

Shorthand notation for lipid structures derived from mass spectrometry

Gerhard Liebisch,^{1,*} Juan Antonio Vizcaíno,[†] Harald Köfeler,^{**} Martin Trötz Müller,^{**} William J. Griffiths,^{††} Gerd Schmitz,^{*} Friedrich Spener,^{§,**} and Michael J. O. Wakelam^{***}

Institute of Clinical Chemistry and Laboratory Medicine,^{*} University of Regensburg, Regensburg, Germany; EMBL-European Bioinformatics Institute,[†] Hinxton, Cambridge, United Kingdom; Institute of Molecular Biology and Biochemistry,[§] and Core Facility Mass Spectrometry,^{**} Medical University of Graz, Graz, Austria; Institute of Mass Spectrometry,^{††} College of Medicine, Swansea University, Singleton Park, Swansea, United Kingdom; and Babraham Institute,^{***} Babraham Research Campus, Cambridge, United Kingdom

Abstract There is a need for a standardized, practical annotation for structures of lipid species derived from mass spectrometric approaches; i.e., for high-throughput data obtained from instruments operating in either high- or low-resolution modes. This proposal is based on common, officially accepted terms and builds upon the LIPID MAPS terminology. It aims to add defined levels of information below the LIPID MAPS nomenclature, as detailed chemical structures, including stereochemistry, are usually not automatically provided by mass spectrometric analysis. To this end, rules for lipid species annotation were developed that reflect the structural information derived from the analysis. For example, commonly used head group-specific analysis of glycerophospholipids (GP) by low-resolution instruments is neither capable of differentiating the fatty acids linked to the glycerol backbone nor able to define their bond type (ester, alkyl-, or alk-1-enyl-ether). This and other missing structural information is covered by the proposed shorthand notation presented here. Beyond GPs, we provide shorthand notation for fatty acids/acyls (FA), glycerolipids (GL), sphingolipids (SP), and sterols (ST). **In summary, this defined shorthand nomenclature provides a standard methodology for reporting lipid species from mass spectrometric analysis and for constructing databases.**—Liebisch, G., J. A. Vizcaíno, H. Köfeler, M. Trötz Müller, W. J. Griffiths, G. Schmitz, F. Spener, and M. J. O. Wakelam. **Shorthand notation for lipid structures derived from mass spectrometry.** *J. Lipid Res.* 2013. 54: 1523–1530.

Supplementary key words lipidomics • nomenclature • abbreviation • fatty acids • glycerolipids • glycerophospholipids • sphingolipids • sterols

A comprehensive classification system for lipids was presented by the International Lipid Classification and

Nomenclature Committee (ILCNC) in 2005 (1) and updated in 2009 (2). This system places lipids into eight categories and is available online on the LIPID MAPS website (<http://www.lipidmaps.org>). The LIPID MAPS nomenclature precisely describes lipid structures.

The key technology for lipid species analysis is mass spectrometry (MS) (3, 4). Typically, MS analysis without intermediate chemical steps does not provide the structural details covered by the LIPID MAPS nomenclature, which led mass spectrometrists to use a variety of different notations for lipid species. Moreover, lipid species are frequently annotated based on assumptions. For example, a precursor ion scan of m/z 184, a standard approach to detect phosphatidylcholine (PC) and sphingomyelin (SM) species, is neither able to differentiate PC species containing an ether bond from diacyl species (Fig. 1A), nor is it able to differentiate the sphingoid base in SM (5). Similarly, annotation of phosphatidylethanolamine (PE) species, particularly plasmalogens, should not be based on head group-specific positive neutral loss (NL 141) or negative precursor ion (PIS m/z 196) scans (6), which identify only the lipid class but do not provide specific mass spectrometric analysis (7) (Fig. 1B).

Although lipidologists possess a “biological intelligence,” which allows them to interpret MS data in a manner that recognizes what is likely or not likely to be the correct structure of a particular lipid species, we think there is need for a standardized, practical shorthand notation of lipid structures derived from MS approaches that enables correct and concise reporting of data and their deposition in databases. Our proposal is based on common, officially accepted terms and on the LIPID MAPS terminology. In addition, it takes

Abbreviations: GL, glycerolipid; GP, glycerophospholipid; ILCNC, International Lipid Classification and Nomenclature Committee; LIMS, laboratory information management system; NL, neutral loss; PC, phosphatidylcholine; PE, phosphatidylethanolamine; PIS, precursor ion scan; SP, sphingolipid; ST, sterol.

¹To whom correspondence should be addressed.
e-mail: gerhard.liebisch@klinik.uni-regensburg.de

This work was supported by the LipidomicNet project, funded within the Seventh Framework Programme of the European Union (Grant 202272) and by the Biotechnology and Biological Sciences Research Council (W.J.G. and M.J.O.W.).

Manuscript received 22 October 2012 and in revised form 2 April 2013.

Published, JLR Papers in Press, April 2, 2013

DOI 10.1194/jlr.M033506

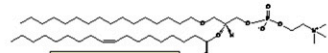
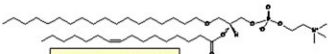
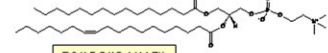
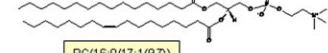
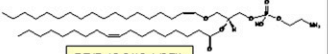
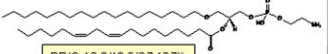
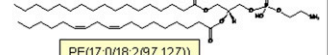
A	Lipid Structure (LIPID MAPS Nomenclature)	Lipid class level mass PIS <i>m/z</i> 184	Lipid species level PIS <i>m/z</i> 184	Bond type level High Resolution	Fatty acyl/alkyl level FA Scans	Fatty acyl/alkyl position level Position Analysis
	 <p>PC(O-16:0/18:1(9Z)) M = 745.60 g/mol</p>	PC (745)	PC 33:1 * PC O-34:1 **	PC O-34:1	PC O-16:0_18:1 ***	PC O-16:0/18:1 ****
	 <p>PC(O-18:0/16:1(9Z)) M = 745.60 g/mol</p>	PC (745)	PC 33:1 * PC O-34:1 **	PC O-34:1	PC O-18:0_16:1 ***	PC O-18:0/16:1 ****
	 <p>PC(15:0/18:1(11Z)) M = 745.56 g/mol</p>	PC (745)	PC 33:1 * PC O-34:1 **	PC 33:1	PC 15:0_18:1	PC 15:0/18:1
	 <p>PC(16:0/17:1(9Z)) M = 745.56 g/mol</p>	PC (745)	PC 33:1 * PC O-34:1 **	PC 33:1	PC 16:0_17:1	PC 16:0/17:1
B	Lipid Structure (LIPID MAPS Nomenclature)	Lipid class level mass NL+ 141 PIS- 196	Lipid species level NL+ 141 PIS- 196	Bond type level High resolution	Fatty acyl/alkyl level FA scans	Fatty acyl/alkyl position level Position analysis
	 <p>PE(P-18:0/18:1(9Z)) M = 729.57 g/mol</p>	PE (729)	PE 35:2 * PE O-36:2 **	PE O-36:2	PE O-18:1_18:1	PE P-18:0/18:1 ****
	 <p>PE(O-18:0/18:2(9Z,12Z)) M = 729.57 g/mol</p>	PE (729)	PE 35:2 * PE O-36:2 **	PE O-36:2	PE O-18:0_18:2 ***	PE O-18:0/18:2
	 <p>PE(17:0/18:2(9Z,12Z)) M = 729.53 g/mol</p>	PE (729)	PE 35:2 * PE O-36:2 **	PE 35:2	PE 17:0_18:2	PE 17:0/18:2

Fig. 1. Ambiguities in interpretation of MS data. Examples for annotation of (A) phosphatidylcholine (PC) species and (B) phosphatidylethanolamine (PE) species and typical MS approaches to identify lipid species. *The annotation is based on the assumption that ester bonds are present. **The annotation is based on the assumption of even numbered carbon chains only. ***Unambiguous identification of an alkyl bond is only possible in the case of a saturated alkyl chain. ****Unambiguous determination of an alk-1-enyl bond requires specific MS experiments; e.g., according to Zemski, Berry, and Murphy (7).

the different levels of structural information provided by MS into account (Fig. 2). At the lowest level of resolution, the respective LIPID MAPS abbreviation (Table 1) is followed either by the detected nominal mass (lipid class level mass) or by the sum of components that are expressed as their total number of carbon atoms and of double bonds (lipid species level); variable components of the species are not identified. In the presence of fatty acids with odd-numbered carbon atoms, ambiguities regarding functional groups or bond types may occur. Therefore, such species are either assigned by their molecular mass or based on assumptions which should be presented with the result.

Bond type level additionally describes the type of linkage of the variable components to the lipid species' backbone without knowing the single components. When MS resolves the variable components of the lipid species (in most cases fatty acids) the fatty acyl/alkyl level notation is applicable. Finally, when a specific analysis for backbone position (*sn*- for

stereospecific numbering) in glycerolipids and glycerophospholipids categories has been carried out, the fatty acyl/alkyl position level is applicable. Fatty acyl/alkyl/sphingoid base structure level describes structural details of these components. Full structural analysis of the lipid species is covered by the LIPID MAPS nomenclature. The proposal presented here covers the major lipid classes of five of the eight LIPID MAPS categories, namely, fatty acyls (FA), glycerolipids (GL), glycerophospholipids (GP), sphingolipids (SP), and sterols (ST), with a focus on mammalian lipids. Other minor lipid classes or lipids from other organisms could be the subject of further proposals.

GENERAL RULES FOR SHORTHAND NOTATION

All presentations of lipid species data include an a priori statement on structural resolution attained by the method of MS analysis. It should be a requirement that lipids are

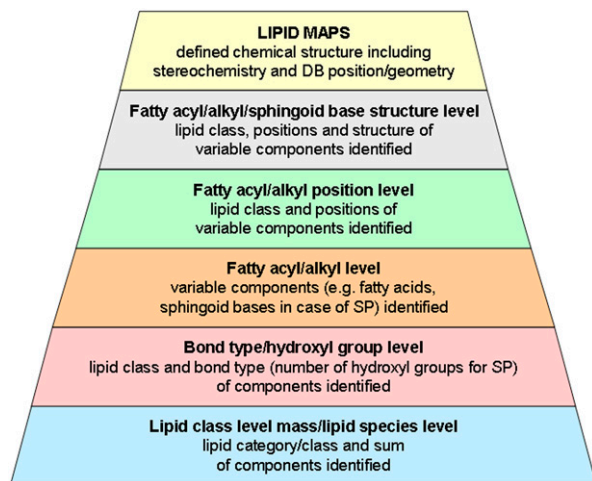


Fig. 2. MS shorthand notation for lipid species aims to add defined levels of information when insufficient data is available to employ LIPID MAPS nomenclature. These levels should cover a correct and concise presentation of the structural information provided by MS analysis.

defined by both their class and their nominal mass (Da). The following rules apply to all lipid categories described below:

- Lipid class abbreviation heads each species description.
- Variable components (constituents), such as fatty acids, are assigned based on their mass as number of C-atoms and number of double bonds (C-atoms:double bonds).
- Only experimentally proven structural details of constituent fatty acids are assigned according to the rules defined for fatty acyls (see below).
- When structural ambiguities are present (e.g., bond type, hydroxyl groups, branched chains, see examples in Fig. 1 and **Tables 2–7**), species are assigned by one of the following rules:
 - Lipid class and the (uncharged) molecular mass (Da) in parentheses (preferred for reporting in databases $\hat{=}$ lipid class level mass). For fatty acyl substituents, the mass of the corresponding free fatty acid is used (see example in Table 6).
 - Annotation based on assumptions must be clearly visible (preferred for publications $\hat{=}$ lipid species level).
- Detailed structures including stereochemistry are covered by LIPID MAPS nomenclature.

FATTY ACYLS (FA)

Fatty acids in free form and as variable fatty acyls in lipids are prevalent lipid structures. Therefore, we use fatty acids and acyls as a paradigm for application of the shorthand notation. For the sake of simplicity, we include some frequent functional groups of fatty acids but do not treat complex FA classes, such as eicosanoids and docosanoids (8). When an annotation of the fatty acid is only based on the mass (low mass resolution), usually it is assumed that a straight-chain fatty acid with no functional groups is present beyond double bond(s). High mass resolution with accurate

mass may identify functional groups. The following rules apply; examples are given in Table 2:

- Shorthand notation: FA number of C-atoms:number of double bonds.
- Functional groups, whose positions in the acyl chain are not known, are shown after the number of double bonds separated by an underscore and followed by the number of groups if more than one.
- Proven positions of functional groups are shown after the number of double bonds (each type of functional group inside a separate pair of parentheses). Positions according to Δ -nomenclature are stated in front of the functional groups that are separated by a comma if more than one.
- Double bond position is indicated by a number according to Δ -nomenclature (geometry unknown) or a number followed by geometry (Z for *cis*, E for *trans*).
- Abbreviations for functional groups:
 - OH for hydroxyl group
 - O for keto group (it is important to note that “O” before the number of carbons refers to an ether bond; see next section)
 - Me for methyl branch
- Order of functional groups: Double bonds - OH - O - Me.

GLYCEROLIPIDS (GL) AND GLYCEROPHOSPHOLIPIDS (GP)

Lipidomic approaches frequently apply direct infusion tandem MS using a low mass resolution analyzer, such as a triple quadrupole mass spectrometer (4, 9, 10). In this way lipid classes and species are identified by selective precursor and neutral-loss scans (6). A major problem of lipid class-specific scans is that bond type, i.e., ester- or ether-linkage to glycerol backbone of constituent acyl-, alkyl-, and alk-1-enyl-chains, cannot be differentiated because these species are quasi-isobaric (Fig. 1).

Frequently, such data are annotated based on the assumption that ester bonds are present. To demonstrate this possibility of incorrect assignment, Fig. 1 includes definition of the lipid species level for glycerophospholipids; this could equally apply to glycerolipids and fatty acyls. An approach to resolve ester and ether bonds is the application of high-resolution MS (11). However, even in high-resolution MS, unsaturated O-alkyl groups cannot be differentiated from O-alk-1-enyl linked residues (Fig. 1B). Yet, specific MS methods exist to clearly identify O-alk-1-enyl linked residues having no or further double bonds in the alkyl-chain (plasmalogens) (7).

The following rules apply for shorthand notation of both the major classes of GL and GP categories; examples are given in Tables 3 and 4:

- Shorthand notation: lipid class abbreviation followed by number of C-atoms:number of double bonds.
- Fatty acids linked to the glycerol are known:
 - Separator $_$: *sn*-position of the fatty acids is not known.
 - Separator $/$: *sn*-position of fatty acids is proven (order *sn*-1/*sn*-2/*sn*-3 for GL; *sn*-1/*sn*-2 or *sn*-2/*sn*-3 for GP); no FA linked 0:0.

TABLE 1. Lipid class abbreviations

LIPID MAPS Category/Class - Common Name	Lipid Class - LIPID MAPS	Abbreviation
Fatty acyls [FA]		
Fatty acids	Fatty acids and conjugates [FA01]	FA
Glycerolipids [GL]		
Monoglycerides	Monoradylglycerols [GL01]	MG
Diglycerides	Diradylglycerols [GL02]	DG
Triglycerides	Triradylglycerols [GL03]	TG
Glycerophospholipids [GP]		
Bis[monoacylglycero]phosphates	Monoacylglycerophosphomonoradylglycerols [GP0410]	BMP
Cardiolipins	Glycerophosphoglycerophosphoglycerols [GP12]	CL
Phosphatidic acids	Glycerophosphates [GP10]	PA
Phosphatidylcholines	Glycerophosphocholines [GP01]	PC
Phosphatidylethanolamines	Glycerophosphoethanolamines [GP02]	PE
Phosphatidylglycerols	Glycerophosphoglycerols [GP04]	PG
Phosphatidylglycerolphosphate	Glycerophosphoglycerophosphates [GP05]	PGP
Phosphatidylinositols	Glycerophosphoinositols [GP06]	PI
Phosphatidylinositol-monophosphate	Glycerophosphoinositol monophosphates [GP07]	PIP
Phosphatidylinositol-3-phosphate	Glycerophosphoinositol monophosphates [GP07]	PIP[3']
Phosphatidylinositol-4-phosphate	Glycerophosphoinositol monophosphates [GP07]	PIP[4']
Phosphatidylinositol-5-phosphate	Glycerophosphoinositol monophosphates [GP07]	PIP[5']
Phosphatidylinositol-bisphosphate	Glycerophosphoinositol bisphosphates [GP08]	PIP2
Phosphatidylinositol-3,4-bisphosphate	Glycerophosphoinositol bisphosphates [GP08]	PIP2[3',4']
Phosphatidylinositol-3,5-bisphosphate	Glycerophosphoinositol bisphosphates [GP08]	PIP2[3',5']
Phosphatidylinositol-4,5-bisphosphate	Glycerophosphoinositol bisphosphates [GP08]	PIP2[4',5']
Phosphatidylinositol-trisphosphate	Glycerophosphoinositol trisphosphates [GP09]	PIP3
Phosphatidylserines	Glycerophosphoserines [GP03]	PS
Lysophospholipids		Prefix L
Sphingolipids [SP]		
Ceramides	Ceramides [SP02]	Cer
Ceramide-1-phosphates	Ceramide-1-phosphates [SP0205]	C1P
Sphingoid bases	Sphingoid bases [SP01]	SPH
Sphingoid base-1-phosphates	Sphingoid bases [SP01]	S1P
Sphingomyelins	Phosphosphingolipids [SP03]	SM
Hexosylceramides	Neutral glycosphingolipids [SP05]	HexCer
Glucosylceramide	Neutral glycosphingolipids [SP05]	GlcCer
Galactosylceramide	Neutral glycosphingolipids [SP05]	GalCer
Dihexosylceramides	Neutral glycosphingolipids [SP05]	Hex2Cer
Lactosylceramide	Neutral glycosphingolipids [SP05]	LacCer
Sterol lipids [ST]		
Sterols	Sterols [ST01]	ST
Steryl esters	Steryl esters [ST0102]	SE
Free cholesterol	Cholesterol [LMST01010001]	FC
Cholesteryl ester	Cholesteryl esters [ST0102]	CE
Bile acids	Bile acids and derivatives [ST04]	
	Cholic acid	CA
	Chenodeoxycholic acid	CDCA
	Deoxycholic acid	DCA
	Ursodeoxycholic acid	UDCA
	Hyodeoxycholic acid	HDCA
	Lithocholic acid	LCA
	Glycocholic acid	GCA
	Taurocholic acid	TCA

Abbreviations are in agreement with Table 3 in Fahy et al. (1) and the updated LIPID MAPS nomenclature (2).

• Bond types other than ester bonds are indicated as follows in front of the sum of C-atoms or fatty acid:

○ O = proven O-alkyl-bond (it is important to note that “O” after the number of carbons designates a keto bond; see previous section)

○ P = proven O-alk-1-enyl-bond (acid-sensitive ether bond in “plasmalogens”).

• More than one “non”-ester bond is indicated in front of the bond type as *d* for *di*, *t* for *tri*.

Additional rules for glycerophospholipids (GP):

• Lysophospholipid classes are abbreviated as stated in the LIPID MAPS nomenclature (Table 1). Where applicable they can be presented formally by their respective phospholipid class indicating the empty *sn*-position by 0:0.

• Four ether bonds are indicated in front of the bond type as *tetra*, abbreviated *e*.

• For BMP and CL classes, the *sn*-position order will be *sn*-2/*sn*-3/*sn*-2'/*sn*-3' and *sn*-1/*sn*-2/*sn*-1'/*sn*-2', respectively.

• Rules for phosphoinositides require additional information if the position of phosphates on the inositol ring are known (see Table 4). The exception is PI3,4,5P₃, as the only known tris isomer is 3,4,5. While phosphoinositides are generally presented as PIP₂ and PIP₃ in Tables 1 and 4, we adopted the annotation PIP2 and PIP3 for ease of handling by databases. Should additional species be identified, this will require further clarification.

TABLE 2. Fatty acids and acyls: shorthand notation examples

Chain Type/Functional Group	Lipid Class Level Mass ^a	Lipid Species Level ^b	Fatty Acyl Level	LIPID MAPS Fatty Acyl Structure Level
Straight chain	FA (304)	FA 20:4	FA 20:4	FA 20:4(4Z,8Z,11Z,4Z)
	FA (282)	FA 18:1	FA 18:1	FA 18:1(9E) FA 18:1(9) ^c
Methyl branched	FA (312)	FA 20:0	FA 16:0_Me4	FA 16:0(3Me,7Me,11Me,15Me)
Hydroxy	FA (300)		FA 18:0_OH	FA 18:0(9OH)
Keto	FA (200)	FA 12:0	FA 11:0_O	FA 11:0(3O)

^aUncharged molecular mass.^bAnnotation based on the assumption of a straight-chain fatty acid with no functional groups except double bond(s).^cUnknown geometry.

SPHINGOLIPIDS (SP)

Several MS methods for sphingolipid species analysis use fragments resulting from the sphingoid base (12), but the commonly used precursor ion scan of m/z 184 for SM analysis is not able to differentiate between the N-linked fatty acid and sphingoid base (13), although there is a more time-consuming procedure that can provide this information (12). In this case, lipid species level annotation could be based on the assumption of the major sphingoid backbone in the respective organism; e.g., sphingosine (d18:1) in mammals or phytosphingosine (t18:0) in yeast. This assumption must be indicated a priori. High-resolution MS allows identification of the number of hydroxyl groups in sphingolipids together with the sum of carbons and double bonds in the sphingoid base and N-linked fatty acid (14).

The following rules apply; examples are given in Tables 5 and 6:

- The sphingoid backbone is annotated by the number of hydroxyl groups in the sphingoid base (m for *mono*, d for *di*, t for *tri*) and separated by a slash from the number of carbons:number of double bonds of the N-linked fatty acid. Positions of hydroxyl groups and double bonds including geometry are indicated as described for fatty acyls (FA).

- If the sphingoid base is not known, the sum of sphingoid base and fatty acid is shown as number of carbons:number of double bonds. Calculations are based on the number of hydroxyl groups of the major sphingoid base for that organism (dihydroxy in mammals). When the number

of hydroxyl groups is known, it is shown in front of the number of carbons (hydroxyl group level, e for *tetra*).

- For further characterization of N-linked fatty acids, rules as described in an earlier section apply. A fatty acid that is ester-bound to an N-linked ω -OH fatty acid is shown in square brackets as [ω FA C-atoms:double bonds].

- Shorthand notation for sugar moieties is stated in Table 1.

This proposal, however, does not cover complex glycosphingolipids, which we suggest could be subject to a separate proposal.

STEROLS (ST)

We use the term sterol to embrace all molecules based on the cyclopentanoperhydrophenanthrene skeleton. All natural mammalian sterols are derived from cholesterol or its precursors, although plant sterols can also be a source. The stereochemistry of the cholesterol molecule is maintained to a large extent by mammalian sterols, which all contain at least one alcohol or oxo group attached to carbon 3. Thus, at the lipid species level, we assume the sterol has at least one alcohol group. High-resolution MS with accurate mass may identify other functional groups, as will precursor ion and neutral loss scans. Stereochemistry can often be defined by comparing the chromatographic retention time to authentic standards and, in some cases, by MS/MS. The following rules for shorthand nomenclature have been adopted in the examples given in Table 7:

- Shorthand notation: ST number of carbon atoms:number of double bonds.

TABLE 3. Glycerolipids: shorthand notation examples

Bond Type	Lipid Class Level Mass ^a	Lipid Species Level ^b	Bond Type Level	Fatty Acyl/Alkyl Level	Fatty Acyl/Alkyl Position Level	LIPID MAPS Fatty Acyl/Alkyl Structure Level
Acyl	MG (358)	MG 18:0	MG 18:0	MG 18:0	MG 0:0/18:0/0:0	
Alkyl	MG (344)	MG O-18:0	MG O-18:0	MG O-18:0	MG 0:0/O-18:0/0:0	
Diacyl	DG (594)	DG 34:1	DG 34:1	DG 16:0_18:1	DG 16:0/18:1/0:0	DG 16:0/18:1(9Z)/0:0
Alkyl-acyl	DG (580)	DG O-34:1	DG O-34:1	DG O-16:0_18:1	DG O-16:0/18:1/0:0	DG O-16:0/18:1(9Z)/0:0
Dialkyl	DG (538)	DG 30:1	DG dO-32:1	DG O-16:0_O-16:1	DG O-16:0/O-16:1/0:0	DG O-16:0/O-16:1(9Z)/0:0
Triacyl	TG (858)	TG 52:2	TG 52:2	TG 16:0_18:1_18:1	TG 16:0/18:1/18:1	TG 16:0/18:1(9Z)/18:1(11Z)
Alkyl-diacyl	TG (844)	TG O-52:2	TG O-52:2	TG O-16:0_18:1_18:1	TG O-16:0/18:1/18:1	TG O-16:0/18:1(9Z)/18:1(11Z)
Dialkyl-acyl	TG (830)	TG 50:2	TG dO-52:2	TG O-18:1_O-16:0_18:1	TG O-18:1/O-16:0/18:1	TG O-18:1(9Z)/O-16:0/18:1(9Z)
Trialkyl	TG (816)	TG O-50:2	TG tO-52:2	TG O-18:1_O-16:0_O-18:1	TG O-18:1/O-16:0/O-18:1	TG O-18:1(9Z)/O-16:0/O-18:1(9Z)

^aUncharged molecular mass.^bAnnotation based on assumption of even-numbered carbon chains only.

TABLE 4. Glycerophospholipids: shorthand notation examples

Bond Type	Lipid Class Level Mass ^a	Lipid Species Level ^b	Bond Type Level	Fatty Acyl/Alkyl Level	Fatty Acyl/Alkyl Position Level	LIPID MAPS Fatty Acyl/Alkyl Structure Level
Phospho- and lysophospholipids containing ester and/or ether bonds						
Diacyl	BMP (690)	BMP 34:2	BMP 34:2	BMP 16:0_18:2	BMP 16:0/0:0/18:2/0:0 <i>sn-2/sn-3/sn-2'/sn-3'</i>	BMP 16:0/0:0/18:2(9Z,12Z)/0:0 <i>sn-2/sn-3/sn-2'/sn-3'</i>
Tetraacyl	CL (1450)	CL 72:7	CL 72:7	CL 18:1_18:2_18:2_18:2 CL 36:3_36:4 (known DG fragments)	CL 18:1/18:2/18:2/18:2 <i>sn-1/sn-2/sn-1'/sn-2'</i>	CL 18:1(9Z)/18:2(9Z,12Z)/ 18:2(9Z,12Z)/18:2(9Z,12Z) <i>sn-1/sn-2/sn-1'/sn-2'</i>
Tetraalkyl	CL (1521)	CL 76:0	CL eO-80:0	CL O-20:0/O-20:0/ O-20:0/O-20:0	CL O-20:0/O-20:0/ O-20:0/O-20:0	CL O-16:0(3Me,7Me,11Me,15Me)/ O-16:0(3Me,7Me,11Me,15Me)/ O-16:0(3Me,7Me,11Me,15Me)/ O-16:0(3Me,7Me,11Me,15Me)
Diacyl	PC (759)	PC 34:1	PC 34:1	PC 16:0_18:1	PC 16:0/18:1	PC 16:0/18:1(9Z)
Alkyl-acyl	PC (745)	PC O-34:1	PC O-34:1	PC O-16:0_18:1	PC O-16:0/18:1	PC O-16:0/18:1(9Z)
Dialkyl	PC (731)	PC 32:1	PC dO-34:1	PC O-16:0_O-18:1	PC O-16:0/O-18:1	PC O-16:0/O-18:1
Diacyl	PE (717)	PE 34:1	PE 34:1	PE 16:0_18:1	PE 16:0/18:1	PE 16:0/18:1(9Z)
Plasmalogen	PE (701)	PE O-34:2	PE O-34:2	PE P-16:0_18:1 ^c	PE P-16:0/18:1 ^c	PE P-16:0/18:1(9Z)
Triacyl	LCL (1188)	LCL 54:5	LCL 54:5	LCL 18:1_18:2_18:2	LCL 18:1/18:2/18:2/0:0	LCL 18:1(9Z)/18:2(9Z,12Z)/ 18:2(9Z,12Z)/0:0
Acyl	LPC (495)	LPC 16:0	LPC 16:0	LPC 16:0 or PC 16:0_0:0	LPC 16:0/0:0 or PC 16:0/0:0	LPC 16:0/0:0 or PC 16:0/0:0
Alkyl	LPC (481)	LPC O-16:0	LPC O-16:0	LPC O-16:0	LPC O-16:0/0:0	LPC O-16:0/0:0
Alkyl/Alk-1-enyl	LPC (479)	LPC O-16:1	LPC O-16:1	LPC O-16:1	LPC O-16:1/0:0	LPC O-16:1(9Z)/0:0
Additional complexity introduced by inositol ring phosphate position						
Diacyl	PIP (966)	PIP 38:4	PIP 38:4	PIP[3'] 38:4	PIP[3'] 18:0/20:4	PIP[3'] 18:0/20:4(4Z,8Z,11Z, 4Z)
Diacyl	PIP2 (1046)	PIP2 38:4	PIP2 38:4	PIP2[4',5'] 38:4	PIP2[4',5'] 18:0/20:4	PIP2[4',5'] 18:0/20:4 (4Z,8Z,11Z, 4Z)

^aUncharged molecular mass.^bAnnotation based on assumption of even-numbered carbon chains only.^cIdentification of plasmalogens (alk-1-enyl bond) requires specific MS analysis (7).

- Annotation at the lipid species level is based on natural sterols possessing 18, 19, 21, 24, or 27 carbons and at least one hydroxyl group at position 3.

- Functional groups, including all hydroxyl groups, are shown after the number of double bonds, separated by an underscore and followed by the number of groups if more than one.

- Proven positions of functional groups are shown after the number of double bonds, separated by a slash. Specific stereochemistry of functional groups is shown in square brackets. (*R*) and (*S*) configurations are preferred for side-chain stereochemistry and are given in italics in parentheses.

- In the case of fully proven structures of cholesterol and cholesteryl esters, abbreviations FC and CE, respectively,

can be used (Table 1). CE is followed by number of carbons:number of double bonds of the fatty acid esterified to the hydroxyl group at position 3 (Table 7).

- In the case of unproven structures and other sterol esters (SE), the shorthand notation is used as above, followed by slash number of carbons:number of double bonds of the fatty acid esterified to the hydroxyl group (Table 7).

- In the case of bile acids, the shorthand notation is prefaced by A to indicate an acid, and at the structure level, the location of the acid group (CO₂H) is indicated.

- Precursor-ion scans reveal the presence of conjugating groups, i.e., taurine (T) or glycine (G), conjugated to the carboxylic acid group of bile acids through an amide bond, sulfuric acid (S) conjugated to a hydroxyl group through

TABLE 5. Sphingolipids with a free amino group: shorthand notation examples

Sphingoid Base	Lipid Class Level Mass ^a	Hydroxyl Group Level	LIPID MAPS Sphingoid Base Structure Level
Sphingosine	SPH (299)	SPH d18:1	SPH d18:1(4E)(1OH,3OH)
Sphinganine	SPH (301)	SPH d18:0	SPH d18:0(1OH,3OH)
Sphingadiene	SPH (297)	SPH d18:2	SPH d18:2(4E,8E)(1OH,3OH)
Phytosphingosine	SPH (317)	SPH t18:0	SPH t18:0(1OH,3OH,4OH)
Sphingosine-C20	SPH (327)	SPH d20:1	SPH d20:1(4E)(1OH,3OH)
Sphingosine	S1P (379)	S1P d18:1	S1P d18:1(4E)(1OH,3OH)
Sphinganine	S1P (381)	S1P d18:0	S1P d18:0(1OH,3OH)
1-Deoxymethyl-sphinganine	SPH (271)	SPH m17:0	SPH m17:0(2OH)
1-Deoxy-sphinganine	SPH (285)	SPH m18:0	SPH m18:0(3OH)

^aUncharged molecular mass.

TABLE 6. Sphingolipids containing an amide bound fatty acid: shorthand notation examples

Lipid Class Level Mass ^a	Lipid Species Level ^b	Hydroxyl Group Level	Fatty Acyl Level	LIPID MAPS Sphingoid Base/Fatty Acyl Structure Level
Cer (537)	Cer 34:1	Cer d34:1	Cer d18:1/16:0	Cer d18:1(4E)(1OH,3OH)/16:0
Cer (539)	Cer 34:0	Cer d34:0	Cer d18:0/16:0	Cer d18:0(1OH,3OH)/16:0
C1P (617)	C1P 34:1	C1P d34:1	C1P d18:1/16:0	C1P d18:1(4E)(1OH,3OH)/16:0
SM (702)	SM 34:1	SM d34:1	SM d18:1/16:0	SM d18:1(4E)(1OH,3OH)/16:0
SM (704)	SM 34:0	SM d34:0	SM d18:0/16:0	SM d18:0(1OH,3OH)/16:0
SM (840)	SM 44:2	SM d44:2	SM d20:1/24:1	SM d20:1(4E)(1OH,3OH)/24:1(15Z)
HexCer (701)	HexCer 34:1	HexCer d34:1	HexCer d18:1/16:0	GlcCer d18:1(4E)(1OH,3OH)/16:0
HexCer (701)	HexCer 34:1	HexCer d34:1	HexCer d18:1/16:0	GalCer d18:1(4E)(1OH,3OH)/16:0
Hex2Cer (861)	Hex2Cer 34:1	Hex2Cer d34:1	Hex2Cer d18:1/16:0	LacCer d18:1(4E)(1OH,3OH)/16:0

^aUncharged molecular mass.^bAnnotation based on assumption of a sphingoid base with two hydroxyl groups.

an ester bond, glucuronic acid (GlcA), *N*-acetylglucosamine (GlcNAc), and hexose (Hex) sugars assumed to be linked to a hydroxyl group through an acetal linkage.

• In the case full stereochemistry is known, the abbreviations given in Table 1 can be used.

DISCUSSION AND CONCLUSIONS

The presented shorthand notation for MS-derived lipid structures provides a system to report MS lipidomic data in a standardized way. The system presents two options to assign species in the presence of ambiguities, e.g., when bond type or functional groups may not be resolved by the analysis. Annotation may be either based on assumptions or according to the molecular mass. When using assumptions, these should be based on current biological knowledge and should be made clearly visible. Self-explanatory annotations based on assumptions are advantageous for publications compared with annotation of molecular mass, which should be reserved for the reporting in databases. However it is desirable and, indeed, must be the long-term aim that assumptions are not used.

Establishment of a standardized approach with which to report results from high-throughput MS lipid experiments would facilitate the interchange of information between

different labs, its interpretation, and more importantly, the storage of the information in LIMS systems or lipid-centric public databases. The existence of such resources would also assist bioinformaticians in developing novel tools and/or approaches required to perform analyses using integrative data-mining approaches. At the moment, such studies are extremely cumbersome. We have experienced this problem in the context of the EU LipidomicNet project (<http://www.lipidomicnet.org>), in which more than 20 different European groups are involved. This problem triggered the concept to generate the presented standard nomenclature system. As an outcome of the LipidomicNet project, a new resource called LipidHome (<http://www.ebi.ac.uk/apweiler-srv/lipidhome>) has been developed specifically to accommodate data from high-throughput MS-based lipidomics approaches (15). LipidHome is a database of theoretical lipid species and not only uses this nomenclature but also organizes the lipids into the same hierarchy.


This proposal should be considered as an important further step building upon and contributing to the work of the ILCNC. Currently, the proposal covers only the major lipid category/classes of mammalian lipids. Future proposals could add minor lipid classes or lipid classes of other organisms. 

TABLE 7. Sterols: shorthand notation examples

Sterol	Lipid Class Level Mass ^a	Lipid Species Level	Bond Type Level	LIPID MAPS Structure Level
Cholesterol	ST (386)	ST 27:1_OH	ST 27:1_OH	ST 27:1/OH [5Z,3βOH]
3β-Hydroxycholestenoic acid	ST (416)	ST A27:1_OH	ST A27:1_OH	ST A27:1/OH [5Z,3βOH,25(R),26CO2H]
Lithocholic acid	ST (376)	ST A24_OH	ST A24_OH	ST A24/OH [5βH,3αOH,24CO2H]
Progesterone	ST (314)	ST 21:3_OH2	ST 21:1_O2	ST 21:1/O2 [4Z,3O,20O]
Testosterone	ST (288)	ST 19:2_OH2	ST 19:1_OH_O	ST 19:1/OH/O [4Z,17βOH,3O]
Dehydroepiandrosterone	ST (288)	ST 19:2_OH2	ST 19:1_OH_O	ST 19:1/OH/O [5Z,3βOH,17O]
17β-Estradiol	ST (272)	ST 18:3_OH2	ST 18:3_OH2	ST 18:3/OH2 [1Z,3E,5E,3OH,17βOH]
Taurocholic acid	ST (515)	ST A24_OH3_T	ST A24_OH3_T	ST A24/OH3/T [5βH,3αOH,7αOH,12αOH,24T]
Glycochenodeoxycholic acid	ST (449)	ST A24_OH2_G	ST A24_OH2_G	ST A24/OH2/G [5βH,3αOH,7αOH,24G]
Dehydroepiandrosterone sulfate	ST (368)	ST 19:2_OH2_S	ST 19:1_OH_O_S	ST 19:1/OH/O/S [5Z,3βOH,17O,3S]
24(S)-Hydroxycholesterol 3-sulfate, 24-glucuronide	ST (658)	ST 27:1_OH2_S_GlcA	ST 27:1_OH2_S_GlcA	ST 27:1/OH2/S/GlcA [5Z,3βOH,24(S)OH,3S,24GlcA]
Cholesteryl palmitate	ST (624)	SE 27:1/16:0	SE 27:1/16:0	CE 16:0
Cholesteryl linoleate	ST (648)	SE 27:1/18:2	SE 27:1/18:2	CE 18:2(9Z,12Z)
Zymosteryl oleate	ST (648)	SE 27:2/18:1	SE 27:2/18:1	SE 27:2 [5α,8E,24,3βFA]/18:1(9Z)

^aUncharged molecular mass.

The authors are particularly indebted to members of the ILCNC for valuable suggestions and positive consideration of this shorthand nomenclature. The ILCNC currently consists of Edward A. Dennis, Chair, (US), Robert C. Murphy (US), Masahiro Nishijima (Japan), Takao Shimizu (Japan), Friedrich Spener (Austria), and Michael Wakelam (UK). Shankar Subramanian serves as Informatics Advisor, and Eoin Fahy serves as Director.

REFERENCES

1. Fahy, E., S. Subramanian, H. A. Brown, C. K. Glass, A. H. Merrill, Jr., R. C. Murphy, C. R. Raetz, D. W. Russell, Y. Seyama, W. Shaw, et al. 2005. A comprehensive classification system for lipids. *J. Lipid Res.* **46**: 839–861.
2. Fahy, E., S. Subramanian, R. C. Murphy, M. Nishijima, C. R. Raetz, T. Shimizu, F. Spener, G. van Meer, M. J. Wakelam, and E. A. Dennis. 2009. Update of the LIPID MAPS comprehensive classification system for lipids. *J. Lipid Res.* **50(Suppl.)**: S9–S14.
3. Wenk, M. R. 2010. Lipidomics: new tools and applications. *Cell.* **143**: 888–895.
4. Blanksby, S. J., and T. W. Mitchell. 2010. Advances in mass spectrometry for lipidomics. *Annu. Rev. Anal. Chem. (Palo Alto Calif.)*. **3**: 433–465.
5. Liebisch, G., B. Lieser, J. Rathenber, W. Drobnik, and G. Schmitz. 2004. High-throughput quantification of phosphatidylcholine and sphingomyelin by electrospray ionization tandem mass spectrometry coupled with isotope correction algorithm. *Biochim. Biophys. Acta.* **1686**: 108–117.
6. Brugger, B., G. Erben, R. Sandhoff, F. T. Wieland, and W. D. Lehmann. 1997. Quantitative analysis of biological membrane lipids at the low picomole level by nano-electrospray ionization tandem mass spectrometry. *Proc. Natl. Acad. Sci. USA.* **94**: 2339–2344.
7. Zemski Berry, K. A., and R. C. Murphy. 2004. Electrospray ionization tandem mass spectrometry of glycerophosphoethanolamine plasmalogen phospholipids. *J. Am. Soc. Mass Spectrom.* **15**: 1499–1508.
8. Murphy, R. C., and E. Fahy. 2010. Isoprostane nomenclature: more suggestions. *Prostaglandins Leukot. Essent. Fatty Acids.* **82**: 69–70.
9. Han, X., and R. W. Gross. 2005. Shotgun lipidomics: electrospray ionization mass spectrometric analysis and quantitation of cellular lipidomes directly from crude extracts of biological samples. *Mass Spectrom. Rev.* **24**: 367–412.
10. Pulfer, M., and R. C. Murphy. 2003. Electrospray mass spectrometry of phospholipids. *Mass Spectrom. Rev.* **22**: 332–364.
11. Schwudke, D., J. T. Hannich, V. Surendranath, V. Grimard, T. Moehring, L. Burton, T. Kurzhalia, and A. Shevchenko. 2007. Top-down lipidomic screens by multivariate analysis of high-resolution survey mass spectra. *Anal. Chem.* **79**: 4083–4093.
12. Shaner, R. L., J. C. Allegood, H. Park, E. Wang, S. Kelly, C. A. Haynes, M. C. Sullards, and A. H. Merrill, Jr. 2009. Quantitative analysis of sphingolipids for lipidomics using triple quadrupole and quadrupole linear ion trap mass spectrometers. *J. Lipid Res.* **50**: 1692–1707.
13. Sullards, M. C., Y. Liu, Y. Chen, and A. H. Merrill, Jr. 2011. Analysis of mammalian sphingolipids by liquid chromatography tandem mass spectrometry (LC-MS/MS) and tissue imaging mass spectrometry (TIMS). *Biochim. Biophys. Acta.* **1811**: 838–853.
14. Schuhmann, K., R. Almeida, M. Baumert, R. Herzog, S. R. Bornstein, and A. Shevchenko. 2012. Shotgun lipidomics on a LTQ Orbitrap mass spectrometer by successive switching between acquisition polarity modes. *J. Mass Spectrom.* **47**: 96–104.
15. Foster, J. M., P. Moreno, A. Fabregat, H. Hermjakob, C. Steinbeck, R. Apweiler, M. J. Wakelam, and J. A. Vizcaíno. LipidHome: a database of theoretical lipids optimised for high throughput mass spectrometry lipidomics. *PLoS ONE*. In press.