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Introduction

Lipids are important components of plants; they are part of cell structures,³ act as energy reserves,⁴ participate in cell signaling,⁵ mitigate stress tolerance⁶ and play a role in both symbiotic and pathogenic interactions.⁷ The entire lipid profile of an organism, a tissue or a cell, is known as the "lipidome",⁸ and the extensive study of lipid molecules including identification, quantification and elucidation of their role in biological systems is called "lipidomics".^{8,9} Lipids are highly diverse both in structure and in composition.^{8,10} Structural diversity arises from the differences in the chemical structures of lipids¹⁰ while the ratio of different lipids in cells, tissues, organelles, membrane leaflets, membrane subdomains or within an organism is known as compositional diversity.¹⁰

For many years, scientists have been studying lipids and their functions in biological systems. It has been shown that even small changes in the structure and composition of lipids can drastically affect vital biological functions.¹⁰ For example, changes in membrane lipids directly affect the function of membrane proteins and the physical properties of the cell membrane including fluidity and permeability.¹¹ Therefore, to correctly interpret the *in vivo* functions of lipids they must be studied as part of an integrated system with metabolites and

The state of the art in plant lipidomics

Cheka Kehelpannala, (1)*** Thusitha Rupasinghe,^b Thomas Hennessy,^c David Bradley,^c Berit Ebert^a and Ute Roessner^a

Lipids are a group of compounds with diverse structures that perform several important functions in plants. To unravel and better understand their *in vivo* functions, plant biologists have been using various lipidomic technologies including liquid-chromatography (LC)–mass spectrometry (MS). However, there are still significant challenges in LC-MS based plant lipidomics, which need to be addressed. In this review, we provide an overview of the key developments in LC-MS based lipidomic approaches to detect and identify plant lipids with emphasis on areas that can be further improved. Given that the cellular lipidome is estimated to contain hundreds of thousands of lipids,^{1,2} many of the lipid structures remain to be discovered. Furthermore, the plant lipidome is considered to be significantly more complex compared to that of mammals. Recent technical developments in mass spectrometry have made the detection of novel lipids possible; hence, approaches that can be used for plant lipid discovery are also discussed.

enzymes.¹² It is also important to determine not only the type of lipid present but also its time-dependent local concentration.¹²

One of the major challenges in uncovering the function of lipids at the cellular level is the existence of thousands of structurally diverse lipids, some of which may be present at trace level concentrations, thus limiting their detection and identification.¹² Consequently, a robust analysis workflow is necessary from sampling to extraction and subsequently to detection and identification of lipids. In this review, we discuss the key developments in liquid chromatography-mass spectrometry-based approaches for the extraction, detection, identification, quantification and discovery of plant lipids (Fig. 1). We conducted a comprehensive and in-depth survey of over 400 journal articles published on lipidomics from 1957 to 2021 to identify studies that have played pivotal roles in advancing plant lipidomics. Our survey indicated that many of the techniques in plant lipid analysis were adapted from lipidomic approaches developed for mammalian lipid analysis, which we have discussed in this review. We have also discussed the recently developed LC-MS based lipidomic approaches to analyse lipids in animal tissues, which can be adapted for plant lipid analysis in the future. The primary goal of this review is to provide an overview of the evolution of plant lipidomics, its current state and potential future developments with special focus on LC-MS based workflows.

Plant lipids

Lipids are classified into different groups by the LIPID MAPS (LIPID Metabolites And Pathways Strategy; https://www.lipid

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^a School of BioSciences, The University of Melbourne, Melbourne, VIC 3010, Australia. E-mail: clkehelpannala@gmail.com; Tel: +61 451821235

^b Sciex, 2 Gilda Ct, Mulgrave, VIC 3170, Australia

^c Agilent Technologies Australia Pty Ltd, 679 Springvale Road, Mulgrave, VIC 3170, Australia

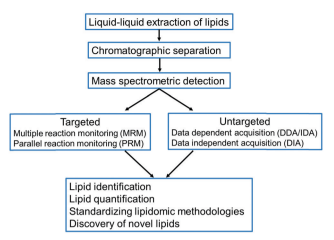


Fig. 1 The workflow applied in liquid chromatography-mass spectrometrybased lipidomics.

maps.org/) classification system based on the distinct hydrophilic and hydrophobic elements that comprise the lipid.¹³ The eight main groups include fatty acyls, glycerolipids, glycerophospholipids, sphingolipids, saccharolipids, polyketides, sterol lipids and prenol lipids, which can be distinguished by their chemically functional backbone structures (Table 1 and Fig. 2).13 The major plant lipids and their functions are summarized in Table 1.

Sample preparation

Given the highly diverse nature of lipid structures (Fig. 2), a comprehensive analysis of the lipidome requires firstly an efficient sample preparation process, then an effective lipid separation strategy followed by an optimal mass spectrometric detection method.33

The first step for lipid analysis is the successful extraction of lipids from the tissue of interest, for which a rapid, simple and efficient extraction method is desirable.³⁴ The most widely used sample preparation method in lipidomics is liquid-liquid extraction.³⁵ Most of the liquid-liquid plant lipid extraction methods are derived from extraction protocols developed for animal tissues, such as the Folch³⁶ and the Bligh and Dyer method.³⁷ In addition to these two, several other protocols using diverse mixtures of organic solvents, including hexane,³⁸

Lipid class	Core structure	Functions in plants	Ref.
Fatty acyls	Repeating series of methylene groups	Main building blocks of complex lipids. ¹³	Fahy <i>et al.</i> , 2005
		Function as signal transduction mediators. ¹⁴	Lim <i>et al.</i> , 2017
Glycerolipids	Glycerol backbone	Major constituents of the thylakoid membranes essential for photosynthesis. ^{15–17}	Dormann and Benning, 2002
		Important molecules in cell communication, signal	Holzl and
		transduction and pathogen responses. ¹⁸	Dormann, 2007
		Major storage reserves in plants. ¹⁹	Hölzl and
			Dörmann, 2019
			Cavaco <i>et al.</i> , 202 Yang and Bennin
			2018
Glycerophospholipids	Phosphate group esterified	Major constituents of cell membranes. ²⁰	Reszczyńska and
Jeer Free Free	to a glycerol backbone		Hanaka, 2020
		Act as signaling molecules. ²¹	Welti <i>et al.</i> , 2007
		Involved in plant defense mechanisms against pathogens. ¹⁸	Cavaco et al., 202
Sphingolipids	Sphingoid long-chain base backbone	Participate in cellular signaling, ²² growth and stress responses. ^{23,24}	Ali <i>et al.</i> , 2018
		Involved in cellular trafficking ²⁵ and defense responses against microbial pathogens ⁷	Hou et al., 2016
			Mamode Cassim
0		<u>Standard</u> 1	<i>et al.</i> , 2021
Sterol lipids	Fused four ring structure	Structural components of cell membranes. ²⁰	Reszczyńska and Hanaka, 2020
		Regulate acyl chain organization whereby the domain structures	Schaller, 2020
		of cell membranes are reinforced. ^{26,27}	Benanci, 2004
		Participate in plant stress responses. ²⁸	Dufourc, 2008
		Involved in resistance mechanisms against pathogens. ⁷	Rogowska and Szakiel, 2020
		Precursors of steroid hormones in plants. ²⁸	Siebers <i>et al.</i> , 201
		r i i i r	Rogowska and
			Szakiel, 2020
Prenol lipids	C ₅ unit	Participate in protective mechanisms supporting the adaptation	Baczewska et al.,
		of plants to environmental stresses. ^{29,30}	2014
Saccharolipids	Fatty acids linked directly	Involved in plant defense responses against insects and fungal	Bajda <i>et al.</i> , 2009 Luu <i>et al.</i> , 2017
accharonpius	to a sugar backbone	pathogen attacks. ³¹	Luu ci ui., 2017
Polyketides	Polyketide backbone	Responsible for the antibacterial, antifungal, antiviral and	Han <i>et al.</i> , 2018
	•	antiparasitic properties of plants. ³²	,

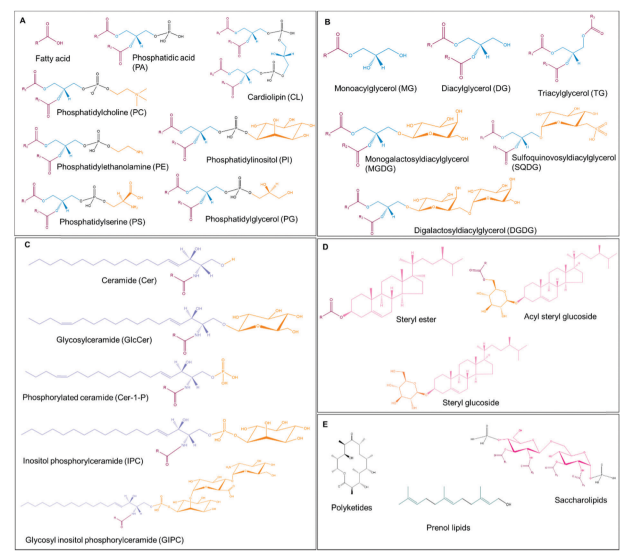


Fig. 2 Representative lipid structures. (A) Fatty acid and glycerophospholipids. The fatty acyl chains are shown in brown, glycerol backbone is indicated in blue, phosphate groups are in black and the head group is in orange. (B) Glycerolipids. The glycerol backbone is indicated in blue, fatty acyl chains are in brown and the hexose group is in orange. (C) Sphingolipids. The long-chain base is denoted in purple, fatty acyl chains are in brown and the head group is in orange. (D) Sterol lipids. The sterol group is indicated in pink, fatty acyl chains are in brown and the sugar moiety is in orange. (E) Polyketides, saccharolipids and prenol lipids. The polyketide backbone is shown in black and the five-carbon groups in prenol lipids are indicated in green. The sugar backbone in saccharolipids is shown in magenta, the phosphate group in black and the fatty acyl chains in brown. R indicates a repeating series of methylene groups. Lipid structures are based on the LIPID MAPS structure database and were generated using ChemDraw 20.0 software.

butanol³⁹ and petroleum ether,⁴⁰ and different treatments such as cold⁴¹ and heat⁴² have been developed, compared and evaluated for plant lipid extraction. Studies comparing these methods concluded that the Folch or Bligh and Dyer methods and those derived from them using chloroform, methanol and water as the extraction solvents are the most efficient methods for plant lipid extraction,^{34,42–45} with the Folch method³⁶ referred to as the "gold standard".⁴⁶ Originally, modifications to the Folch method³⁶ led to the development of the Bligh and Dyer method.³⁷ Later, significant improvements were made to the Bligh and Dyer protocol³⁷ when it was adopted for plant lipid extractions, leading to several new protocols. For example, in the study by de la Roche *et al.*⁴² wheat, *Triticum aestivum* L. cv, seeds were boiled with isopropanol to

inactivate the lipolytic enzymes and extract neutral lipids more efficiently before applying the Bligh and Dyer method.⁴² Ryu and Wang⁴⁷ simplified the protocol published by de la Roche *et al.*⁴² and introduced 0.01% butylated hydroxytoluene (BHT) to the extraction solvent to reduce lipid oxidation.⁴⁷ The Ryu and Wang⁴⁷ method was adapted and modified with additional steps by Welti *et al.*⁴⁸ and since then this method has been extensively used in plant lipidomics. Since the method by Ryu and Wang⁴⁷ was relatively labor-intensive, a shorter high-throughput single-step extraction method with a 24 h extraction period was developed by Vu *et al.*⁴⁹ Additional modifications to this single-step extraction method of Vu *et al.*⁴⁹ resulted in an extraction protocol that uses 30 parts chloroform, 25 parts isopropanol, 41.5 parts methanol, and 3.5 parts water (v/v/v)

with 0.01% BHT.³⁴ This protocol developed by Shiva et al.³⁴ has been shown to be highly efficient in extracting lipids from Arabidopsis thaliana and Sorghum bicolor leaf tissues.³⁴ To inactivate the lipolytic enzymes, fresh leaves were harvested directly into isopropanol pre-heated to 75 °C.42,50 However, this single step protocol³⁴ does not include steps for tissue homogenization. Tissue homogenization, however, is crucial to obtain a representative sample as it allows the extraction solvent to access lipids even in rigid plant tissues, increases the surface area in contact with the extraction solvent⁵¹ and solubilizes the lipids. While an extraction method should contain minimal sample preparation to achieve high-quality results, homogenization and metabolic quenching are vital steps.⁵² Thus, the efficiency of a lipid extraction method can be further improved through the addition of a mechanical grinding step.53 A recent comparison of four popular plant lipid extraction methods undertaken by us included the single step extraction method published by Shiva et al.,³⁴ the multistep extraction method of Welti et al.,⁴⁸ the biphasic extraction method using methyl tert-butyl ether (MTBE)⁴⁶ derived from the Folch method³⁶ and a rapid method employing an extraction temperature of 4 °C.⁴¹ This study revealed that overall, the single-step extraction method³⁴ is the most efficient for extracting most of the major lipid classes from Arabidopsis tissue. However, due to the diverse relative hydrophobicity of lipid molecules, the development of a single protocol efficient in extracting all lipid species in a plant sample is impossible.⁴⁵ For example, several studies have shown that chloroform/ methanol mixtures are not suitable for the extraction of plant sphingolipids such as glycosylinositol phosphorylceramides (GIPCs).^{34,54,55} Due to the highly amphiphilic nature of sphingolipids,⁵⁵ the standard lipid extraction methods are inefficient in solubilizing them.⁵⁴ It is therefore recommended to use a mixture of isopropanol, hexane and water for the extraction of plant sphingolipids^{55,56} as described by Markham et al.⁵⁴

Even though chloroform and methanol are considered ideal solvents for lipid extraction, they can have detrimental effects on human health due to their lipo-solubility.⁴⁵ Given the proven carcinogenicity of chloroform, attempts have been made to replace it with methyl-*tert*-butyl ether (MTBE), which extracts lipids into the organic layer while the polar compounds remain in the aqueous layer.⁴⁶ However, this biphasic extraction method is labor-intensive, which limits its applicability in large-scale lipidomic studies. Furthermore, significant technical errors may be introduced during the separation of the organic layer from the aqueous layer, thus ultimately affecting the extraction reproducibility.⁵³ Substituting the chloroform used in the single-step extraction protocol of Shiva *et al.*³⁴ with MTBE and comparing the extraction efficiencies are worth testing to further improve the existing extraction protocols.

Recent studies have focused on developing lipid extraction methods using environmentally friendly "green solvents", which can be organic, ionic liquids or supercritical fluids, such as carbon dioxide under supercritical conditions.⁴⁵ Supercritical fluids are substances that are not in a distinct gas or liquid phase but can be gradually compressed from low to high density.⁵⁷

By adjusting the density, its properties can be manipulated for the required process.⁵⁷ Organic green solvents are low in toxicity, readily biodegradable, easily recyclable and have high boiling points and low miscibility, while ionic liquids are nonaqueous organic salts, which are liquid at room temperature.^{45,58} The use of green solvents is widely explored for lipid extraction in biomass processing for biofuel production where large amounts of chemicals are routinely used.⁵⁹ Thus, there could be the potential to replace chloroform with one or some of these green solvents to efficiently extract plant lipids. For example, a biphasic system of cyclopentyl methyl ether (CPME):methanol: water was as efficient as the Bligh and Dyer method in extracting triacylglycerols from wet biomass of the oleaginous yeast *Lipomyces starkeyi*,⁶⁰ thus the study concluded that CPME can be used as an alternative to chloroform.⁶⁰ However, a comparative study applying the green solvents 2-methyl tetrahydrofuran (2-MeTHF) and cyclopentyl methyl ether (CPME) revealed that the Bligh and Dyer method was more effective in extracting total lipids from the microalgae Chlorella pyrenoidosa for biodiesel production.⁶¹ Although the potential of green solvents such as isoamyl acetate,⁶² terpenes, ionic liquids,⁵⁸ 2-methyltetrahydrofuran,⁶³ supercritical and subcritical fluids^{64,65} for lipid extraction from yeast and microalgae has been evaluated, their applicability for plant lipid extraction is yet to be demonstrated. Future work will require the comparison of the traditional plant lipid extraction methods with the green solvent-based extraction methods to evaluate the applicability of green solvents in routine plant lipidomic analysis.

Even though current plant lipidomic analyses rely on manual extraction of lipids, automated workflows for lipid analysis will be developed in the future. Automation for sample preparation would be beneficial since the quality, reproducibility and efficiency would be enhanced when compared with manual execution.

Liquid chromatography-mass spectrometry (LC-MS) in lipidomicsã

Electrospray ionization (ESI) mass spectrometers are the predominant instruments used for lipid analysis.⁶⁶ Lipid samples can be analyzed either by directly injecting the extracts into the mass spectrometer or by first separating them through chromatographic methods before injection into the mass spectrometer.¹ In 1994, Han and Gross⁶⁷ first proposed lipidomic analysis using ESI mass spectrometry, when they analyzed human erythrocyte plasma membrane phospholipids using a triple quadrupole with syringe pump sample injection. This method was further extended to characterize and quantify membrane lipids from cells or subcellular structures by Brügger et al. 1997,⁶⁸ using a nano-ESI source. This is now widely known as "shotgun lipidomics" and has seen continuous improvement over the past 25 years. The strategy used by Brügger et al. 1997⁶⁸ for the high-throughput profiling of complex lipids from unfractionated extracts of animal cell membranes was later adapted by Welti et al.48 to profile membrane lipids in Arabidopsis tissues.48,69

Because of the relative simplicity, high sample throughput and cost-effectiveness of the shot-gun lipidomic methods³⁴ developed by Welti and colleagues over the past 20 years,^{48,49,69–71} they are still widely used in plant lipidomic analysis.^{72,73} However, the application of direct infusion methods is not without drawbacks. They show higher ion suppression effects compared to methods with chromatographic separation, consequently hindering the analysis of low-abundant lipid species.¹ The selectivity of the direct infusion approach is limited by the existence of isobaric precursor and fragment ions.⁷⁴ The predominantly targeted analysis involving these methods makes the detection of unknowns impossible.⁷⁴ Due to these factors, the shot-gun approach was rapidly followed by LC-MS for plant lipidomics.⁷⁵

The LC-MS based methods were highly sensitive with reduced ion suppression and matrix effects,⁷⁶ able to detect novel lipids,⁷⁷ able to yield reliable identities of lipids even at low levels and able to separate isomers and isobaric compounds.⁷⁸ LC-MS is now considered a comprehensive and powerful technique for the analysis of complex lipid classes such as glycerolipids, glycerophospholipids, and glycolipids and is successfully used in plant lipid research.⁷⁹ To maximize the performance of the LC-MS systems, both the LC and MS methods must be meticulously optimized.33 An ideal LC method should possess high peak capacity, good chromatographic separation and resolution, reproducibility, selectivity and sensitivity in addition to being compatible with the data acquisition cycles in mass spectrometers.33 In general, lipidomic methods take long chromatographic analysis times to separate the large number of lipids, which are present over a wide concentration range in biological samples, and to separate isomeric and isobaric compounds.⁸⁰ Quicker chromatographic methods can be used; however, several limitations need to be addressed such as analyte overlap, isotopic interferences and ion suppression effects.⁸⁰ To mitigate these effects, ion mobility can be introduced into the analytical workflow. For example, a separation method employing an elution time of 3.7 min per sample has resulted in a reduced number of features compared to longer chromatographic methods.⁸⁰ The introduction of ion mobility to the analytical workflow, increased the peak capacity and helped to resolve co-eluting species, thereby increasing the number of detected features.80

The LC technologies used in modern lipidomics include reversed-phase chromatography (RPLC), normal phase chromatography (NPLC), hydrophilic interaction liquid chromatography (HILIC) and supercritical fluid chromatography (SFC).⁷⁶ RPLC is one of the most often used techniques utilizing a nonpolar stationary phase and a polar mobile phase where lipid molecules are separated based on their hydrophobicity, length of fatty acyl chain, and number and position of double bonds.⁷⁶ In general, the retention time of lipids increases with increasing carbon number and decreases with increasing number of double bonds. For example, PC 34:6 elutes before PC 36:6 while PC 34:5 elutes after PC 34:6 (Fig. 3). Non-polar lipids, such as TGs are longer retained in the column while polar lipids such as lysophospholipids elute faster.

The stationary phase of RPLC consists of silica microparticles containing hydrophobic functional groups such as C_{30} , C_{18} , C_8 , C_4 hydrocarbon chains or ligands, which may be cyano, phenyl or amino derivatives with C_{18} being the most widely used.⁷⁶ The polar mobile phase generally contains water and a water-miscible organic solvent such as acetonitrile, methanol and isopropanol.⁷⁶ The development of ultrahigh performance liquid chromatography (UHPLC) columns and core–shell materials for columns significantly improved traditional HPLC techniques providing higher resolution and requiring less time.⁸¹

NPLC separates lipid classes based on polarity and consists of a polar stationary phase and a non-polar mobile phase such as hexane or heptane with a miscible organic solvent like chloroform, isopropanol or ethylacetate.⁷⁶ NPLC can only separate non-polar lipids like DGs and TGs while phospholipids cannot be separated using this method.⁷⁶ In addition, NPLC methods are lengthy and the highly non-polar organic solvent systems used in NPLC are incompatible with ESI-MS.⁸² HILIC is a more efficient type of NPLC technique that uses less apolar solvent systems compatible with ESI-MS and separates lipid classes based on their head group polarity.82 The polar stationary phase consists of silica or functionalized silica while the mobile phase consists of water and water-miscible organic solvents.⁷⁶ A comprehensive description of the theories behind different chromatographic approaches has been recently published by Lange et al. 2019.⁷⁶ HILIC and RPLC are currently the

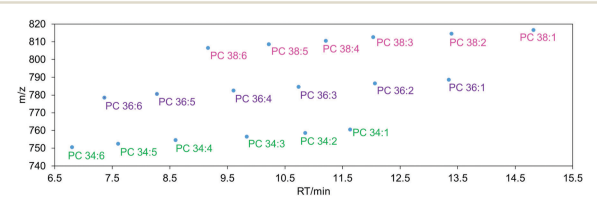


Fig. 3 The elution order of phosphatidylcholine (PC) 34, 36 and 38 carbon lipid species in reverse-phase liquid chromatography (RPLC).

two most widely applied chromatographic separation techniques in LC-MS based lipidomics including plant lipidomics.^{78,83} A direct comparison of HILIC-MS and RPLC-MS based lipidomic workflows concluded that both can be equally used to accurately quantify LPCs, LPEs, PCs, PEs and sphingomyelins in human blood plasma;⁸³ however, no such comparison has been reported for lipids from plant tissues. An evaluation of HILIC-MS and RPLC-MS based workflows for plant lipid analysis will be of great benefit for future studies on plant lipids.

In addition to traditional one dimensional LC methods, recently two dimensional LC (2DLC) methods have also been investigated for high-throughput lipidomic analysis of various samples including human plasma⁸⁴ and plant tissues.⁸⁵ In LC, only one liquid phase separation system is applied to the analytes in a sample while 2DLC techniques utilize two orthogonal liquid phase separation systems to separate sample components.⁸⁶ A study comparing several LC and 2DLC methods coupled with high-resolution time-of-flight mass spectrometry (ToF-MS) revealed that the C18xHILIC approach was the most effective in comprehensive lipid profiling of extracts of zebrafish embryos.⁸⁷ However, the data generated by 2DLC coupled with mass spectrometry are highly complex and make an untargeted analysis challenging,⁸⁵ consequently, advanced chemometric data analysis has been proposed for such studies.85 For example, a 2DLC setup combining RPLC and HILIC coupled to a triple quadrupole (QQQ) mass spectrometer with chemometric data analysis was able to evaluate the effects of arsenic exposure on the rice lipidome, resolving a larger number of lipids compared to the LC system.85 However, the identification of lipids was difficult due to poor detection sensitivity, likely caused by the dilution due to two successive chromatographic separations or because it was impossible to optimize the collision energy.⁸⁵ While the application of 2DLC systems in routine lipidomic analysis may be challenging,³ where simple, rapid, user-friendly and proven lipidomic data acquisition strategies are required, future developments particularly in data processing workflows may improve its applicability. It is worthwhile to also investigate if the 2DLC approach can be used to discover novel lipids.

The separation of lipids by LC is followed by transfer to a mass spectrometer. Mass spectrometers have been used in the analysis of lipids since the 1980s⁸⁸ and over the past decades have seen substantial improvements in the instrumentation and data acquisition methodologies. Mass spectrometric analysis of complex biological samples can be performed using untargeted or targeted approaches.89 Untargeted methods aim to comprehensively profile all detectable compounds in a sample while targeted methods focus on analyzing a predefined set of metabolites.⁸⁹ In general, targeted methods are more sensitive and more quantitative, however, only a limited number of compounds can be analyzed.⁸⁹ On the other hand, untargeted methods can provide an unbiased analysis of the changes in a system and allow the discovery of novel lipids.⁹⁰ While targeted methods require prior knowledge of sample matrix and LC-MS properties of the analytes, untargeted methods do not rely on prior information about the sample matrices and

are ideal for discovery-focused lipidomic approaches.⁸³ However, due to the complexity of the data retrieved using untargeted methods and the unspecific nature of the analysis, some analytes may go undetected. Furthermore, targeted MS conditions can specifically be tuned to detect individual compounds, for example through the selection of optimized collision energies, while this option is rather limited for the discovery approach.

The commonly used mass spectrometers in lipidomics are triple quadrupoles (QQQ), quadrupole ion trap (QIT), triple quadrupole-linear ion trap (QTRAP), orbitraps and quadrupole time-of-flight (Q-ToF) instruments.35,91 Instruments such as Q-ToF and Orbitrap provide high mass accuracy at MS and MS/MS levels and offer both accurate identification and guantification of lipids, although at a lower sensitivity.^{89,90} In contrast, OOO mass spectrometers show low mass accuracy but offer high sensitivity when operated in the multiple reaction monitoring (MRM) mode.^{90,91} This makes them the preferred instruments for quantitative targeted studies including the analysis of lowabundant lipid species.^{90,91} However, parallel reaction monitoring also known as MRM high-resolution (MRM^{HR}) performed on a high-resolution instrument such as Q-ToF is a more comprehensive, targeted quantitative strategy where all detectable product ions of a pre-selected precursor ion are scanned in the highresolution mode.⁹² In parallel reaction monitoring a scheduled algorithm is used, in which a targeted/predefined set of precursor ions are isolated in the quadrupole followed by fragmentation and subsequent detection of all the product ions.⁹² This yields more accurate m/z and narrower peak widths, and simultaneously detects up to 100 precursor ions per duty cycle; the intensities of multiple fragments can be summed for better sensitivity.92 Untargeted LC-MS based lipidomic analyses were found to be independent of the type of high-resolution mass spectrometer used and led to nearly identical results when lipidomic data acquisitions across nine different platforms (one single ToF, one Q/orbital ion trap, and seven Q-ToF instruments) were compared.93 This suggests that the up- and down-regulation patterns of lipids in biological samples can be studied using any type of mass spectrometer.³⁵

MS-based lipidomic analyses utilizing high-resolution mass spectrometers can apply data-dependent acquisition (DDA) or data-independent acquisition (DIA) strategies.⁸⁰ With DDA methods, full scan spectra are recorded at the MS level and MS/MS spectra are recorded by automatically fragmenting the precursor ions with an intensity above a certain threshold or as specified by a user-defined precursor list,³⁵ a given mass range or by mass defect filters.⁹⁴ Since the precursor ions for fragmentation are selected based on the MS survey scan, the DDA approach is also known as information-dependent acquisition (IDA),⁹⁴ data directed acquisition or data-directed analysis.⁹⁵ Generally, the most abundant 10-20 precursor ions are selected for fragmentation using the DDA approach,⁹⁴ which is also known as the top-N data-dependent MS/MS approach⁹⁶ with N referring to the number of the most abundant precursor ions used for fragmentation. Therefore, MS/MS spectra of lowintensity ions will be excluded³⁵ and the reproducibility of the data is low.94 To increase the coverage of low abundant lipids using the DDA approach, an iterative exclusion is proposed where precursors previously selected for fragmentation are excluded in subsequent injections.^{97,98} The application of the iterative exclusion to DDA workflows has shown to increase lipid identification by over 50% in some cases.⁹⁸

By contrast, the DIA methods collect spectral data of all fragments without the selection of specific precursor ions, yielding highly complicated spectra.⁹⁴ DIA methods include collision energy switching MS of everything (MS^E) also known as MS^{All} or all ion fragmentation $(AIF)^{35,99}$ and sequential window acquisition of all theoretical fragment ion spectra (SWATH).¹⁰⁰ In the MS^{All} approach, the data-dependent acquisition is performed on all product ions regardless of the precursor ion. As all precursor ions are sent to the collision cell for fragmentation, MS^{All} relies on good chromatographic separation techniques, such as ultrahigh-performance liquid chromatography (UHPLC) to obtain quality MS/MS spectra.⁹⁵

To reduce the complexity of the spectra generated using the MS^{All} approach, SWATH was introduced by Gillet et al.¹⁰⁰ for proteomic experiments and was quickly adapted for metabolomic and lipidomic studies.94 However, SWATH still provided complex spectral data and special software tools were necessary to deconvolute the data.³⁵ This issue was solved when an open-source data processing software called mass spectrometry-data independent analysis (MS-DIAL) software¹⁰¹ was introduced to deconvolute the data accumulated by SWATH, which incorporated the LipidBlast¹⁰² library to identify lipids.¹⁰¹ At present, the SWATH acquisition method is immensely popular for untargeted lipidomic analyses¹⁰³ and has been used to analyze lipids in human platelets,⁹⁴ mouse plasma,¹⁰⁴ and human plasma.¹⁰⁵ More recently it was adapted for plant lipidomics to analyse lipids from diverse Arabidopsis tissues.¹⁰⁶

SWATH is considered to be advantageous over the DDA methods as it offers full coverage of all MS/MS fragments, yields more lipid identifications and shows high sensitivity, which is comparable to MRM reactions on a QQQ.⁹⁹ Furthermore, the SWATH approach extracts chromatograms not only at the MS level but also at the MS/MS level whereby reliable quantification can be achieved at either the ToF-MS or SWATH-MS/MS levels using precursor or product ions, respectively.¹⁰⁴ A comparison of the IDA, SWATH and MS^{AII} approaches revealed that the IDA methods yield qualitatively better MS/MS spectra but a lower number of lipid annotations, while SWATH outperformed the MS^{AII} approach with better quality MS/MS spectra and identical MS/MS acquisition hit rates.⁹⁵

Here, it should be noted that the method of choice for lipidomics depends on the purpose of the study⁹⁵ and available resources (Table 2). The IDA approaches such as SWATH and MS^{All} are ideal for untargeted lipidomic approaches.^{99,106,107} However, while SWATH can be used for quantitative analyses,⁹⁹ lipid identification and quantification may be far more simple with targeted MRM methods where quantification of a targeted set of lipids is the primary focus.

Lipid identification

High-throughput untargeted data acquisition strategies on high-resolution mass spectrometers generate a large amount of spectral data,¹⁰⁸ which makes the accurate annotation of the mass spectrometric features challenging. Since the launch of the LipidBlast library in 2013,¹⁰² over 25 commercial and opensource software tools have been developed to process data and identify lipids.¹⁰⁸ However, there are currently no lipid libraries developed specifically for the identification of plant lipids. A customized library for plant lipid identification will be of great value for plant lipid researchers.

Most of the developed software tools use in silico lipid libraries, which can be rapidly generated by computing all possible combinations of fatty acids, head groups, linkages and backbones with fragments predicted using a simple set of rules.¹⁰⁸ Commercial software tools include for example SimLipid (PREMIER Biosoft), Lipid Search[™] (Thermo Fisher Scientific),¹⁰⁹ Lipid Annotator software⁹⁷ (Agilent Technologies), and LipidyzerTM platform¹¹⁰ (Sciex) while freely available openaccess tools include lipid data analyzer 2 (LDA2),¹¹¹ Lipid-Blast,¹⁰² Competitive Fragmentation Modeling-ID (CFM-ID),¹¹² LIQUID,¹¹³ MS-DIAL,¹⁰¹ Greazy,¹¹⁴ lipidr,¹¹⁵ LipiDex,¹¹⁶ LipidMS¹¹⁷ and LipidMatch Flow.¹¹⁸ From these tools, MS-DIAL is the most commonly used open-source software, which is compatible with many LC-MS workflows.¹⁰⁸ The acquisition modes, data processing pipelines and features of the abovementioned LC-MS based lipidomic software tools have recently been reviewed and compared by Züllig and Köfeler, 2020³⁵ and Züllig et al., 2020.¹¹⁹ Despite all these software solutions, unambiguous structure identification is still a challenging task, mostly due to the over annotation of lipid identities that cause false-positive results and demand manual correction of the lipid annotations.^{35,120} This problem may be circumvented by developing a lipid library where the lipid identities have been manually curated and validated at least for a model organism such as Arabidopsis. Recently, we have compiled a manually curated database containing over 800 lipids in Arabidopsis tissues with information on the precursor m/z values, adducts, product ions, characteristic fragmentation patterns and retention times.⁵³ The information therein can be used to confirm lipid identities and develop targeted MS methods for lipid analysis.

Lipid quantification

Qualitative analyses focus on the profiling and comparison of lipids in biological samples while quantitative analyses aim to provide the concentrations of specific lipids in a biological matrix using meticulously developed quantification methods and lipid standards.⁸⁰ Qualitative comparisons between biological samples use a limited number of internal standards or no standards at all; instead, the peak intensities or area of detected lipids are compared.² This approach is generally known as lipid profiling and can provide comprehensive comparisons between

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	Ref.	Welti <i>et al.</i> , 2002, Devaiah <i>et al.</i> , 2006 ^{150,151}	Markham and Jaworski, 2007 ⁵⁵	Hummel <i>et al.</i> , 2011 ¹⁵²	Burgos <i>et al.</i> , 2011 ⁴¹	Okazaki <i>et al.</i> , 2013a, Oka- zaki <i>et al.</i> , 2013b ^{148,153}
	Lipids identified	Phospholipids, polar glycerolipids	Sphingolipids	Phospholipids, oxidized lipids, glycerolipids, sphingolipids (less comprehensive compared to other lipid classes)	Phospholipids, galactolipids	Novel plant lipid, glucur- onosyldiacylgłycerol, phospholipids, polar głycerolipids, sterol lipids, sphingolipids
	Internal standards	di14:0-PC, di24:1-PC, 13:0- LPC, 19:0-LPC, di14:0-PE, di24:1-PE, 14:0-LPE, 18:0- LPE, di14:0-PG, di24:1-PG, 14:0-LPG, 18:0-LPG, di14:0- PA, di20:0 (phytamoyl)-PA, di14:0-PS, di20:0 (phytamoyl)-PA, pS, 16:0-18:0-0 (phytamoyl)- PS, 16:0-18:0-0 (di18:0-PI, 16:0-18:0-0 di18:0-DGDG and di18:0-DGDG and	GM1, 12-GlcCer, C12-Cer, sphingosine (C17 base), and sphingosine1-phosphate (C17 base)	PE 34:0 (17:0, 17:0) and PC 34:0 (17:0, 17:0)	Not given	10:0/10:0 PC
	Lipid identification	MRM transitions	RT of known standards and by comparison with other peaks	Matching the <i>m/z</i> values with previously reported data	Targeted search based on <i>m</i> /z values, expected RT and manual inspection of MS/MS data	<i>m/z</i> values and RT of authen- tic compounds
	MS conditions	QQQ in positive and negative modes	QTRAP using MRM mode in positive and negative modes	Orbitrap with full scan and all-ion frag- mentation in positive and negative modes	SYNAPTTM using MS ^E in negative ion mode	IT-TOF oper- ated at scan mode in negative ion mode
ant lipid analysis	LC conditions	N/A	SUPELCOSIL ABZpPlus column (150 mm \times 3 mm \times 5 µm) A: THF: MeOH: 5 mM a.f. (3:2:5) + 0.1% f.a. B: THF: MeOH: 5 mM a.f. (7:2:1) + 0.1% f.a.	C8 RP column (100 mm \times 2.1 mm \times 1.7 µm) A: 1% 1 M NH ₄ Ac +0.1% a.a. B: ACN : IPP (7:3) + 1% 1 M NH ₄ Ac +0.1% a.a.	C8 RP column (150 mm \times 2.1 mm \times 1.8 µm) A: 1% 1 M NH ₄ Ac +0.1% a.a. in H ₂ O B: MeO- H:IPP (5:2) + 1% 1 M NH ₄ Ac, 0.1% a.a.	HILIC silica column (100 mm $\times 2.1$ mm $\times 3$ µm) A: MeOH : H ₂ O (95 : 5) + 0.2% a.f. B: ACN : MeOH : H ₂ O (95 : 2 : 3) + 0.2% a.f.
Mass spectrometry-based workflows used for plant lipid ar	Lipid extraction method	Extracted with CHCl ₃ : H ₂ Ofollowed by a five-fold extrac- tion with a mixture of CHCl ₃ : MeOH (2:1). The solvent was eva- porated under nitro- gen and resuspended in CHCl ₃	Extracted with IPP: hexane: H_2O (55:20:25). The dried extract was de-esterified using methylamine, dried under nitrogen and dissolved in THF: MeOH: H_2O (2:11:2) containing 0.1% f.a.	Extracted with MeOH: MTBE $(1:3)$ followed by H_2O : MeOH $(3:1)$. Upper organic phase was removed, dried in a speed-vac and resus- pended in a buffer		Bligh and Dyer method with slight modifications. Extracted with CHCl ₃ : MeOH : H ₂ O and the lower layer evaporated by concentration and reconstituted in CHCl ₃
spectrometry-base	Homogenization method	None	The glass plun- ger of the DUALL tissue grinder, attached to a Ryobi D45CK power drill	Retsch mill	Powdered using a cryogenic grind- ing robot	Mixer mill
Table 2 Mass	Plant tissue	Arabidopsis thaliana rosettes, leaves, flowers, flower stalks, siliques, roots, and seeds	Arabidopsis thaliana leaves	Arabidopsis thaliana leaves	Arabidopsis thaliana rosettes	Arabidopsis thaliana leaves and roots. leaves of rice (Oryza sativa),

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Table 2 (con	(continued)							
Plant tissue	Homogenization method	Lipid extraction method	LC conditions	MS conditions	Lipid identification	Internal standards	Lipids identified	Ref.
Arabidopsis thaliana leaves	Mixer mill cooled with liquid nitrogen	Extracted with a hot solvent mixture containing 60% IPP	ACQUITY UPLC HSS T3 column (100 mm \times 1 mm \times 1.8 µm) A: MeOH : 20 mM NH ₄ Ac (3:7) + 0.1% a.a. B: THF:MeOH : 20 mM NH ₄ Ac (6:3:1) + 0.1%	QTRAP using MRM in positive and negative modes	MRM transitions	Not given	Phospholipids, sphingolipids, sterol lipids, glycerolipids	Tarazona et al., 2015 ⁷⁵
Arabidopsis thaliana leaves	None	Extracted with CHCl ₃ : MeOH : 300 mM NH ₄ Ac in H ₂ O (30 : 41.5 : 3.5)	N/N	QQQ in positive and negative modes	MRM transitions	A mixture of 22 internal standards	Phospholipids, glycer- olipids, acylated galactolipids, sterol lipids	Shiva <i>et al.</i> , 2020 ⁷²
Arabidopsis thaliana leaves and roots. Mature leaves and new shoots of <i>Camellia</i> <i>sinensis</i> (tea) plants.	Lyophilized sam- ples were homo- genized in a Retschmill	Modified Matyash et al., 2008 method. Extracted with MeOH: MTBE: H_2O (1:3:1) followed by MeOH: water (1:3). Upper phase was dried down in a speed vac and resuspended in	C8 column (100 mm × 2.1 mm × 1.7 μ m) A: H ₂ O + 1% 1 M NH ₄ Ac + 0.1% a.a. B: ACN + IPP (7:3)+ 1% 1 M NH ₄ Ac + 0.1% a.a.	tive ull scan i in e and e	In-house library	None. Content of each lipid species is presented as mean intensity	Phospholipids, galactolipids and sphingolipids	Giavalisco et al., 2011, Liu et al., $2017^{154,155}$
Barley (<i>Hor-</i> deum vulgare L.) roots	Ground to a fine powder using a mortar and pes- tle in liquid nitrogen	ALM: ILP? $(/:3)$ Sphingolipid extrac- tion in plant tissues as described by Markham <i>et al.</i> , 2006. Extracted with IPP : hexane : H ₂ O (60 : 26 : 14), evapo- rated under liquid nitrogen and resus- pended in IPP:	Poroshell EC-C18 col- umn (100 mm \times 2.1 mm \times 2.7 µm) A: MeOH:20 mM NH ₄ Ac (3:7) B: IPP:MeOH:20 mM NH ₄ Ac (6:3:1)	QToF using SPRM in positive and modes. Oxidized lipids were analyzed based on precursor- production	Precursor- product ion transitions	PE(12:0/12:0) and Cer(d18:1/12:0)	Oxidized lipids, phospho- lipids, polar and neutral glycerolipids, sterol lipids, sphingolipids	Yu <i>et al.</i> , 2018, Yu <i>et al.</i> , 2020 ^{92,156}
Arabidopsis seeds	Not given	MeOH: H_2O (4: 4: 1) Extracted with IPP: hexare: H_2O (55: 20: 25) accord- ing to the protocol by Markham <i>et al.</i> , 2006 followed by alkaline hydrolysis, dried down and recon- stituted in BuOH: MEOH (1: 1)	Ascentis Express RP Amide column (50 mm \times 2.1 mm \times 2.7 µm) H ₂ O:MCOH: THF (50:20:30 to 5:20:75)	transitions. QQQ using MRM transitions	MRM transitions	16: 0-d31 SM	Ceramides, hydro- xyceramides, glucosylceramides and GIPCs	Ebert <i>et al.</i> , 2018 ⁵⁶

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Review

Table 2 <i>(con</i> t	(continued)							
Plant tissue	Homogenization method	Lipid extraction method	LC conditions	MS conditions	Lipid identification	Internal standards	Lipids identified	Ref.
Rice (Aerial parts and roots)	Ground to a fine powder using a mortar and pestle in liquid hitrogen and lyophilized for 24 h to dryness	Modified Matyash et al. 2008 method extraction with pre- cooled MeOH: MTBE (1:3) followed by MeOH: water (1:3), upper-phase dried down and resus- pended in MeOH: water (1:0.85)	Column 1: RP ZORBAX Eclipse XDB-C18 col- umn (150 mm \times 2.1 m \times 5 µm) A: ACN : IPP (1:2) + 0.1% f.a. B: H ₂ O + 0.1% f.a. B: H ₂ O + 0.1% f.a. Column 2: KINETEX HILIC col- umn (30 mm \times 3 mm \times 8.5 µm NH-AC	QQQ in full scan mode followed by MRM transi- tions in negative and positives modes.	Matching the experimental MS/MS spectra with the refer- ence spectra available in Metlin	17:1 LPC, 17:0 LPC, 1,3- 17:0 D5 DG, 17:0 CE, 1,2,3- 17:0 TG, 16:0 D31-18:1 PC, 16:0 D31-18:1 PS, <i>N</i> - dodecanoylsphingosine, <i>N</i> -dodecanoylglucosyl- sphingosine and <i>N</i> - sylphosphorylcholine.	Phospholipids, glycerolipids	Navarro-Reig et al., 2018 ⁸⁵
Halophytes: Salicornia ramosissima (fresh branch tips) and Halimione portulacoides (leaves)	Ground to a powder	Bligh and Dyer protocol with modifications. Extracted with CHCl ₃ : MeOH (1:2) followed by the addi- tion of water. The lower phase was dried under nitrogen and resuspended in mobile phase B	Ascentis Si column (150 mm \times 1 mm, 3 µm) A: ACN : MeOH : H_2O (50: 25 : 25) + 1 mM NH_4AC B:ACN : MeOH (60: 40) + 1 mM NH_4AC	Q-Exactive with full scan MS and DDA in positive and negative modes	MS/MS spectral data	None	Phospholipids, glycolipids and glycosphingolipids	Maciel <i>et al.</i> , 2018 ¹⁵⁷
Wheat leaves	Cryo- homogenization (Cryomill)	Folch method with modifications. Extracted with MeOH, CHCl ₃ vacuum dried and resuspended in BuOH: MeOH (1:1)	Ascentis Express RP- Amide column (50 mm \times 2.1 mm, 2.7 μ m) A: 10 mM a.f. in H ₂ O: MeOH: THF (50: 20: 30) B: 10 mM a.f. in H ₂ O: MeOH: THF (5: 20: 75)	QQQ in positive and negative modes	MRM transitions	LPC(17:0), PC(34:1), PE(34:0), PG(34:1), PI(36:2), PS(34:0).	Phospholipids	Cheong <i>et al.</i> , 2019, Cheong <i>et al.</i> , 2020 ^{158,159}
Vascular plants (kale leaves and corn roots)	Cryo- homogenization (Cryomill)	Bligh and Dyer method. Samples were incubated in hot IPP prior to extraction. Extracted with MeOH and CHCl ₃ followed by the addition of H ₂ O. The bottom organic layer was dried under a nitrogen stream and reconstituted in CHCl ₃ : MeOH (1:1)	HILIC column (Luna 100 mm \times 2 mm \times 3 µm) to screen for any novel lipid classes A: ACN: H ₂ O (97:3) + 10 mM NH ₄ Ac B: 10 mM NH ₄ Ac in pure H ₂ O C30-RPLC (150 mm \times 2 × 2.6 µm) to confirm the HILIC results A: ACN: H ₂ O (60:40) +10 mM a.f. + 0.1% f.a. B: IPP:ACN: H ₂ O (90:10:1) + 10 mM a.f. + 0.1% f.a.	Q-extractive orbitrap using a top-20 DDA in positive and negative modes	MS/MS frag- mentation patterns	SQDG 16:0/16:0, MGDG 16:3(72,102,132)/ 18:3(92,122,152), PA 18:1(92)/18:1(92), PG 18: 0/20:4(52,82,112,142), PC PE 18:0/ 20:4(52,82,112,142), diether PC 0-18:0/0-18:0, plasma- logen PC P-18:0/ 20:4(52,82,112,142), mMPE logen PC P-18:0/ 20:4(52,82,112,142), MMPE 20:4(52,82,112,142), MMPE 20:4(52,82,112,142), MMPE 20:4(52,82,112,142), MMPE 20:4(52,82,112,142), MMPE 16:0/16:0 and DMPE 16: 0/16:0, LPA 20:4(52,82,112,142), MMPE 18:1(92), LPE 18:0, Plasma- logen LPE P-18:0; Pl 18:0/ 20:4(52,82,112,142), MMPE 18:2(92,122)/18:2(92,122), and SM d18:1/18:0.	Phospholipids, polar and neutral glycerolipids, phytosphingolipids, glycolipids. Acylpho- sphatidylglycerols, a modified class of acylpho- sphatidylglycerols and their isomeric forms in con roots. Regioisomers of lysophospholipids in kale leaves	Pham <i>et al.</i> , 2019 ⁹⁶

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Table 2 (con	(continued)							
Plant tissue	Homogenization method	Lipid extraction method	LC conditions	MS conditions	Lipid identification	Internal standards	Lipids identified	Ref.
Root, stem, leaf, and petiole tis- sues of Rapeseed	Intact plant tis- sues were used without homogenization	Plant tissue har- vested directly into hot IPP and extracted with $CHCl_3$: H_2O as described by Welti <i>et al.</i> , 2002	Acquity UPLCTM BEH C18 column (100 mm \times 2.1 mm \times 2 µm) A: H ₂ O:MeOH: ACN: 300 mM NH ₄ Ac (20: 20: 20: 1) B: IPP: MeOH: 300 mM NH ₄ Ac (100. 00. 3)	TripleTOF using DDA and MS/MS ^{ALL} scan in positive and negative modes	RT, accurate m/z , and frag- mentation ion patterns	PC-12: 0/12: 0, PA-14: 0/14: 0, PE-12: 0/12: 0, PG-20: 0/20: 0, PI-16: 0/18: 0, PS-20: 0/20: 0, LPC-19: 0, LPG-18: 0, LPE-18: 0, MGDG18: 0/18: 0, DGDG- 18: 0/18: 0, TAG-17: 0/17: 0/17: 0, DMC-17: 0/17: 0/17:	Phospholipids, glycerolipids	Lu <i>et al.</i> , 2019 ¹²⁰
Salad (Lac- tuca sativa var. capitata nidus tenerimma), deep frozen Spinach (Spinach (Spinach oleracea), raspberries (Rubus idaeus) and	Salad was manu- ally cut into small pieces and homogenized using an ultra- turax. Raspber- ties and straw- berries were homogenized using a hand blender	Extracted with IPP, n -hexane and H_2O according to the pro- tocol by Markham <i>et al.</i> , 2006, followed by alkaline hydro- lysis, dried down and reconstituted in IPP: H_2O (65:35)	(100:20:3) HSS T3 column (150 mm \times 2.1 mm \times 1.8 µm) A: ACN: H ₂ O (3:2) + 0.1% f.a. + 10 mM a.f. B: IPP: ACN (9:1) + 0.1% f.a. + 10 mM a.f.	Q-Exactive using top-10 DDA in posi- tive and nega- tive modes	Lipid Data Analyzer and manual inspection of data.	(d18:1/16:0) (d18:1/16:0)	GIPC	Panzenboeck et al., 2020 ¹⁶⁰
(<i>Fragaru</i>) Green alga, <i>Arabidopsis</i> tissues, maize, tomato and grey poplar and common beech trees	Bead beater	Extracted with pre- cooled MeOH: MTBE (1:3) solution fol- lowed by MeOH: water (1:3) extrac- tion. The upper- phase was dried down and resus- pended in ACN: IPP (70, 20)	C8 column (100 mm × 2.1 mm × 1.7 μ m) A: H ₂ O: IPP: ACN (446 g:130 g:300 g)+ 10 ml 1 M NH ₄ Ac+ ml a.a. B: IPP: ACN (236 g:546 g) + 10 ml 1 M NH ₄ Ac + 1 ml a.a.	Q-Exactive with full scan mode in positive and negative modes	Matching the exact <i>mlz</i> and RT to a list of potential compounds	PC 34:0	Phospholipids, glycerolipids	Lapidot- Cohen <i>et al.</i> , 2020 ¹⁶¹
Banana fruit peel	Crushing	Extracted with MeOH Extracted with MeOH followed by MTBE and H_2O . The upper phase was dried with nitrogen stream and resursanded in IDD	C18 column (100 mm \times 2.1 mm \times 1.7 µm) A: ACN: H ₂ O (6:4) + 10 mM a.f. B: ACN: IPP (1:9) + 10 mM a.f.	Q-Exactive using full scan MS in positive and negative modes	LipidSearch	None. Content of each lipid species is presented as mean intensity	Plycerophospholipids, saccharolipids, and glycerolipids	Liu <i>et al.</i> , 2020 ¹⁶²
Arabidopsis thaliana leaves, rosettes, internodes, roots, seed- lings, seeds and siliques	Cryo- homogenization (Cryomill)	Extracted with CHCl ₃ : MeOH: 300 mM NH ₄ Ac in H_2O (30:41.5:3.5), dried down in a speed vac and resuspended in BUOH:MeOH (1:1)	InfinityLab Poroshell C18 column (100 mm \times 2.1 mm \times 2.7 µm) A: ACN:H ₂ O (6:4) + 10 mM a.f. B: IPP:ACN (9:1) + 10 mM a.f.	QToF using SWATH in positive mode	MS-DIAL, <i>m/z</i> values, RT pattern and man- ual inspection of the MS/MS spectra	None	Phospholipids, glycerolipids, sphingolipids	Kehelpannala <i>et al.</i> , 2020, Kehelpannala <i>et al.</i> , 2021, Schillaci <i>et al.</i> , 2021 ^{53,106,163}
RT: retention quadrupole t dependent ac BuOH: butan	time, THF: tetrahy ime-of-flight mass (quisition, SWATH: ol ACN: acetonitrile	drofuran, a.a. acetic ac spectrometer, QQQ: trip sequential window acq , IPP: isopropanol, MTF	RT: retention time, THF: tetrahydrofuran, a.a: acetic acid, a.f.: ammonium forms quadrupole time-of-flight mass spectrometer, QQQ: triple quadrupole mass spec dependent acquisition, SWATH: sequential window acquisition of all theoretical BuOH: butanol ACN: acetonitrile, IPP: isopropanol, MTBE: methyl- <i>tert</i> -butyl-ether.	ate, f.a.: formic a ctrometer, SPRM l fragment ion sp r.	acid, NH₄Ac: am : scheduled para bectra, GIPC: gly	RT: retention time, THF: tetrahydrofuran, a.a: acetic acid, a.f.: ammonium formate, f.a.: formic acid, NH ₄ Ac: ammonium acetate IT-TOF: Ion trap-time-of-flight mass spectrometer, QToF: quadrupole time-of-flight mass spectrometer, QQQ: triple quadrupole mass spectrometer, SPRM: scheduled parallel reaction monitoring, MRM: multiple reaction monitoring, DDA: data dependent acquisition, SWATH: sequential window acquisition of all theoretical fragment ion spectra, GIPC: glycosyl inositol phosphoryl ceramide, CHCl ₃ : chloroform, MeOH: methanol, BuOH: butanol ACN: acetonitrile, IPP: isopropanol, MTBE: methyl- <i>tert</i> -butyl-teher.	rap-time-of-flight mass spec 4: multiple reaction monito nide, CHCl ₃ : chloroform, M	trometer, QToF: rring, DDA: data eOH: methanol,

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biological samples when combined with substantive statistical analyses.^{2,121–126}

For the absolute quantification of lipids, ideally, one stable isotope-labelled internal standard per compound should be used to account for the ion suppression effects that can arise from matrix effects and changes in mobile phase composition.³⁵ Absolute quantification will immediately indicate whether a lipid is a major or a minor species and allows the direct comparison of lipidomic results across different laboratories, which increases the confidence in lipidomic data.¹²⁷ However, using a lipid standard for each lipid species is almost impossible, particularly in large-scale lipidomic studies where several thousand lipids are present in the biological sample of interest.83 Therefore, instead of one lipid standard for each lipid species, the "minimum standards" established by the Lipidomics Standards Initiative¹²⁸ require that at least one internal standard per class should be used for lipid quantification. However, it is important to note that even then only a relative quantification can be achieved where the increasing and decreasing patterns of lipids between two samples are compared.¹²⁹ While the use of a single internal standard per lipid class may account for errors in extraction efficiency and systematic errors, concentrations of individual species cannot be established.¹²⁹ This is because the instrument's response for different lipid species even within the same lipid class can differ significantly depending on various factors.^{129,130} For example, the instrument's response for phospholipids is dependent on acyl chain length, acyl chain unsaturation, the structure of the polar head group, total lipid concentration, solvent composition, and instrument settings.130 Consequently, multiple internal standards per lipid class distributed across the entire retention time range and acyl chain composition would be required for an accurate quantitative analysis where the concentrations of individual lipids need to be recorded.35,129 Hence, the choice of quantification strategy depends on the scope of the study, and it is necessary to carefully consider all the different strategies before undertaking a lipidomic analysis. The principles of quantification, selection of internal standards for different analytical platforms and the factors that affect the accurate quantification of lipids are extensively reviewed by Wang et al.²

Standardizing lipidomic methodologies

The lipidomics standards initiative (https://lipidomics-standards initiative.org/) was established in 2018 to provide guidelines and standards for lipidomics.¹²⁸ The first draft of guidelines is now provided, which covers the entire lipidomic workflow, including pre-analytics, lipid extraction, data acquisition, lipid identification, quantification, method validation, quality control and data reporting.¹²⁸ If the guidelines introduced by the Lipidomics Standards Initiative¹²⁸ are followed by the lipid research community, this could enable the cross-comparison of lipidomic data across many different laboratories. For example, we recently developed a comprehensive lipid map of *Arabidopsis* across development; this includes a catalogue of the major lipids

observed in different *Arabidopsis* tissues and provides the initial framework for a plant lipid database.¹⁰⁶ This *Arabidopsis* lipid database has great potential to be expanded, for example, by including profiles of sphingolipids, sterol lipids, oxylipins and prenol lipids detected in diverse *Arabidopsis* tissues such as leaves, stems, flowers and roots, during development. Many laboratories around the world conduct studies comparing the lipid profiles of *Arabidopsis* wild-type and mutants under standard and various stress conditions. If the data generated from all these studies can be collated to build a common platform similar to the *Arabidopsis* lipid map, this would enable an effective sharing and comparison of data and results. Following the minimum standards set by the Lipidomics Standards Initiative¹²⁸ would be a first step towards achieving such a resource that assists the lipid research community.

Furthermore, to enable the efficient cross-comparison of lipidomic data, it is advisable to follow the quality control (OC) guidelines set by the lipidomics standards initiative (https://lipidomics-standards-initiative.org/). QC in lipidomic analyses is an important aspect that must be used at all times to ensure data quality. QC samples are used to stabilize the analytical instrument, monitor the instrument performance¹³¹ and account for within and between batch variations in mass spectrometric responses.¹³² The occurrence of systemic variations of instrument's response within and between batches, for example, due to variations in mobile phase composition, system calibration, differences in MS detection sensitivity and resolution, is often encountered in untargeted LC-MS metabolomics.¹³² The qualitative and quantitative tools that can be used to account for batch variations have previously been evaluated and discussed by Sanchez-Illana et al., 2018¹³² while various QC strategies in mass spectrometry-based lipidomics have been reviewed by Xie et al., 2017.131

Identifying lipid functions

Plant scientists have studied lipids for over 50 years and their initial focus was mainly directed at understanding their biosynthesis.¹³³ The development of Arabidopsis thaliana as a genetic plant model, and the sequencing of its genome,¹³⁴ enabled the exploration of many lipid biosynthetic pathways including the synthesis of membrane glycerolipids, sphingolipids, triacylglycerols, cutins, waxes and suberins.^{133,135} With the advent of innovative mass spectrometric technologies in the 1990s,⁷¹ plant biologists started applying mass spectrometry to study lipid metabolic pathways involved in plant development and stress responses,^{71,136} thereby identifying lipids that act as substrates and products of enzymes137 and detecting lipidmetabolizing enzymes.²¹ To gain a clear understanding of the in vivo functions of lipids in plants, often several different approaches are required including genetic, genomic¹⁹ and biochemical studies. These studies can provide novel insights into the roles of lipids in plants and the information may help in designing engineering strategies in biotechnology.¹⁹ For example, TGs have been long known to be the major energy

reserves in seeds;^{19,138} however, newer studies have shown that they are also involved in cell division and expansion,¹³⁹ stomatal opening,¹⁴⁰ membrane lipid remodelling,¹⁹ organ formation,¹⁹ stress response¹⁴¹ and pollination.¹⁴² This suggests that strategies aiming to increase the amount of TGs in plant tissues for biodiesel and industrial chemical production must be designed carefully,¹⁹ as an overaccumulation of TGs could prove detrimental to the growth of such engineered plants.¹⁹

It has been estimated that the cellular lipidome comprises hundreds of thousands of diverse lipids^{1,143} and the plant cellular sphingolipidome alone may comprise at least 500 and perhaps thousands of different species of sphingolipids.¹⁴⁴ Theoretically, possible lipid structures exceed 180000 without even considering all the position isomers, backbone substitutions and stereochemistry. Indeed several million potential lipids could exist when all structural differences are taken into account.98 This suggests that much of the plant lipidome is yet to be fully described and that there are likely many lipid species to be discovered. However, new species are most often present in low levels in complex biological matrices compared to other more common lipids,¹⁰⁷ which is one of the main reasons why they are yet to be discovered. Due to their low abundance, the precursor ions will not be selected for fragmentation by datadependent acquisition methods in the LC-MS/MS approaches, 107 which makes it difficult to identify them. In shotgun lipidomics, the MS/MS spectra of novel species are difficult to interpret due to the presence of product ions of co-fragmented lipids and chemical noise.¹⁰⁷ Furthermore, identifying the structures of all unknown lipids in a complex biological sample is challenging since thousands of unidentified mass spectrometric features are detected,⁹² some of which may be artifacts generated from insource fragmentation¹⁴⁵ while some peaks could also be the result of solvent masses.¹⁴⁶ Multiple adduct formation by the same lipid species and isotopic peaks may further complicate the identification of novel lipids.¹⁴⁶ Together these factors hinder the systematic discovery of novel lipids.

A strategy that can be used to detect novel lipids is to look for new peaks in the mass spectra of the treated samples, which are absent in control samples. Since plant cells often produce lipids in response to environmental stresses and during the developmental process,¹⁴⁷ there is potential to discover new lipid species through a comparison of lipid profiles from treated and control plant tissues. Given that the respective biosynthetic genes are identified first, the functions of newly discovered lipids can be studied in the corresponding mutants. For example, a new class of lipids, glucuronosyldiacylglycerols, was discovered in phosphorus-starved Arabidopsis plants, and further studies showed that they are essential for protection against phosphorus depletion.¹⁴⁸ The authors observed a new peak in the mass spectra of lipid extracts from the leaves of phosphorus-starved wild-type plants but not in the extracts of the plants grown with sufficient phosphorus.¹⁴⁸ Further investigations into the new peak by tandem mass spectrometry revealed the building blocks to be a glucuronosylglycerol and two fatty acids, which suggested that the structure is a glucuronosylated diacylglycerol.148 The novel compound was purified

from Arabidopsis leaf material and its structure was established by nuclear magnetic resonance spectroscopy (NMR).¹⁴⁸ Subsequent studies revealed that the glucuronosyldiacylglycerols mitigate phosphorus depletion stress as sqd2 mutant plants, which were unable to accumulate SQDGs and glucuronosyldiacylglycerol, showed severe growth defects under phosphorus limiting conditions.¹⁴⁸ In another study, mechanical disruption of leaf tissue caused by wounding, pathogen attack and cold stress stimulated the production of an extensive variety of oxidized and acylated galactolipids in Arabidopsis.149 The enzyme responsible for the acylation of the galactolipid head group was purified from oat leaves and subsequently identified using tandem MS. This led to the identification of the Arabidopsis ortholog, which was subsequently confirmed through lipid analysis of loss-offunction mutants.¹⁴⁹ These examples highlight that by monitoring the lipid changes of plants subjected to stress conditions and by comparing the lipid profiles of loss-of-function mutants and wild-type plants, novel lipids essential for the survival of plants under challenging environmental conditions can be detected and their specific functions can be uncovered.

Conclusions

LC-MS based technology is now considered as a powerful tool to explore lipid biosynthetic pathways, identify lipid-metabolizing enzymes, and uncover lipid functions.

Plant lipid extraction methods, which were initially adapted from animal lipid extraction methods, have also vastly improved and modified to better suit the requirements for extracting lipids from plant tissue.

Shotgun lipidomics and LC-MS based lipidomics are the two lipidomic approaches widely used to effectively analyse plant lipids. For LC-MS based plant lipidomic workflows, there is a shift towards using high-resolution mass spectrometers such as QToF and orbitrap coupled with RPLC and HILIC. Most highresolution mass spectrometers now offer high sensitivity with higher resolution that can be used for both targeted and discovery lipidomic approaches. Nevertheless, the unambiguous identification of lipids is still extremely challenging and requires the manual curation of data sets, and yet, no lipid library specifically built for plant lipid identification is available.

The development of high-resolution mass spectrometers and comprehensive data acquisition methods enable the detection of not only known lipid species but also a plethora of unidentified features. It is currently impossible to identify all unknown mass spectrometric features that can be detected and many of them may be due to artefacts. However, recent studies have shown that by considering lipids as part of an integrated system with enzymes and metabolites and in the context of plant function, new lipids essential for the survival of plants can be identified and studied.

Conflicts of interest

There are no conflicts of interest to declare.

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