (Continued)
D HLA-B*2705

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues	R									L F	a, b
Other preferred residues	A	L	K	I	I	I	K	Y			
	G	I	Q	V	A	T	N	M			
	K	F	E	L	N	Y	R	I			
	R		G	P	Q	M	E	R			
			G	D	L	Q	H				
				V	W		K				
					K	N					
					V						
					P						
Examples for ligands	G	R	L	T	K	H	T	K	F	Rat ribosomal protein L 36 36–44	b
	A	R	L	F	G	I	R	A	K	HBBCP 190–198	a, b
	R	R	L	G	X	Q	Y	R	R	Unknown	b
	R	R	F	G	D	K	L	N	F	Immediate early response gene 87–95	b
	G	R	v	F	I	I	K	E		Homologous to IL-1 receptor antagonist	b
	G	R	X	X	i	f	X	r	f	Unknown	b
	T	R	Y	P	I	L	A	G	H	Cytochrome P450 20–28	b
	r	r	i	s	g	V	D	R	Y	Unknown	a, b
	K	R	F	S	F	K	K	S	F	Cattle MARCKS 155–163	b
	A	R	L	Q	T	A	L	L		Rat core histone 188–196	b
	R	R	X	P	I	F	S	R	L	TIS 11B protein 325–333 (X = L)	b
	R	R	F	M	q	Y	Y	V	Y	Homologous to proteasome subunit C5 127–135	b
	R	R	I	K	E	I	V	K	K	HSP 86 200–209	b
	R	R	M	G	P	P	V	G	G	Ribonucleoprotein L 312–322	b
	R	R	I	K	E	I	V	K	K	HSP 89 a 200–208	a
	G	R	I	D	K	P	I	L	K	Ribosomal protein L8 173–181	a
	R	R	S	K	E	I	T	V	R	ATP-dependent RNA helicase 77–85	a
	K	R	F	E	G	L	T	Q	R	Unknown	a
	R	R	V	K	E	V	V	K	k	HSP 89 B 195–203	a
	F	R	Y	N	G	L	I	H	R	60 S ribosomal protein L28 37–45	a
	R	R	Y	Q	K	S	T	E	L	Histon H3.3 52–60	a
	R	R	W	L	P	A	G	d	a	Elongation factor 2 341–349	a
	R	R	I	S	G	V	D	R	Y	Unknown	a
	R	R	F	T	R	P	E	H		Unknown	a
	K	K	Y	Q	K	S	T	E	L	Unknown	c
	XXLNSQDQQCDSSLVE									Draft-1 protooncogene 1–16	i
B*270x-restricted T-cell epitopes	S	R	Y	W	A	I	R	T	R	Influenza NP 383–391	d
	R	R	R	W	R	R	L	T	V^x	EBNA LMP2 236–244	e
	R	R	Y	P	D	A	V	Y	L	Measles F protein 438–446	f
	R	R	I	Y	D	L	I	E	L	EBNA 3C 258–266	e
	K	R	W	I	I	L	G	L	N	HIV-1 gag p24 265–274	d, g
	G	R	A	F	V	T	I	G	K	HIV-1 gp120 314–322	d
	R	R	K	A	M	F	E	D	I	HSP 60 284–292	h

* B*2704-restricted

References:

- a: Jardetzky et al. 1991; b: Rötzschke et al. 1994; c: Shepherd et al. 1993; d: Huet et al. 1990; e: Brooks et al. 1993; f: van Binnendijk et al. 1993; g: Buseyne et al. 1993; h: Cerrone et al. 1991; i: Frumento et al. 1993

Table 3 (Continued)**E HLA-B*3501**

	Position										Source	Ref.
	1	<u>2</u>	3	4	5	6	7	8	9	10		
Anchor or auxiliary anchor residues		P						Y	Y			a, b
								F				
								M				
								L				
								I				
Other preferred residues	M	A	I	K	D	I	V	E				
	V	L	D	I	Q	N	Q					
	Y	F	E	V	K	E	V					
	R	V	G	T	V	Q	T					
	D	M	P	E	L	T						
		E		G	M	K						
		T		L								
		Y		M								
		N										
T-cell epitopes	K	P	K	D	E	L	D	Y			P. falciparum CSP 368–375	a
	K	S	K	D	E	L	D	Y			P. falciparum CSP 368–375	a
	K	P	N	D	K	S	L	Y			P. falciparum LS 1850–1857	a
	A	S	R	C	W	V	A	M			HCV E1 235–242	c

References:

a: Hill et al. 1992; b: Falk et al. 1993 b; c: Koziel et al. 1992

F HLA-B*3701

	Position									Source	Ref.
	1	<u>2</u>	3	4	<u>5</u>	6	7	8	9		
Anchor or auxiliary anchor residues		D			V			F	I		a
		E			I			M	L		
								L			
Other preferred residues	K	H			T		Q	T			
	Q	P			R		K	E			
		G			A		Y	N			
		S			D		L	D			
		L			G		Q				
					H			G			
					M			H			
T-cell epitope	E	D	L	R	<u>V</u>	L	S	F	I	Influenza NP 339–347	b

References:

a: Falk et al. 1993 b; b: Townsend et al. 1986

G HLA-B*3801

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues		H	D				F				a
			E				L				
Other preferred residues	I	F	I	G	M	V	Y	K	I		
	P	A	E	T	I	V	Y				
	W	S	P	V	T	N	N				
	Y	N	L	A	K		R				
	M	V	E	R			T				
	V		G	N							
			L	H							
			K								
			S								
Examples for ligands	E	<u>H</u>	A	G	V	I	S	V	L	Unknown	a
	T	<u>H</u>	D	E	L	E	D	K	L	Unknown	a
	Q	<u>Y</u>	D	E	A	V	A	Q	F	Histone binding protein 627–635	a
	Y	P	D	P	A	N	G	K	F	Elongation factor 2 265–273	a
	S	<u>H</u>	I	G	D	A	V	V		Cyclin 152–159	a
	Y	<u>H</u>	E	D	I	H	T	Y	L	Cyclin A 178–186	a
	T	F	<u>D</u>	V	A	P	S	R	L	Pm5 protein 270–278	a

References:

a: Falk et al. 1995b

H HLA-B*39011

	Position									Source	Ref.
	1	2	3	4	5	<u>6</u>	7	8	9		
Anchor or auxiliary anchor residues	R				I			L			a
	H				V						
					L						
Other preferred residues	A	D	V	N	N	S	V				
	D	E	Y		Y	K	I				
	I	G	I		F	R	M				
	L	P	L			E					
	F	K	F			T					
	V		T								
	M		G								
	S		K								
	T		N								
	Y		P								
Examples for ligands	S	<u>H</u>	I	G	D	A	V	V		Cyclin 152–159	a
	I	<u>H</u>	E	P	E	P	H	I		CKShs1 protein 59–66	a
	S	R	D	K	T	<u>I</u>	I	M		GBLP 35–42	a

References:

a: Falk et al. 1995b

Table 3 (Continued)**I HLA-B*3902**

	Position									Ref.
	1	2	3	4	5	6	7	8	9	
Anchor or auxiliary anchor residues	K		I			L				a
	Q		L							
		F								
		V								
Other preferred residues	K	A	G	N	V	V	T	F		
	A	I	P	E	Y	L	S	M		
		F	G	T	T	R				
		V	P	H	Y					
		N	Q	F	N					
		L	S	I	D					
		T	T	M	H					
		Y		P						
		E		R						
		H								
		S								

References:

a: Falk et al. 1995 b

K HLA-B40*

	Position											Source	Ref.	
	1	2	3	4	5	6	7	8	9	10	11			
Anchor or auxiliary anchor residues	E	F				L							a	
	I					W								
	V					M								
						A								
Examples for ligands	T	E	F	P	K	E	R	H	L	R	L	Unknown	a	
	G	E	F	P	N	K	N	X	L			Unknown	a	
	G	E	F	P	N	K	N	X	L	Y	A	Unknown	a	
	G	E	F	P	G	K	I	F	L	Y	A	Unknown	a	
	W	E	F	L	Q	P	I	L	L			Unknown	a	
	G	E	F	I	P	G	N	D	L	H	R	Unknown	a	
	G	E	F	P	P	X	D	N	W			Unknown	a	
	E	E	F	Y	V	D	L	E	R			HLA-DQ α 33-41	a	
	N	E	F	P	D	I	D	I	R			Unknown	a	
	A	E	F	P	K	X	E	A	R			Unknown	a	
	A	E	I	G	E	V	I	V	L	W	X	W	Unknown	a
	A	E	I	P	G	E	I	A	L			Unknown	a	
	G	E	I	L	D	V	F	D	A			IRE-BP 695-703	a	
	F	E	I	P	X	L	D	V	A			Unknown	a	
	D	E	V	T	P	Q	P	Q	L	V		Unknown	a	
	K	E	V	G	V	D	V	A	L	Y	A	Unknown	a	
	K	E	S	T	L	H	L	V	L			Ubiquitin 63-71	a	
	G	E	V	D	V	E	Q	H	T			Cyclin B 313-321	a	
	H	E	E	T	P	P	T	T	S			c-myc 241-249	a	

* Motif and ligands deduced by exclusion: Class I ligands from a *c-myc* transfected B-cell line expressing A2, A68, and B40 were sequenced. Those not containing an A2 or A68 motif were thought to contain B40 ligands.

References:

a: Harris et al. 1993

Table 3 (Continued)
L HLA-B*4402

	Position										Ref.
	1	2	3	4	5	6	7	8	9	10	
Anchor or auxiliary anchor residues	E										a
Preferred residues	A S	M I L D	I	V	Y			F Y	F Y		
Others	D	N R K	P								

References:

a: Fleischhauer et al. 1994

M HLA-B*4403

	Position										Source	Ref.	
	1	2	3	4	5	6	7	8	9	10			
Anchor or auxiliary anchor residues	E										F F	a	
Preferred residues	A S	M I L V D											
Others		N R K	P V K	I F									
Examples for ligands	A A	E E	D M	K G	E K	N G	Y S	K F	K K	F Y	HSP 90 427–436 Elongation factor 2 48–57	a a	
B*440x-restricted T-cell epitope		E	E	N	L	L	D	F	V	R	F	EBNA 6 130–139	b

References:

a: Fleischhauer et al. 1994; b: Khanna et al. 1992

Table 3 (Continued)
N HLA-B*5101

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	A						F				a
	P						I				
	G										
Other preferred residues	I	W	I	G	V	N	K	T	W		
	L	F	L	V	T	I	Q		M		
	V	M	I	G	L	R			V		
	Y	F	K	A	K	E			L		
	D	W	E	I	Q						
		Y	D	S							
		V									
		E									
		H									
		D									
		R									
		N									
Examples for ligands	Y	P	F	K	P	P	K	V		UBC5, yeast 61–68	a
	D	A	H	I	Y	L	N	H	I	Thymidylate synthase 253–261	a
	T	G	Y	L	N	T	V	T	V	GBLP 192–200	a
	d	A	Y	A	L	N	H	T	L	Unknown	a
	I	P	P	E	V	N	R	Q	L	Unknown	a

References:

a: Falk et al. 1995a

O HLA-B*5102

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	P	Y					I				a
	A						V				
	G										
Other preferred residues	F	G	V	I	R	T					
	V	E	Q	N	E	R					
	L	K	N	Q	Q	Y					
	I	L	G	T	K						
		T	T								
		Q									
		R									
		N									
		H									
Examples for ligands	Y	A	Y	D	G	K	D	Y	I	MHC I α chain 140–148	a
	Y	P	F	K	P	P	K	V		UBC5, yeast 61–68	a
	L	P	P	G	R	I	I	K	X	Unknown	a
	L	P	F	T	V	I	L	v		CDC25 homol. 560–567	a
	T	G	Y	L	N	T	V	T	V	GBLP 192–200	a
	F	A	Y	D	G	K	D	Y	I	MHC I α chain 140–148	a
	F	P	S	E	I	V	G	K	R	Ribosomal protein S7/S8A 135–144	a
	M	P	W	F	K	G	w	K	V	Elongation factor 1 a 208–216	a

References:

a: Falk et al. 1995a

Table 3 (Continued)
P HLA-B*5103

	Position									Comments	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	A		Y					V		Anchor at 9 deduced from individual ligands	a
	P							I			
	G							F			
Other preferred residues	T	F	F	E	G	I	V				
	V	W	D	L	A	K	M				
	D		L	N	V	T					
				R	N						
				G	Q						
				Q	M						
				T	R						
				V							
										Source	
Examples for ligands	T	G	Y	L	N	T	V	T	V	GBLP 192–199	a
	D	A	H	I	Y	L	N	H	I	Thymidilate synthase 253–261	a
	Y	F	D	d	t	L	E	D	F	Unknown	a

References:

a: Falk et al. 1995a

Q HLA-B*5201

	Position									Comments	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	Q	F		L			I	I		C-terminal anchor at 8 or 9	a
	Y		I			V	V				
	W		V								
Other preferred residues	V	M	I	L	M	K	K	M	M		
	L	F	L	I	F	N	E	F	F		
	I	P	P	V	A	L	Q				
				D	P	T	T	Y			
				K	K	G	S				
					E						
					A						
										Source	
Examples for ligands	T	G	Y	L	N	T	V	T	V	GBLP 192–200	a
	G	Q	F	K	T	Y	A	I		Ribos. prot. S21 60–67	a
	H	S	T	I	M	P	R	L		P1-CDC21 259–266	a
	G	F	Y	P	G	S	I	E	V	MHC II β chain 150–158	a
	V	Q	I	F	G	N	K	M		RBAP-2 266–273	a
	Y	P	D	P	A	N	G	K	F	Elongation factor 2 265–273	a
	L	Q	F	P	V	G	R	I		Histone 2a Z 25–32	a
	H	M	Y	I	F	L	H	T	V	Unknown	a

References:

a: Falk et al. 1995a

Table 3 (Continued)**R HLA-B53**

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues		P									a
T-cell epitope	K	P	I	V	Q	Y	D	N	F	P. falciparum LSA-1 1786–1794	a

References:

a: Hill et al. 1992

S HLA-B*5801

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	A		P	V					F		a
	S		E	I					W		
	T		K	L							
				M							
				F							
Other preferred residues	K	G	G	D	A	I	L	N	Y		
	R	T	Q	D	V	Y	R				
	I	I	R	N	L	M	K				
				L	T	F	N	T			
				V		Y					
				F		W					
				Y		Q					
				N							
				Q							
Examples for ligands	K	A	G	Q	V	V	T	I	W	Lamin C 490–498	a
	A	G	D	R	T	F	Q	K	W	MHC class I 260–268	a
	I	t	T	<u>K</u>	A	I	S	R	F	Unknown	a
	R	T	D	G	K	V	F	Q	F	Ribosomal protein L30 23–31	a
	I	T	S	Q	D	V	L	H	S	Cytochrome C oxidase 154–163	a
	I	S	D	S	N	P	F	L	T	Unknown	a
	K	t	D	e	V	V	T	L	F	Unknown	a
	V	T	S	P	<u>L</u>	T	V	E	W	MHC class II β 209–217	a
	g	A	V	N	<u>Y</u>	V	M	T	f	Glucose transporter 5 322–330	a

References:

a: Falk et al. 1995c

Table 3 (Continued)
T HLA-B60 (B*40012)

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues		E				I		L			a
						V					
Other preferred residues	A	P	L	K	L	K					
	V	K	I	N	Y	R					
	I	D	V	P	M	Q					
	L	G	D	V							
	M	N	T	I							
	F	Q	N	D							
	S	T	P	R							
	D	G	Q								
	N	K									
		Q									
Examples for ligands	K	E	S	T	L	H	<u>L</u>	V	L	Ubiquitin 63–71	a
	H	E	A	T	L	R	c	w	A	MHC class I 221–230	a
	Y	E	I	H	D	G	<u>M</u>	N	L	HSHMO2C05	a
	S	E	S	P	I	V	<u>V</u>	V	L	Signal peptidase 45–54	a
	I	E	V	D	P	D	T	K	E	Ribosomal protein S17 95–105	a

References:

a: Falk et al. 1995c

U HLA-B61 (B*4006)

	Position									Comments	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	E	F		I			V				a
	I										
	L										
	V										
	Y										
	W										
Other preferred residues	G	P	M	E	V	N	Y	K	A	P1 deduced from individual ligands	
	R	T	G	I	V	S	P				
			P	L		L					
			S	M		W					
			N	D		I					
			D	G		T					
			K	V		R					
			A	F		D					
			R	N		Q					
			N	S		G					
			Q	K							
Examples for ligands	G	E	F	G	G	F	G	S	V	IEF (mRNA) 9306 127–135	a
	E	E	F	Q	F	I	K	K	A	Associated-microfibril. protein 72–80	a
	G	E	F	V	D	L	Y	V		Ribosomal protein S21 6–13	a
	R	E	R	R	D	N	Y	V		Ribosomal protein S17 77–84	a
	R	E	I	I	I	N	A	V		Ribonucl. reductase 290–297	a
	G	E	F	S	I	T	Y	K		Ribosomal protein S15 116–123	a
	G	E	H	G	L	i	I	R	V	Unknown	a
	R	E	<u>M</u>	I	P	F	A	D	i	Unknown	a

References:

a: Falk et al. 1995c

Table 3 (Continued)
V HLA-B62 (B*1501)

	Position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or auxiliary anchor residues	Q		I			F				a		
	L		V			Y						
Other preferred residues	I	M	K	P	G	V	V	Y				
	V	A	E	L	T	T	V					
	N	G	F	G	L	T						
	F	D	T	I	I							
	P											
	Y											
	H											
	R											
Examples for ligands	V	L	K	P	<u>G</u>	M	V	V	F	Elongation factor 1 α 271–280	a	
	Y	L	G	E	<u>F</u>	S	I	T	Y	Ribosomal protein S15 114–122	a	
	G	Q	R	K	<u>G</u>	A	G	S	V	Ribosomal protein L8 (rat) 7–15	a	
	K	I	K	S	<u>F</u>	V	K	V	Y	Ribosomal protein L27 66–74	a	
	I	Q	P	G	<u>R</u>	G	F	V	L	Unknown	a	
	S	Q	F	G	<u>G</u>	G	S	Q	Y	Unknown	a	
	G	Q	R	K	<u>P</u>	A	T	S	Y	Ribosomal protein L28 (rat) 68–76	a	
	V	Q	G	P	<u>V</u>	G	L			Collagen $\alpha 1$ 1106–1112	a	
T-cell epitopes	I	L	G	N	K	I	V	R	M	Y	HIV gag 267–276	b

References:

a: Falk et al. 1995 c; b: Buseyne et al. 1993

W HLA-B*7801

	Position								Comments	Ref.
	1	2	3	4	5	<u>6</u>	7	8		
Anchor or auxiliary anchor residues	P			I		A			This motif is only partial; the C-terminal anchor has not been determined	a
	A			L						
	G			F						
				V						
Other preferred residues	Y	F	D		A	K				
	D	D	G		V	S				
	W	G	V		N					
		L	N		K					
		V	R		Q					
		S	Q		E					
		Q	S							
		R	T							
		N								

References:

a: Falk et al. 1995 a

Table 4 HLA-C motifs
A HLA-Cw*0301

	Position									Source	Ref.	
	1	2	<u>3</u>	<u>4</u>	5	<u>6</u>	7	8	9			
Anchor or auxiliary anchor residues			V	P		F			L		a	
			I			Y			F			
			Y						M			
			L						I			
			M									
Other preferred residues	A	E	E	N	M	Q	T					
	R	N	R			K						
						S						
						M						
T-cell epitopes	H	Q	A	I	S	P	R	T	L	HIV gag 144–152	b	
or	Q	M	<u>V</u>	H	Q	A	I	S	P	R	HIV gag 141–152	

References:

a: Falk et al. 1993 a; b: Littaua et al. 1991

B HLA-Cw*0401

	Position									Source	Ref.
	1	2	3	4	5	<u>6</u>	7	8	9		
Anchor or auxiliary anchor residues		Y				V			L		a
		P				I			F		
		F				L			M		
Other preferred residues	D	D	A		A	K					
	H	E	H			S					
		P	M			H					
			T								
			R								
T-cell epitope	S	F	N	C	G	G	E	F	F	HIV-1 gp 120 380–388	b

References:

a: Falk et al. 1993 a; b: Johnson et al. 1993

C HLA-Cw*0602

	Position									Source	Ref.	
	1	2	3	4	<u>5</u>	<u>6</u>	7	8	9			
Anchor or auxiliary anchor residues					I	V			L		a	
					L	I			I			
					F	L			V			
					M				Y			
Other preferred residues	I	P	P	P	K	A	R	Y				
	F	R	I	E		T	K	E				
	K	G	D		S	Q	Q					
	Y	F	Q			N	N					
		Y	L					R				
					K			G				
					N			T				
					A			S				
								K				
Examples for ligands	Y	Q	F	T	G	I	K	K	Y	Unknown	a	
	V	R	H	D	G	G	N	V	L	Unknown	a	
	F	A	F	p	I	<u>I</u>	q	R	V	Unknown	a	
	X	Q	r	T	P	k	A	g	l	Y	Unknown	a

References:

a: Falk et al. 1993 a

Table 4 (Continued)
D HLA-Cw*0702

	Position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor or auxiliary anchor residues	Y P		V Y I I L L M F M		V I L M		Y F L			a	
Other preferred residues	R D G E A V Q P S G	P G E R M A N F R D V K F E	D A V P S G	T R M N F R D V K F E	A M A	Y E Y E Y	E A L				
Examples for ligands	K R N I I N	Y Y K Y R Y	F R A P K G	D P D q P G	E G Y n Y G	H T I v I N	Y V L i w Y	E A K L E G	Y L Y Y Y S	CKS-2 11–19 Histone H3.3 40–48 Protein synthesis factor eIF-4C 87–95 Unknown Glutamyl-tRNA synthetase 343–351 Homologous hnRNP A2 or B1 (S11 = N) 277–288 Unknown Unknown	a a a a a a a
	F X	Y M	P P	P f	y l	1 L	Y d		G		a a

References:

a: Falk et al. 1993 a

Table 5 Processing motif for all MHC class II ligands

Absolute position																	Ref.
1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	
P												p	p	p	p	p	a, b, c

References:

a: Falk et al. 1994 b; b: Kropshofer et al. 1993; c: Malcherek et al. 1993

Table 6 Human MHC class II motifs
A HLA-DRB1*0101

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor residues				L,A		A,G			L,A			
	Y,V,				S,T			I,V			a, b, c	
	L,F,		I,V									
	I,A		M,N		P			N,F				
	M,W		Q					Y				
Examples for ligands	VGSD	W	R	F	L	R	G	Y	H	Q	YA	HLA-A2 103–117
	VGSD	W	R	F	L	R	G	Y	H	Q	YAYDG	HLA-A2 103–120
	VGSD	W	R	F	L	R	G	Y	H	Q	Y	HLA-A2 103–116
	GSD	W	R	F	L	R	G	Y	H	Q	YA	HLA-A2 104–117
	LPKPPKPVSK	M	R	M	A	T	P	L	L	M	QALPM	Invariant chain 97–120
	IPAD	L	R	I	I	S	A	N	G	C	K	Na ⁺ -K ⁺ -ATPase 199–216
	RVE	Y	H	F	L	S	P	Y	V	S	PKESP	Transferrin receptor 680–696
	YKHT	L	N	Q	I	D	S	V	K	V	WPRRPT	Cattle fetuin 56–74
	AILE	F	R	A	M	A	Q	F	S	R	KTD	Unknown
	PK	Y	V	K	Q	N	T	L	K	L	AT*	Influenza HA 306–318

* Alignment determined by structural analysis

References:

a: Hammer et al. 1992; b: Falk et al. 1994b; c: Chicz et al. 1992; d: Kropfhofer et al. 1992; e: Stern et al. 1994.

Table 6 (Continued)
B HLA-DRB1*0301 (DR17)

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or auxiliary anchor residues	L,I F,M V		D		K,R E,Q N			Y,L F			a, b, c	
Examples for ligands	ISNQ	L	T	L	D	S	N	T	K	Y	FHKLN	Apolipoprotein B 2877–2894
	ISNQ	L	T	L	D	S	N	T	K	Y	FHKL	Apolipoprotein B 2877–2893
	ISNQ	L	T	L	D	S	N	T	K	Y	FHK	Apolipoprotein B 2877–2892
	VDT	F	L	E	D	V	K	N	L	Y	HSEA	α 1-Antitrypsin 149–164
	KPRA	I	V	V	D	P	V	H	G	F	MY	LDL-Receptor 518–532
	KQT	I	S	P	D	Y	R	N	M	I		IgG2a, Membrane domain
	YPD	F	I	M	D	P	K	E	K	D	KV	Unknown
	NIQ	L	I	N	D	Q	E	V	A	R	FD	Unknown
	LLS	F	V	R	D	L	N	Q	Y	R	ADI	Transferrin receptor 618–632
	LPKPPKPVSK	M	R	M	A	T	P	L				Invariant chain 97–113
	LPKPPKPVSK	M	R	M	A	T	P	L	M			Invariant chain 97–119
	LPKPPKPVSK	M	R	M	A	T	P	L	M			Invariant chain 97–120
	PKPPKPVSK	M	R	M	A	T	P	L				Invariant chain 98–113
	PKPPKPVSK	M	R	M	A	T	P	L	M			Invariant chain 98–117
	KPPKPVSK	M	R	M	A	T	P	L	M			Invariant chain 99–116
	KPPKPVSK	M	R	M	A	T	P	L	M			Invariant chain 99–119
	VDDTQF	V	R	F	D	S	D	A	A	S		HLA-A30 28?
	ATKYGN	M	T	E	D	H	V	M	H	L	LQNA	Invariant chain 131–149
	VFL	L	L	A	D	K	V	P	E	T	SLS	ACh receptor 289–304
	LNK	I	L	L	D	E	Q	A	Q	W	K	ICAM-2 64–76
	GPPKLD	I	R	K	E	E	K	Q	I	M	IDIFH	IFN- γ receptor 128–147
	GPPKLD	I	R	K	E	E	K	Q	I	M	IDIFHP	IFN- γ receptor 128–148
	GKFA	I	R	P	D	K	K	S	N	P	IIRTV	Cyt-b5 155–172
	YAN	I	L	L	D	R	R	V	P	Q	TDMTF	Apolipoprotein B 1207–1224
	NLF	L	K	S	D	G	R	I	K	Y	TLNKNSLK	Apolipoprotein B 1276–1295
	IPDNL	L	K	S	D	G	R	I	K	Y	TLNKN	Apolipoprotein B 1273–1292
	IPDNL	L	K	S	D	G	R	I	K	Y	TLNK	Apolipoprotein B 1273–1291
	IPDNL	L	K	S	D	G	R	I	K	Y	TLN	Apolipoprotein B 1273–1290
	IPDNL	L	K	S	D	G	R	I	K	Y	TL	Apolipoprotein B 1273–1289
	NLF	L	K	S	D	G	R	I	K	Y	TLNK	Apolipoprotein B 1276–1291
	NLF	L	K	S	D	G	R	I	K	Y	TLN	Apolipoprotein B 1276–1290
	VTT	L	N	S	D	L	K	Y	N	A	LDLTN	Apolipoprotein B 1294–1810
	V	G	S	D	W	R	F	L	R		GYHQYA	HLA-A2 103–117

References:

a: Malcherek et al. 1993; b: Geluk et al. 1994; c: Geluk et al. 1992; d: Riberdy et al. 1992; e: Chicz et al. 1993; f: Sette et al. 1992

Table 6 (Continued)
C HLA-DRB1*0401 (DR4Dw4)

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	F,Y W,I L,V M			F,W I,L V,A D,E	N,S T,Q H,R ali.*	pol.* chg.* ali.*		pol.* K			a, b, c, d	
Examples for ligands				no R,K								
VDDTQ	F F F F	V V V V	R R R R	F F F F	D D D D	S S S S	D D D D	A A A A	SQRMEP SQRM SQRM SPRGE...	HLA-A2 33–47 HLA-A2 28–45 HLA-A2 33–45 HLA-C 28–?	a	
DGKD	Y	I	A	L	N	E	D	L	S		HLA-B44 143–156	a
LSS	W	T	A	A	D	T	A	A	Q	ITQ	HLA-B44 154–168	a
LSS	W	T	A	A	D	T	A	A	Q	IT	HLA-B44 154–167	a
IY	F	R	N	Q	K	G	S	H	S	GLQPTGFL	HLA-DR4β 252–270	a
DVA	F	V	K	D	Q	T	V	I	Q	NTD	Cattle transferin 68–82	a
YDHN	F	V	K	A	I	N	A	I	Q	KSW	Cathepsin C 170–185	a
KHKV	Y	A	C	E	V	T	H	Q	G	...	Igκ chain C region 80–?	a
HKV	Y	A	C	E	V	T	H	Q	G	L...	Igκ chain C region 81–?	a
DGP	F	R	I	I	T	V	P	A	A	LDY	Unknown	a
TGN	Y	R	I	E	S	V	L	S	S		Sphingolipid activator protein 3 165–176	a
GERA	M	T	K	D	N	N	L	L	G	...	HSC 70 445–?	a
XXX	Y	E	X	A	L	S	L	P	S	K...	Unknown	a
GSLF	V	Y	N	I	T	T	N	K	Y	KAFLKQ	VLA-4 229–247	e
SPEDF	V	Y	Q	F	K	G	M	C	Y	F	HLA-DQβ 3.2 chain 24–38	e
AAPYEKEVP	L	S	A	L	T	N	I	L	S	AQL	PAI-1 261–281	e
GVYF	Y	L	Q	W	G	R	S	T	L	VSVS	Ig heavy chain 121–?	e
AEALERM	F	L	S	F	P	T	T	K	T		Cattle hemoglobin 26–41	e
LRS	W	T	A	A	D	T	A	A	Q	ITQRKWEAA	HLA-Cw9 130–150	e
DLSS	W	T	A	A	D	T	A	A	Q	ITQRKWEAA	HLA-Bw62 129–150	e
APSP	L	P	E	T	T	E	N	V	V	CALG	HLA-DRα chain 182–198	e

* pol.: Polar; chg.: charged; ali.: aliphatic

References:

a: Friede and co-workers, submitted; b: Sette et al. 1993; c: Hammer et al. 1993; d: Hill et al. 1994; e: Chicz et al. 1993

D HLA-DRB1*0402 (DR4Dw10)

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	V,I L,M			Y,F W,I L,M R,N	N,Q S,T K	R,K H,N Q,P; rare D,E		pol.* ali.* H			a	
Examples for ligands				no D,E								
GPDR	L	L	R	G	H	N	Q	F	A	YDGKD	HLA-B38 128–146	a
GR	L	L	R	G	H	N	Q	F	A	YDGK	HLA-B38 131–145	a
I	I	K	G	V	R	K	S	N	A	AERRG	HLA-DRα 238–252	a
I	Y	F	R	N	Q	K	G	H	H	SGLQPTGFLS	DR4β 248–266	a
F	I	Y	F	R	N	Q	K	G	H	SGLQP	DR4β 250–261	a
Y	V	R	F	D	S	D	V	G	H	SGLQPTGFLS	DR4β 249–266	a
LPKPPKPVSK	M	R	M	A	T	P	L	L	Q		Invariant chain 97–?	a
FDQK	I	V	E	W	D	S	R	K	S	KYFE	BLAST-1 62–78	a
DQK	I	V	E	W	D	S	R	K	S	KYF	BLAST-1 63–77	a
IKI	I	S	K	I	E	N	H	E	G	VRR	Pyruvate kinase 264–278	a
IKI	I	S	K	I	E	N	H	E	G	VR	Pyruvate-kinase 264–277	a
FGR	I	G	R	L	V	T	R	A	A	FNSG	GAPDH 11–25	a
FGR	I	G	R	L	V	T	R	A	A	FN	GAPDH 11–23	a
GFGR	I	G	R	L	V	T	R	A	A	FNSG	GAPDH 10–25	a
CNE	I	I	N	W	L	D	K	N	Q		HSC 70 574–585	a
QPD	L	R	Y	L	F	L	N	G	N		Leucine-rich α2-glycoprotein 200–211	a

References:

a: Friede and co-workers, submitted

Table 6 (Continued)
E HLA-DRB1*0404 (DR4Dw14)

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	V,I L,M			F,Y W,I L,V M,A D,E no R,K		N,T S,Q R	pol.* chg.* ali.*		pol.* ali.* K		a	
Examples for ligands	GSHS SHS YDNS	M M L	R R K	Y F I	F H S	H T N	A A A	M M S	S RPGRGE TTN	RPGRGE RPGRGE GAPDH 139-154	HLA-B60 1-? HLA-B60 2-? GAPDH 139-154	a a a

* pol.: Polar; chg.: charged; ali.: aliphatic

References:

a: Friede and co-workers, submitted

F HLA-DRB1*0405 (DR4Dw15)

* pol.: Polar; chg.: charged; ali.: aliphatic

References:

a: Friede and co-workers, submitted; b: Matsushita et al. 1994; c: Kinouchi et al. 1994

Table 6 (Continued)
G HLA-DRB1*1101

	Relative position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues											a, b
	W,Y			M,L		R,K					
	F			V,I							
Examples for ligands	IDF	Y	T	S	I	T	R	A	R	EE	HSC 70 291–305
	CPAG	Y	T	C	N	V	K	A	R	CEK	Granulin D 41–56
	VNH	F	I	A	E	F	K	R	K	H	KKD
	VNH	F	I	A	E	F	K	R	K	H	K
	MR	Y	F	H	T	S	V	S	R	P	Homol. HSC 70 238–250
	KHKV	Y	A	C	E	V	T	H	Q	G	GRGEP
										LS	HLA-Bw61 5–20
											Homol. Ig κ-chain 190–204

References:

a: Hammer et al. 1993; b: Newcomb and Cresswell 1993

H HLA-DRB1*1201

	Relative position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues	I,L		L,M		V,Y			Y,F			a
	F,Y		N,V		F,I			M,I			
	V		A		N,A			V			
Examples for ligands	GPDGRL	L	R	G	Y	D	Q	F	A	Y	DGK
	GPDGRL	L	R	G	H	N	Q	Y	A	Y	HLA-B38 104–121
	TGT	I	K	L	L	N	E	N	S	Y	HLA class I 104–119
	T	I	K	L	L	N	E	N	S	Y	Transferrin receptor 142–155
	FTGT	I	K	L	L	N	E	N	S	Y	Transferrin receptor 144–156
	DFTGT	I	K	L	L	N	E	N	S	Y	Transferrin receptor 141–156
	SDEK	I	R	M	N	R	V	V	R	N	NLR
	SSV	I	T	L	N	T	N	V	G	L	Valosin-cont. protein p97 78–93
	EAL	I	H	Q	L	K	I	N	P	Y	Homol. to apolipoprotein
	AHL	F	K	Q	N	K	V	V	H	V	Unknown
										NG	Dihydrolipoamide dehydrogenase 138–152

References:

a: Falk et al. 1994 b; b: Falk and co-workers, unpublished

I HLA-DRB1*1501 (DR2b)

	Relative position									Source	Ref.
	1	2	3	4	5	6	7	8	9		
Anchor residues	L,V			F,Y			I,L				a, b
	I			I			V,M,				
							F				
Examples for ligands	EAEQ	L	R	A	Y	L	D	G	T	G	HLA-A3 152–166
	L	E	E	F	G	R	F	A	S	FEAQG	HLA-DRα 45–58
	D	V	G	V	Y	R	A	V	T	P	QGRPDA
T-cell epitope	PV	V	H	F	F	K	N	I	V	T	HLA-DQw6 43–58
											MBP 85–95

References:

a: Vogt et al. 1994; b: Wucherpfennig et al. 1994

Table 6 (Continued)
K HLA-DRB5*0101 (DR2a)

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	F,Y L,M									R,K		
Examples for ligands	DVGV	Y	R	A	V	T	P	Q	G	R	P	HLA-DQw6 43–56
	DVGV	Y	R	A	V	T	P	Q	G	R	PDA	HLA-DQw6 43–58
	DSDVGV	Y	R	A	V	T	P	Q	G	R	PD	HLA-DQw6 41–57
	DSDVGV	Y	R	A	V	T	P	Q	G	R	PDA	HLA-DQw6 41–58
	DSDVGV	Y	R	A	V	T	P	Q	G	R	PDAEY	HLA-DQw6 41–60
	AAD	M	A	A	Q	I	T	K	R	K	WEAAH	HLA-A3 135–151
	TAAD	M	A	A	Q	I	T	K	R	K	WEA	HLA-A3 134–149
	DVGE	F	A	A	V	T	E	K	R	R	PDAEYW	HLA-DR2b 43–61
T-cell epitopes	PK	Y	V	K	Q	N	T	L	K	L	AT	HA 307–319
		L	Q	A	A	P	A	L	D	K	L	HSP65 418–427
	VHF	F	K	N	I	V	T	P	R	T	P	MBP 87–99
	ASD	Y	K	S	A	H	K	G	F	K	GVD	MBP 131–145
	KG	F	K	G	V	D	A	Q	G	T	LSKI	MBP 139–153

References:

a: Vogt et al. 1994; b: Wucherpfennig et al. 1994; c: O'Sullivan et al. 1991; d: Anderson et al. 1988; e: Martin et al. 1991

L HLA-DQA1*0501/DQB1*0301

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor residues	F,Y I,M L,V									V,L I,M Y		
										Y,F M,L V,I		
Preferred residues	A		A	A								
Examples for ligands	TPL	L	M	Q	<u>A</u>	L	P	M	G	A	LPQG	Invariant chain 111–126
	TPL	L	M	Q	<u>A</u>	L	P	M	G	A	LPQ	Invariant chain 111–125
	KPPKPVSKMR	M	<u>A</u>	T	P	L	L	M	Q	A		Invariant chain 99–117
	LPKPPKPVSKMR	M	<u>A</u>	T	P	L	L	M				Invariant chain 97–115
	IPE	L	N	K	V	A	R	A	A	A		Transferrin receptor 579–597
	DVEV	Y	R	<u>A</u>	V	T	P	L	G	P	EVAGQF	DQ β chain 43–55

References:

a: Falk et al. 1994b

M HLA-DPA1*0201/DPB1*0401

	Relative position										Source	Ref.	
	1	2	3	4	5	6	7	8	9	10			
Anchor residues	F,L Y,M I,V A										F,L Y,M V,I A		
											V,Y I,A L		
Examples for ligands	EKK	Y	F	A	A	T	Q	F	E	P	L	AARL	Unknown
	KK	Y	F	A	A	T	Q	F	E	P	L	AARL	Unknown
	EKK	Y	F	A	A	T	Q	F	E	P	L	LNVANRR	Unknown
	GPG	A	P	A	D	V	Q	Y	D	L	Y	IL-3 Receptor α -chain 127–146	a

References:

a: Falk et al. 1994b

Table 6 (Continued)
N HLA-DPA1*0102/DPB1*0201

Anchor residues	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
ADEKKF GEP LPSQA	F,L M,V W,Y	W	G	K	Y	L	Y	E	I	A	RRHP RQVDG	Cattle serum albumin 152–170 Transferrin receptor 15–31 Cathepsin H 185–198
Examples for ligands		L	S	Y	T	R	F	S	L	A		a
		F	E	Y	I	L	Y	N	K	G		a

References:
a: Rötzschke and Falk 1994

Table 7 Other human class II ligands

MHC molecule	Peptide sequence	Source	Ref.
HLA-DR2 (DRB5*0101 or DRB1*1501)	NIVIKRSNSTAATNEVPEVTVF S NIVIKRSNSTAATNEV SDVGVYRAVTPQGRPDAE DVGVYRAVTPQGRPDAE DVGVYRAVTPQGRPD RVQPKVTVYPSKTQPLQH RVQPKVTVYPSKTQ LSPIHIALNFSLDPQAPVDSHGLRPALHYQ DGILYYYQSGGRLRPN IQNLIKEEAFLGITDEKTEG EHIFI LGATNYIYVLNEEDLQKV QELKNKYYQVPRKGIA FPKSLHTYANILLDRRVPTD FPKSLHTYANILLDRRVQ LWDYGMSSSPHVLRNR	HLA-DQ α HLA-DQ α HLA-DQ β HLA-DQ β HLA-DQ β HLA-DQ β HLA-DRB1*1501 HLA-DRB1*1501 Fibronectin receptor α K^+ channel protein Mannose binding protein MET protooncogene Guanylate binding protein 2 Apolipoprotein B100 Apolipoprotein B100 Factor VIII	97– 119 97– 112 42– 59 43– 59 43– 57 94– 111 94– 108 586– 616 173– 190 174– 193 59– 81 434– 450 1200–1220 1200–1218 1775–1790
HLA-DRB1*0701	RPAGDGTFOKWASVVVPSQ RPAGDGTFOKWASVV GDGTFOKWASVVVPSQEQRYT GDGTFOKWASVVVPSQ GTFOKWASVVVPSG GTFOKWASVVVPSQ GTFOKWASVVVPSQEQRYTCHV RETOISKTNTQTYREN RETOISKTNTQTYREN RETOISKTNTQTYRE RSN \tilde{Y} TPITNP \tilde{P} EVTVLTNSPVELREP GALANI AVDKANLEIMTKRSN SIQSPITVEWRAQSESAQS KMLSGIGGFVL VTQYLNATGNRWCSWLSQAR VTQYLNATGNRWCSWLS TSILCYRKREWIK PAFRFTREAAQDCEV GDMYPK TWSGMLVGALCALAGVLTI TPSYVAFTDTERLIGDA TPSYVAFTDTERLIG VPGLYSPCRAFFNKEELL VPGLYSPCRAFFNK KVDLTFSKQHALLCSDYQADYES KVDLTFSKQHALLCS FSHDYRGSTSHRL LPKYFEKKRNII APVLISQKLSPYINLVPK VGSDWRFRLRGYHQYAYDG PKPPKPVS KMRM ATPLLMOALP APSPLPETENVVCALGLTV KHKVYACEVTHQGL	HLA-A29 HLA-B44 HLA-DR α chain HLA-DQ α chain 4F2 LIF receptor Thromboxane-A synthase K^+ channel protein Hsp 70 EBV MCP Apolipoprotein B 100 Complement C9 HLA-A2 Invariant chain HLA-DR α chain Ig kappa chain	234– 253 234– 249 237– 258 237– 254 239– 252 239– 253 239– 261 83– 99 83– 98 83– 97 101– 126 58– 78 179– ? 318– 338 318– 334 854– 866 406– 420 492– 516 38– 54 38– 52 1264–1282 1264–1277 1586–1608 1586–1600 1942–1954 2077–2089 465– 483 103– 120 98– 119 182– 200 188– 201

Table 7 (Continued)

MHC molecule	Peptide sequence	Source	Ref.
HLA-DRB1*0801	APSPLPETTENVVCALG SETVFLPREDHLFRKFHYLPFLP RHNYELDEAVTLO DPQSGALYISKVQKEFDNSTYI GALYISKVQKEFDNSTYI DPVPKPVKIEKIEDMDD DPVPKPVKIEKIED FTFTISRLEPEDFAVYYC FTFTISRLEPEDFAV DPVEMRRLNYQTPG YQLLRS咪GTYEELAPIV GNHLYWKWQIPDCENVK LPFFLFQAYHPNNSSPVCY RPSMLQHLLR DDFMGQLLNGRVLFPVNQLQGA IPRLQKIWKNYLSMNKY EPFLYILGKSRVLEAQ NRSEEFLLIAGKLQDGLL RSEEFLLIAGKLQDGLL SEEFLIAGKLQDGLL NRSEEFLLIAGKL QAKFFACIKRSDGSCAWYRGAAPPKQEF QAKFFACIKRSDGSCAWYR DRPFLFVVVRHNPTGTFLFM MPHFFRLFRSTVKQVD QNFTVIFDTGSSNWLWPSVYCTSP QNFTVIFDTGSSNLWV TAFQYIIDNKGDSDAS DEYÝRRLRLVLRAREQIV EAIYDICCRNLDIERPT EAIYDICCRNLDI HELEKIKKQVEQEKCIEQAAL AEVYHDVAASEFF ... KRSFFALRDQIPDL ROYRLKKISKEEKTPGC KNIFHKVNQEGLKLNSNDMM KNIFHKVNQEGLKL YKQTVSLSDIQPYSLVTTLNS STPEFTILNLTIHIPSFT TPEFTILNLTIHIPSFTID TPEFTILNLTIHIPSFT SNTKYFHKLNIQLDF LPFFKFLPKYFEKKRNT LPFFKFLPKYFEKRR WNFYYSPQSSPDKKL DVIWELLNHAQEHFQKDKSKE DVIWELLNHAQEFG DVIWELLNHAQEHE IAIALLMASQEPEORMSRNFVR IAIALLMASQEPEORM	HLA-DR α chain HLA-DR α chain HLA-DP β chain LAM Blast-1 Ig κ chain LAR LIF receptor IFN- α receptor IL-8 receptor Ca $^{2+}$ release channel CD35 CD75 Calcitonin receptor TIMP-1 TIMP-2 PAI-1 Cathepsin E Cathepsin S Cystatin SN Tubulin α -1 chain Myosin β heavy chain α -enolase <i>c-myc</i> K-ras Apolipoprotein B-100 Cattle transferrin von Willebrand factor	182– 198 158– 180 80– 92 88– 108 92– 108 129– 146 129– 143 63– 80 63– 77 1302–1316 709– 726 271– 287 169– 188 2614–2623 359– 380 106– 122 38– 53 101– 118 102– 117 103– 117 101– 112 187– 214 187– 205 378– 396 133– 148 89– 112 89– 104 189– 205 41– 58 207– 223 207– 219 1027–1047 23–? 371– 385 164– 180 1724–1743 1724–1739 1780–1799 2646–2662 2647–2664 2647–2662 2885–2900 2072–2088 2072–2086 4022–4036 261– 281 261– 275 261– 273 617– 636 617– 630
HLA-DR11 or Dw52	SXVITLNTNVGLYXQS DPXQDELQKLNAXDP XPELNKVÅRAAAEVAG	Homol. Apolipoprotein Unknown Homol. Transferrin receptor	3345–3360 b b 580– 595 b
DR17 or DRw 52	TFDEIASGFRQGGASQ YGYTSYDTFSWAFL GOVKKNNHQEDKIE TGHGARTSTEPTTDY KELKROYEKKLRQ SPLQALDFFGNGPPVNYKTGNL	Glucose transporter Na $^+$ channel protein CD45 EBV gp220 EBV tegument p140 IP 30	459– 474 384– 397 1071–1084 592– 606 1395–1407 38– 59

References:

a: Chicz et al. 1993; b: Newcomb and Cresswell 1993

Table 8 Mouse class II motifs
A H-2E^k

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	I,L V,F Y,W		I,L V,F S		Q,N A		K,R				a, b, c	
Examples for ligands	HPPHIE DNRM TPTL	I V V	Q H E	M F A	L I A	K A R	N E N	G F L	K K G	K R R VG	β_2m 42–56 HSC70 234–248 Serum albumin 347–361	c c c
	VNKE GFPT IP YDRN	I I L T	Q Y I K	N F M S	A P L P	V A N L	Q A K F	G N K V	V K A G	K K R R V	C cyt inhib. 41–55 ER60 448–461 Unknown $\alpha 1$ -antitryp. 397–410	c a a
		F LH	A L	E G	F Y	G P	T N	L Q	K L	K F	AAVHYDRSG (human) dead box protein	a
										R	Cattle fetuin 342–	a
T-cell epitopes	RADL RADL LEDARR QD VTV	I I L I L	A Y K L T	Y L A I A	L K A R L	K Q Y F G	Q A E K A	A T K S I	T A K H L	K A K PETL K	MCC 91–103 PCC 91–104 λ rep 12–26 SWMb 26–40 SWMb 66–78	b b e e d
											EqMb 69–77 MoMb 69–77 MoHb 68–76	b b b
											HEL 1–18 HEL 81–96	e d
	KVFGR SALLSSD VEK RTDKYGRG HEHQ	C I W Y L	E T V G A	L A S P Y	A A V E I	A A N A Y	A M C A S	G K A F D	L K K T G	HGLD GTD KKMVENAK MVN KEKLNW	HEL 108–119 SNase 51–70 SNase 81–100 SNase 121–140 LLO 218–226	d e e f b

References:

a: Schild and co-workers, submitted; b: Reay et al. 1994; c: Marrack et al. 1993; d: Spouge et al. 1987; e: Altuvia et al. 1994; F: Sette et al. 1989

B H-2E^d

	Relative position									Source	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	W,Y F,I, L,V		K,R I		I,L V,G		K,R				a	
Examples for ligands	SQLELR LELR ERAEE RAEA AQ SLDEH	W W W W F Y	K K R R M	S S Q Q W H	R R K K I	H H L L I	I I H H I	K K G R R	E R R L	IL-2R. γ chain 168–182 IL-2R. γ chain 170–182 Apo-E prec. 222–236 Apo-E prec. 223–236 Unknown Similar Apolipoprotein B 2211–2224	a a a a a a	
										GQFY LV	C. elegans cDNA homol. 74–87 Actin B 8–21	a a
										R		a
T-cell epitopes	ALWFRRNH KYLEFISEA NKALE W A SS LEDARR EK	F I L W Y F I	V I F A V E R	F H R W Y R A	G V D R Y E P	G L I N K I G	G H A R P F K	T S A C N P K	K R A K H P K	TV R Y G TD EQHLRKSE AGF V TD H K K K	Ig lambda 91–108 SWM 102–118 SWM 132–146 HEL 108–119 SNase 112–129 FLU PR/8 HA 109–119 λ rep 12–26 HIV-1 gag p17 17–28	b c c d c c c c f

References:

a: Schild and co-workers, submitted; b: Bogen et al. 1986; c: Spouge et al. 1987; d: O'Sullivan et al. 1991; e: Chicz et al. 1992; f: Sette et al. 1989

Table 8 (Continued)**C H-2E^a**

	Relative position									Comments	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	I,V L			L,I V		Q,N		K,R		This motif has been predicted based on prediction of pocket structure and comparison with H-2E ^k and H-2E ^d motifs	a	
Source												
Examples for ligands	HPPHIE EGEC MQKEITA CT EGSLI	L I V L F V	Y Q E A A	V M W P I K	L L L T C I	K N H M W M	I G R K L P	K K Y I F S	K K L K H S	DG NG II VFFL E	Carboxypeptidase A 44–54 β_2 42–56 H-2L ^d 160–174 β -actin 286–303 Substance P receptor 255–269 HSP60 478–492	b b b b b
T-cell epitope	DL	I	A	Y	L	K	Q	A	T	K	MCC 93–103	c, d

References:

a: Schild and co-workers, submitted; b: Marrack et al. 1993; c: Altuvia et al. 1994; d: Reay et al. 1994

D H-2E^b

	Relative position									Comments	Ref.	
	1	2	3	4	5	6	7	8	9			
Anchor or preferred residues	W,F Y			L,I F,V		Q,N, A		K,R		This motif has been predicted based on prediction of pocket structure and comparison with H-2E ^k and H-2E ^b motifs	a	
Source												
Examples for ligands	SPSYV SPSYV SPSYV GK XPQS	Y Y Y Y Y	H H H L L	Q Q Q Y I	F F F E H	E E E I E	R R R A X	R R R R X	A A A R X	K K K H I	YK YKREPVSL PYFY BSA 141–155 Unknown	b b b b b
T-cell epitopes	RTDKYGRG DL	L I	A A	Y Y	I L	Y K	A Q	D A	G T	K K	MVN MCC 93–103	c, d c, d

References:

a: Schild and co-workers, submitted; b: Rudensky et al. 1991; c: Altuvia et al. 1994; d: Reay et al. 1994

Table 9 Other mouse class II ligands

MHC Molecule	Peptide sequence	Source	Ref.
H-2A ^b	HNEGFYVCPGPHRP	MuLV env	145–158
	ASFEAQGALANIAAVDKA	H-2E α	52–68
	KPVSQMRMATPLLMR	Invariant chain	86–100
	NYNAYNATPATLAVD	Unknown	a, b
	RPDAEYWNSQPE	H-2A β	55–66
	XNADFKTPATLTVDKP	IgG V μ	59–74
H-2A ^s	IRLKITDSGPRVPIGPN	MuLV env	255–269
	IRLKITDSGPRVPIG	MuLV env	255–267
	WQSQSITCNVAHPASS	IgG2a	194–210
	NVEVHTAQTOOTHREDY	IgG2a	281–296
	KPTEVSGKLVHANFGT	Transferrin receptor	203–218
	XPYMFADKVVHLPGSQ	Unknown	b
H-2A ^d	WANLMEKIQASVATNP	Apo-E	268–284
	WANLMEKIQASVATNP	Apo-E	268–283
	DAYHSRAIQVVRARKQ	Cys-C	40–55
	ASFEAQGALANIAAVDKA	H-2I-E α^d	52–68
	ASFEAQGALANIAAVDK	H-2I-E α^d	52–67
	EFTQQIQLQAEIFQAR	Apo-E	236–252
	EFTQQIQLQAEIFQAR	Apo-E	237–252
	KPVSQMRMATPLLMRPM	Li	85–101
	VPQLNQMVRTAAEVAGQX	Tf recip.	442–459
	ISQAVHAAHAEINE	Ovalbumin	323–336
	LEDARRLKAIYEKKK	λ repressor	12–26
	DGSTDYGILQINSR	Hen egg lysozyme	48–61
H-2A ^k	DGSTDYGILQINS		48–60
	DGSTDYGILQINSRW		48–62
	DYGILQINSRW (C)		52–63 (64)
	IIANDQGNRTTPSY	hsp70	28–41
	TPRRGEVYTCHVEHP	H-2I-A κ β chain	165–179
	KVHGSLARAGKVRGQTPKVAKO	S30 ribosomal protein	75–96
	AGKVRGQTPKVAQEEKKKKKT		83–103
	EPLVPLDNHIPENAQPG	Ryudocan	84–100
	XQLGAQNEMLXPL	Unknown	e
	XXKKGTDFQLNQLE	Transferrin	100–113
	KGTDQFQNLQLEGKKG	Transferrin	103–117
	YVRFDSFVG \bar{Y} RAVT	H-2A β^k	37–51
	XPLALQFAELPVNKG	Unknown	e
	XNLRFDSDVGEFRAV	H-2E β^k	33–47
	EDENLYEGLNLDGXSMYE	MBI	177–194
	XXLYNKGIMGEDSYPY	Cathepsin H	77–92
	SYLDAXVXEQLAT	Fce-Receptor II	298–310
	XXXHFVHQFQPFcyF	H-2A β^k	3–17
	QFQPFXYFTNT	H-2A β^k	10–20
H-2A ^{g7}	KPKATAEQLKTVMDD	Serum albumin	560–574
	GHNYVTAIRNQQEG	Transferrin	55–68
	ETTEESLRNYY \bar{E} Q	hnRNP B1 & A2	31–43
	VVMRDPAKRSR \bar{G} FGF	hnRNP A2 & B1	51–66
	VVMRDQTKRSRGFGF	hnRNP A1	44–59
	PKEPEQLRKLFIGGL	hnRNP A1	7–21
	VVYPWTQRYFDSF	β Globin major	33–45

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a: Rudensky et al. 1991; b: Rudensky et al. 1992; c: Hunt et al. 1992b; d: Nelson et al. 1992; e: Marrack et al. 1993; f: Reich et al. 1994

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