All-in-one Data-Processing and Interactive Visualizations of Lipid LC-HRMS/MS Data using LipidMatch 4.0

Introduction

Lipid pathways are altered in virtually all disease states, making lipidomics a valuable tool for biomarker discovery and understanding mechanisms of disease. As application of lipidomics gains prevalence, it is essential that software tools adapt to provide high-confidence and high-coverage annotations and be designed to be user friendly. Currently, manual review of lipidomics data is necessary for confident assignment. To address this need, LipidMatch 4.0 was developed. This software provides confident annotations which have been benchmarked against Lipid Annotator, MS-DIAL, and GREAZY, and to our knowledge, provides the most indepth interface for validating annotations and discovering new lipid species

Methods (see Application Note 5994-1356EN)

Lipidomics profiling workflow was used to analyze lipid alterations the Acute Myeloid Leukemia (AML) K562 cell line in response to different drug combinations. Data was acquired on a 6546 LC/Q-TOF with an Agilent 1290 Infinity II LC. Reverse phase chromatography was applied using an Agilent InfinityLab Poroshell 120 EC-C18 (3.0 × 100 mm, 2.7 µm) with a polar phase consisting of water:methanol (9:1) and non-polar phase consisting of acetonitrile:methanol:isopropanol (2:3:5) both with 10 mM ammonium acetate.



MS/MS automatically, and purple are annotated PS species with overlapping masses

Right Top: Each homologous series is defined by a specific lipid class and unsaturation can be viewed in retention time vs m/z plots where outliers (false positives) can easily be determined

Right Bottom: EICs of selected files can also be viewed simultaneously for each selected series, readily showing peak shape, any isomers, and the most dominant members of a class or series

The interactive dataset can be review for lipidomics can be viewed at innovativeomics.com/datasets

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Entire Acquisition and Software Workflow



Annotation Using Isotopic Pattern and MS/MS

TG(16:0_18:1_20:0)+NH4, TG(14:0_18:0_22:1)+NH4, TG(12:0_18:0_24:1)+NH4, TG(14:0_20:0_20:1)+NH4, TG(14:0_18:1_22:0)+NH4, TG(14:0_16:0_24:1)+NH4, TG(16:0_16:0_22:1)+NH4, TG(16:0_16:1_22:0)+NH4, TG(12:0_20:1_22:0)+NH4, TG(12:0_20:0_22:1)+NH4, were all annotated with that rank, and MS/MS evidence shows why all species likely exist under the peak

Right: Annotations of NL peaks

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Conclusions

LipidMatch can be used to rapidly annotate lipids in an automatic fashion and determine unknowns and expand annotation using an interactive visualizer

- Incorporates MS/MS, MS, EICs, homologous series, and retention time
- Has over 300,000 species with fragmentation in libraries; fragment screening and substructure assignment for unknowns
- Many previously unidentified species exist in datasets which can be discovered using the visual interface
- < 5% False Positive



To install the software please visit: Innovativeomics.com/software Questions? Trainings? Collaboration? Contact: jeremykoelmel@gmal.com

Fragment Screening						
Fragment Filter	Intensity					
	71K					
NL_NH3_(18:0)	16K					
NL_NH3_(18:1)	6K					
NL_NH3_(16:0)	6K					
NL_NH3_(20:0)	3K					
NL_NH3_(20:1)	2K					
NL_NH3_3H2O	0K					
NL_NH3_(14:0)	0K					
FA(42:7)-H	0K					
NL_NH3_3H2O	0K					
FA(44:4)-OH-H	0K					
NL_NH3_(22:1)	0K					
NL_NH3_(22:0)	0K					
FA(44:3)-OH-H	0K					
NL_NH3_2H2O	0K					
NL_NH3_15:1(ок 🎽					
<	>					

algorithms cover file conversion, blank filtering

Discovery of Unknowns: Fragment Screening

Ret	ention Time vs	Fragment Scree	agment Screening		
Nam	ne_or_Cl 🖣 🗖	PC ● PE ● PG ● Plasma	nyl 🔶 PS 🌒 SM	Fragment Filter	Intensity
					84,098K
				Phosphorylcholine	30,998K
m/z	1.000			NL_PEheadC2H8N	932K
	1,000			CholineC5H14NO	854K
		Tene o o a		CH2CHN(CH3)3	472K
	800			NL_NH3_(16:0);CLA	404K
		•	NL_NH3_(15:0);CLA	369K	
	600	•	NL_NH3_2H2O_(22	296K	
				NL_NH3_(18:1);CLA	266K
				C2H6O4P	236K
				C41H71O4	219K
	400			NL_PIhead;CLASS:	213K
				FA(9:0)-OH-H2O	188K
	200			NL_NH3_(18:0);CLA	163K
	200			NL_NH4_C3H8O6P	109K
		5 10	15	exp_1;CLASS:PGM	96K
	0			NL_TrimethylAmin	84K
	Retention Time CholesterolC27H45		CholesterolC27H45	82K	
			NU DOL LOOLIONI	C717	

phosphorylcholine m/z 184.073 fragment was used to filter features, indicating PC, SM, ether and oxidized derivates, and other species containing this head group.

Light blue dots are unknowns showing that a significant number of species which were potentially polar lipids with a phosphocholine head group were left unidentified. This signifies the wealth of information which is missing in traditional lipidomics approaches without unknown discovery. It is important to note that some (but not all) of these unknown features, which overlap in *m/z* and retention time with PCs or SMs may be isobaric and hence the 184 peak may come from another species.

