

Inal Chem. Author manuscript; available in PMC 2013 May 20.

Published in final edited form as:

Anal Chem. 2012 January 3; 84(1): 283–289. doi:10.1021/ac202450g.

CAMERA: An integrated strategy for compound spectra extraction and annotation of LC/MS data sets

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Abstract

Liquid chromatography coupled to mass spectrometry is routinely used for metabolomics experiments. In contrast to the fairly routine and automated data acquisition steps, subsequent compound annotation and identification require extensive manual analysis and thus form a major bottle neck in data interpretation. Here we present CAMERA, a Bioconductor package integrating algorithms to extract compound spectra, annotate isotope and adduct peaks, and propose the accurate compound mass even in highly complex data. To evaluate the algorithms, we compared the annotation of CAMERA against a manually defined annotation for a mixture of known compounds spiked into a complex matrix at different concentrations. CAMERA successfully extracted accurate masses for 89.7% and 90.3% of the annotatable compounds in positive and negative ion mode, respectively. Furthermore, we present a novel annotation approach that combines spectral information of data acquired in opposite ion modes to further improve the annotation rate. We demonstrate the utility of CAMERA in two different, easily adoptable plant metabolomics experiments, where the application of CAMERA drastically reduced the amount of manual analysis.

Introduction

Mass spectrometry (MS) is one of the dominant analysis methods for metabolomics experiments. In metabolite profiling studies, a large number of complex samples is analyzed. Typically, samples are separated in prior to ionization and MS-based detection, mostly chromatographically either by gas chromatography (GC) or liquid chromatography (LC). An overview of techniques and applications was given by Dunn¹. Depending on the sample preparation method and the analyzed organism, samples contain anywhere between dozens to thousands of compounds, e.g. the estimated number of metabolites in *E.coli*² is just above 1 000, in human serum³ above 4 000 or 5 000 to 25 000 for higher plants⁴. The coverage within an experiment is much lower due to analytical limitations.

The typical metabolomics data processing pipeline first performs a feature detection step. The term feature describes a two-dimensional bounded signal: a chromatographic peak

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Supporting Information Available Experimental procedures and characterization data for all new compounds. This material is available free of charge via the Internet at http://pubs.acs.org/.

(retention time) and a mass spectral peak (m/z). Several software packages exist for feature detection, for example the closed-source but freely-available MetAlign⁵ or frameworks with open-source licenses such as OpenMS⁶, MZmine⁷ and XCMS⁸. Other packages, some of them specific for LC/MS-based proteomics, have been reviewed elsewhere⁹.

Upon ionization, an individual chemical compound gives rise to one or more ion species, which can be observed in the same mass spectrum. Those ion species include isotopologue ions, fragment ions, and in particular for electrospray ionization (ESI) adduct and cluster ions. A summary can be found in Keller et al. ¹⁰.

For biological interpretation, users are mainly interested in the compounds, rather than the redundancy of the different ion species, which induce an undesired bloat in the number of observed features, e.g. for an *Arabidopsis thaliana* seed extract Böttcher et al. ¹¹ reported 434 features for 180 compounds. The complexity of both the downstream statistical analysis and subsequent compound identification especially in untargeted metabolite profiling experiments is unduly increased.

To address these problems, two additional processing steps are desired for LC/MS data analysis: 1) *grouping* all features which are derived from the same analyte, and 2) *annotation* of the type of ion species. The first step alone achieves both a data reduction and a first estimation of the total number of detectable compounds in a MS analysis. Such an estimate can be used for the optimization of the analytical protocol, similar to Yanes et al. ¹² where the authors used the feature number as optimization criterion. Both steps together can reveal quasi-molecular ions, whose annotation is essential for further metabolite identification, such as elemental composition calculation based on accurate mass and isotope pattern or tandem-MS analysis.

The authors of Brown et al. ¹³ have developed a workflow using the retention time, m/z difference and intensity correlation across samples to group related features, both reducing the number of relevant features down to 50%, and matched 60% of the remaining features against the Manchester Metabolome Database (MMD). Intensity correlation across samples is also used in ?, and a data-reduction of 86% is reported.

Alternatively, similarity across chromatographic peak shapes allows the grouping of related features. Ipsen et al. 15 use a χ^2 test to check for exact co-elution. In case of LC/MS data acquired on TOF instruments with a time-to-digital converter, the test provides p-values for the (un-)certainty of co-elution. The test works best with low ion counts, and the instruments' detector saturation correction had been disabled for this evaluation. ACD/ IntelliXtract 16 is a commercial software solution to cluster features based on their retention time, and the annotation of ion species according to a given rule table.

Both correlation across samples and peak shape analysis techniques are used in the R package ESI¹⁷. A fixed mass difference rule table is used for annotation and detection of isotope peaks. The same approach was later used by Scheltema et al. ¹⁸ for high-resolution LC/MS data. By explicitly removing features exhibiting both similar peak shapes and intensity correlation across samples, they achieved a 60% size-reduction of the feature list.

In this paper we present the CAMERA package, which integrates multiple methods for grouping related features, and uses a dynamic rule table for the annotation of ion species. We evaluate the performance of CAMERA with several validation experiments and demonstrate the analysis of two metabolomics experiments.

Theory, architecture and algorithms

The analysis workflow with CAMERA is shown in Figure 1. The numbering (1–5) describes the typical workflow order. In the next paragraphs the steps are explained in more detail.

Creating compound spectra based on retention time ①

The initial creation of compound spectra has to be fast, if dozens to hundreds of samples with thousands of features have to be processed. We select the most intense feature from the feature table not yet assigned to a compound spectrum and calculate a feature specific retention time window, typically 60% of the chromatographic peak FWHM (full width at half maximum) around the centroid. All features within this range are then included into a new compound spectrum. This step is repeated until all features are assigned to a compound spectrum. The most intense feature usually has the highest signal-to-noise (S/N) ratio, and often provides the most accurate estimate of the centroid and retention time.

Isotopic peak detection and charge state calculation 2

The detection of isotopic patterns is required to deduce the charge states. Within each compound spectrum we calculate a pairwise m/z distance matrix and detect isotopes which exhibit a m/z difference of $1.0033/z^{19}$ and also pass an additional intensity ratio check, described in detail in Supplementary Information S1.

Compound spectrum refinement graph 3

Depending on the chromatographic separation, the resulting compound spectra might still encompass features of two or more closely co-eluting compounds. We use a graph-based algorithm to integrate three more cues for an improved separation (see Figure 2 for an example):

First, we use the chromatographic peak shape similarity. CAMERA uses the raw data to obtain the extracted ion chromatograms (EIC) for each feature, and calculates a pointwise pearson correlation of the intensities between the chromatographic peak boundaries for all pairs of features in a compound spectrum. CAMERA uses the EICs from the sample which had the most intense feature, often the one with the best S/N ratio. Alternatively, the peak shape correlation can be performed for all samples in the experiment. Second, we include the pearson correlation of intensities across all samples for each pair of features in a compound spectrum. Finally we encode the isotope relationship between two features detected in step as 1, and 0 otherwise. These three values are combined as shown in Eq. (1).

Score
$$(x, y) = CAS_{xy} + ISO_{xy} + \frac{1}{N} \sum_{i=1}^{N} CPS_{ixy}$$
 (1)

The score which represents the relationship between two features x and y is the combination of the intensity correlation across samples (CAS) for these two features, the binary encoded presence or absence of an isotope relationship, and the peak shape correlation (CPS_i) calculated for sample i.

In a graph, all features in a compound spectrum, which could still include features of two or more closely coeluting compounds, are represented as nodes, connected by edges with this score as edge weight. Several algorithms for graph separation have been developed, we employ the "Highly-connected-subgraphs" (HCS²⁰) from the R package RBGL or the "label propagation community" (LPC²¹) from the R package igraph. After the graph clustering the initial compound spectrum is split into one refined compound spectrum for each subgraph.

Figure 2 shows an example for a relationship graph before and after separation. Both coeluting compounds were separated completely.

Annotation of adducts, common neutral losses and cluster-ions ①

For ESI, uncharged compounds are ionized through adduct formation with cations or anions or abstraction of protons. In addition, neutral losses occur leading to the formation of fragment ions. An annotation of these ion species reduces the number of features which have to be considered further in the downstream analysis. From at least two annotated ions, the molecular mass can be calculated, which is necessary to search in compound libraries, or to calculate the elemental composition of the neutral compound.

CAMERA uses a dynamic rule set, which is created from the combination of lists of observable ions. Each rule describes a specific ion species with the mass difference to the molecular mass, ion charge and the number of molecules the ion species contains. All m/z differences within a compound spectrum are matched against the dynamic rule set. Matches with the same molecular mass hypothesis (below a given relative error) are combined into hypothesis groups. If no peaks can be explained via the rules, a reliable annotation is impossible. CAMERA does not use ad-hoc heuristics such as assuming that the most intense feature in a spectrum is the [M+H]+-ion. Afterwards conflicting hypothesis groups are resolved as described in Supplementary Information S2.

Combining data from opposite ion modes for verification (5)

In metabolite profiling samples are often measured in both positive and negative ion mode, to increase the metabolite coverage. Although some compounds ionize in only one mode, many compounds are detectable in both. In these cases the complementary ions provide further evidence for the quasi-molecular ion.

CAMERA includes a novel annotation verification algorithm using compound spectra measured in both ion modes. The algorithm calculates m/z differences for all features of corresponding compound spectra from both modes within a retention time window. These differences are matched against a second, cross-polarity rule table. If a cross-polarity rule matches, it will either 1) annotate two previously un-annotated ions, e.g. [M+H]⁺ and [M-H]⁻, or 2) verify an existing annotation or 3) conflict with an existing annotation. In the latter case the existing annotation is replaced. The cross-polarity rule table should only contain common and trusted combinations, because these rules can override the single-polarity annotations.

Documentation and availability

CAMERA is implemented in R, the packages for Windows (both 32 and 64 bit), Mac OS and Linux are available from the Bioconductor repository²² since release 2.4 in 2009.

Experimental section

Reagents and Materials

All solvents used for sample preparation and analyses were of LC/MS-grade quality (CHROMASOLV, Fluka). A list of standard compounds used for the recovery experiment including sum formulas, molar masses, PubChem IDs and suppliers can be found in the Supplemental Information S3. L-Tryptophan-2',4',5',6',7'- D_5 (98%) was purchased from Cambridge Isotope Laboratories. *Arabidopsis thaliana* (ecotype Col-0) was grown for six weeks on a soil/vermiculite mixture (3/2) in a growth cabinet with 8 h light (150 μ E $m^{-2}s^{-1}$) at 22°C and 16 h dark at 20°C. Seeds of *Brassica napus*, *Brassica oleracera* and *Brassica rapa* were kindly provided by D. Strack, Department of Secondary Metabolism,

Leibniz Institute of Plant Biochemistry, Halle. All other seeds were obtained from local distributors. Procedures for extraction of leaf and seed material are provided in Supplemental Information S4.

Feeding Experiments

Plants were sprayed with 5 mM aqueous silver nitrate solution 1 h after the beginning of the light period. After 5 h, twenty-five rosette leaves originating from five individual plants were excised at the petiole and immersed in PCR tubes containing either 200 μL water or 200 μL of an aqueous [RING-D₅]-Trp solution (1 mM), respectively. Leaves were incubated for an additional 2 d in a growth cabinet under the same conditions as described above. Individual leaves of the same treatment were pooled, frozen in liquid nitrogen and stored at -80° C until analysis.

UPLC/ESI-QTOFMS Analysis

Chromatographic separations were performed on an Acquity UPLC system (Waters) equipped with a HSS T3 column (100×1.0 mm, particle size $1.8 \,\mu$ m, Waters) applying the following binary gradient at a flow rate of $150 \,\mu$ L min⁻¹: 0–1 min, isocratic 95% A (water/formic acid, 99.9/0.1 (v/v)), 5% B (acetonitrile/formic acid, 99.9/0.1 (v/v)); 1–16 min, linear from 5 to 95% B; 16–18 min, isocratic 95% B; 18–20 min, isocratic 5% B. The injection volume was 2.7 μ L (full loop injection). Eluted compounds were detected at a spectra rate of 3 Hz from m/z 100–1000 using a MicrOTOF-Q-I hybrid quadrupole time-of-flight mass spectrometer (Bruker Daltonics) equipped with an Apollo II electrospray ion source in positive and negative ion mode. We made sure that the concentration of the samples do not lead to saturation of the MS detector system, which is known to cause shifts of m/z and retention time centroids of the features, leads to truncated chromatographic peak profiles and distorted isotopic patterns. For detailed instrument settings and acquisition of collision-induced dissociation mass spectra see Supplemental Information S4.

LC/MS data preprocessing

Processing of raw data was performed with XCMS package⁸. For the feature detection we used the XCMS *centWave*²³ algorithm with the following parameters: snthresh = 6, ppm = 30, peakwidth = (5,12), prefilter = (2,200). The feature alignment was performed with the standard *group.density* algorithm from XCMS with bw = 3 and mzwid = 0.015. Afterwards, each dataset was processed with CAMERA functions in the following order *groupFWHM*, *findIsotopes*, *groupCorr*, *findAdducts* using standard parameters. Supplemental Information S9 provides runtime measurements of CAMERA.

Results and discussion

We evaluated CAMERA with several experiments. First, using standards we analyzed the performance of compound spectrum creation and success rate of molecular mass annotation. Then, we processed the output from two different experiments, where the CAMERA results were used to perform targeted profiling of phenolic choline esters and tryptophan-derived metabolites, respectively.

Evaluation on known compound mixture

For the evaluation we used a mixture of 39 known compounds (short: MM39), covering a broad mass range between 161 and 822 Da and different physico-chemical properties (see Supplementary Information S3). The mixture was measured as pure solution and spiked in different concentrations $(20,5,1,0.2 \, \mu M)$ into methanolic extracts of *Arabidopsis thaliana* leaves, to simulate a realistically complex matrix.

The first evaluation focuses on the extraction of compound spectra, which requires a data set and a gold standard of true positive and true negative cases, i.e. pairs of peaks which should or should not be part of the same compound spectrum. Because it is very tedious to manually create a gold standard of a sufficiently large number of features from different compounds which co-elute, we altered the retention times in the raw data files to artificially force "co-elution" for this evaluation. We used only those peaks in the compound spectra of the MM39 for which a reliable annotation exists, to rule out false positives, and randomly collected peaks from the remaining file with unrelated retention times to assemble a negative set. These data sets allowed us to calculate the precision and recall for the collection of compound spectra. The default peak shape correlation threshold of 0.75 results in a recall of 0.93, with a precision of 0.48. We also analyzed the influence of different acquisition parameters (scan rates varied from 0.5 Hz to 6 Hz). Precision and recall had a standard deviation of 0.07 and 0.03 respectively across the different conditions, see the Supplementary Information S5 for details, including the ROC curves.

We then evaluated how successfully CAMERA could annotate the different ion species from a compound, which is required for the calculation of the molecular mass. We created baseline values for all *annotatable* compounds: we define an annotatable compound as observed to produce 1) the protonated molecular ion, 2) its first isotopic peak (required to calculate the charge state), and 3) the most prominent adduct ion (observed at $20\mu M$). This strategy serves as gold standard to determine the number of annotatable compounds in the MM39 measurement; 35 out of 39 compounds pass the above requirements for the $20\mu M$ positive mode measurement. If the mixture is diluted two orders of magnitude to $0.2\mu M$, many peaks drop below the detection limit and only 10 compounds remain annotatable.

CAMERA was able to detect the correct molecular mass in 90% of all annotatable compounds in either positive or negative mode across all concentrations. After combining results from both ionization modes, CAMERA correctly determined molecular mass for all annotatable compounds, and additionally for four compounds that were not on the gold standard list. Because the manual assignment of corresponding features in both positive/negative mode data is quite cumbersome, the combined annotation promises to annotate more compounds than a human operator could do on a routine basis.

It is remarkable that the complex leaf matrix does not have an observable negative effect on the annotation performance. Table 1 shows the results for the individual concentrations. On closer inspection, the missing molecular mass annotations have few common causes: they occurred either because the compound spectrum did not contain enough explained features, or in other cases several hypotheses had the same precedence scores and we did not count those as successful. For some compounds the compound spectra contained only the molecular ion and fragment ions, but no further adducts. In this case the compound cannot be annotated directly, unless the neutral loss is added to the rule set. Supplemental Information S10 shows an overview of the frequency of annotated adducts we observed.

In the measurements with different scan rates we found that CAMERA missed up to two correct annotations in those cases where either an essential (albeit low abundant) feature was not found by the feature detection algorithm, or features were assigned to a different compound spectrum, especially in the case of lower scan rates where chromatographic peaks were covered by only a few scans. This suggests that CAMERA can also be used for LC/MS measurements with a low scan rate, e.g. on Orbitrap instruments at high resolution.

Case Study I: Screening for phenolic choline esters in brassicaceous seeds

In this section we use untargeted LC/MS profiles of seeds from some *Brassicacea*, and demonstrate how CAMERA can be used to perform a neutral loss screen for phenolic choline esters as a targeted analysis strategy on a TOF instrument.

Phenolic choline esters accumulate in considerable amounts in seeds of many plant species within the *Brassicacea*²⁴ family. Representatives of this compound class structurally characterized so far include substituted cinammoyl and benzoyl cholines, which are further diversified by glycosylation or oxidative coupling to monolignols. A total of 30 phenolic choline esters could be identified in seeds of the model plant *Arabidopsis thaliana* and the oil crop *Brassica napus* using LC/ESI-tandem mass spectrometry. A study of the fragmentation behaviour of phenolic choline esters under positive-ion electrospray-CID conditions revealed a loss of trimethylamine as initial fragmentation step (see Supplemental Information S6). The formation of the corresponding fragment ion [M–C₃H₉N]⁺ requires different collision energies depending on the compound. However, it is also inducible by insource CID allowing a systematic screening for phenolic choline esters even by single-stage MS. For that purpose, the neutral loss detection has to be performed *in silico* after data acquisition by searching for a given *m/z*-difference between pairs of peaks within a set of extracted compound spectra.

We prepared extracts from seeds of twelve different *Brassicacea* species and cultivars and analyzed each extract by UPLC/ESI(+)-QTOFMS at four different in-source CID voltages (0, 30, 60 and 90 V) to induce fragmentation of a broad range of phenolic choline esters. All 48 raw data files were preprocessed with XCMS with snthresh = 5 and ppm = 20, other parameters analogous to evaluation section. Due to the large number of chromatographically unresolved compounds eluting near the void time, compounds with a retention time below 45 s were excluded from further analysis. Afterwards CAMERA was used to create the compound spectra. Each compound spectrum was then screened for peak pairs displaying a m/z-difference of 59.074 ± 0.015 corresponding to a neutral loss of trimethylamine.

In addition, we included a m/z-difference of 221.126±0.015, related to a successive loss of trimethy lamine and anhydrohexose (162.053 Da), because [M-C₃H₉N]⁺-type ions formed from 4-O-hexosylated phenolic choline esters are known to readily eliminate their hexose moiety²⁵. After alignment of positively screened peak pairs, the elimination of isotopic peak pairs and application of a reasonable intensity threshold (1000 counts) we detected a total of 90 putative choline esters. A data matrix including mass-to-charge ratios of proposed molecular ions, retention times and intensities can be found in Supplemental Information S6. It should be noted, that the number of putative candidates is rapidly increasing when tolerance thresholds for m/z-differences were increased. Therefore, use of mass analyzers providing adequate resolution and mass accuracy, such as TOFMS, is mandatory for this type of screening approach in order to assure a highly specific neutral loss detection. In order to evaluate the obtained candidate list, previously published analytical data of choline esters from seeds of A. thaliana and B. napus was used for compound annotation²⁵. Out of 31 choline esters described recently, we were able to retrieve 22 from our list. Seven choline ester were consistently detected across all samples. Five of them could be annotated (Table 2), including sinapoylcholine, which is known to occur as major phenolic choline ester in seeds of numerous *Brassicacea* species²⁴. Although a rigorous evaluation is not possible because the choline ester composition of an alyzed seeds is unknown, recovery of the majority of compounds described in the literature demonstrates the usability of CAMERA for such a screening approach. An additional advantage of this approach compared to triple quadrupole MS-based neutral loss scanning techniques, is that any number of neutral losses can be simultaneously detected after data acquisition, allowing screening for a broad range of compound classes.

Case study II: Identification of Trp-derived metabolites from *Arabidopsis thaliana* after [ring-D₅]-Trp feeding

In vivo administration of isotope-labeled substrates combined with mass spectrometry-based analysis represents a powerful tool to investigate biochemical pathways. The detection of an isotope-labeled substrate incorporated into a known metabolite allows to deduce a biosynthetic relationship between the fed precursor and the metabolite under study. Nontargeted screening for metabolites and their isotopologues after partial isotope-labeling of an endogenous precursor pool has been applied to explore unknown biosynthetic pathways and to discover novel intermediates and products. ²⁶

To demonstrate the applicability of the CAMERA package for such an analytical approach the metabolic fate of the aromatic amino acid Trp was studied in the model plant $Arabidopsis\ thaliana\ using\ [ring-D_5]$ -Trp as isotope-labeled tracer. In Arabidopsis, Trp represents an important precursor for a variety of secondary metabolites including the phytoanticipin indol-3-ylmethyl glucosinolate and the phytoalexin camalexin (3-thiazol-2'-ylindole).

Arabidopsis leaves were sprayed with silver nitrate to induce expression of Trpmetabolizing enzymes, detached from plants and fed with [ring-D₅]-Trp or water as control. Methanolic extracts of label-fed and control leaves were analyzed in duplicate by UPLC/ ESI-OTOF-MS in positive and negative ion mode. In order to identify Trp-derived metabolites, the raw data was processed with XCMS and CAMERA to extract compound spectra and annotate isotopic peaks within these spectra. Afterwards, deisotoped compound spectra extracted from data sets of label-fed leaves were screened for feature pairs that exhibit an m/z-difference of 5.031, reflecting the exchange of five hydrogen atoms by deuterium. Since deuterium labelling can slightly shift retention times, we searched for these feature pairs between compound spectra within a sliding retention time window of 8 s. For this purpose we created a dedicated script using CAMERA functionality for the positive/ negative polarity combination. We also included the m/z-difference of 4.025, because indole ring hydroxylation (a frequently observed transformation in Trp metabolism in Arabidopsis) results in a loss of one of the five deuterium labels. The retention time for Trp-candidates was restricted between 45 and 600 s. All features related to unlabeled Trp-metabolites have to be detectable in both label-fed and control samples whereas the labeled ones in label-fed samples only, see Figure 3. After those filtering steps 46 putative Trp-derived metabolites could be identified in positive ion mode and 34 in negative ion mode. Corresponding candidate lists including compound annotation can be found in Supplemental Information S7.

To verify the obtained candidate lists tandem mass spectra of quasimolecular ions of putative pairs of non-labeled and labeled metabolites were acquired and compared (Supplemental Information S8). Because of low peak intensities or low incorporation rates, only 19 candidate pairs could be rigorously verified following this strategy. Together with literature data, a total of 23 Trp-metabolites could be identified, of which 20 were already known from the literature. This case study clearly demonstrates applicability of CAMERA for such a screening approach, even in case of a retention time shifts when using deuterium labels.

Conclusion

The CAMERA package is designed to post-process XCMS feature lists, and to collect all features related to a compound into a compound spectrum. For this, a set of algorithms has been implemented in CAMERA, such as the fast retention time-based grouping, but also a novel, graph-based algorithm to integrate the peak shape analysis, isotopic information and

intensity correlation across samples. The automatic sample selection avoids poor results if compounds have a low intensity (or are absent) in some samples. The ion species annotation uses a dynamic rule set, and a new strategy to combine spectral information from samples measured in positive and negative ion mode, resulting in both more and more reliable ion species annotation. We evaluated the reliability of the molecular mass calculation, and found a 90% success rate for MM39 in different concentrations, both pure and after spiking the mixture at various concentrations into a complex *A. thaliana* leaf extract.

Finally, we performed two experiments, demonstrating advanced analyses which can be performed with CAMERA. The first case study essentially performed a neutral loss screen for putative phenolic choline esters using multiple in-source voltages to induce fragmentation. 90 putative choline esters were detected. The second case study demonstrated the search for mass differences as a result of [ring-D₅]-Trp feeding in *A. thaliana* leaves. CAMERA was used to detect pairs of features indicating 46 Trp-derived metabolites. In addition to 20 already known compounds, three new ones were found and verified with tandem-MS. Both studies can easily be adopted to other compound classes and metabolites.

The CAMERA packages for Windows, Mac OS and Linux, manuals and tutorials are freely available from the Bioconductor repository and its mirrors under the open source GPL license.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This work was supported by the California Institute of Regenerative Medicine (Grant TR1-01219), the National Institutes of Health (Grants R24 EY017540-04, P30 MH062261- 10, and P01 DA026146- 02), and the Department of Energy (Grants FG02-07ER64325 and DEAC0205CH11231).

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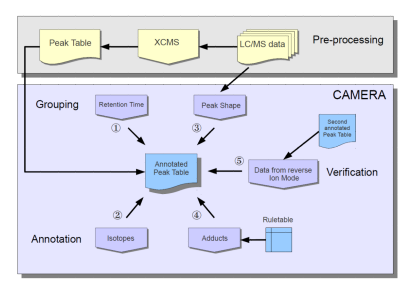


Figure 1.
The CAMERA workflow for LC/MS data analysis. Raw files are preprocessed with XCMS (upper part) and the resulting feature lists are passed to CAMERA. The feature grouping steps integrate Retention Time ① and Chromatographic Peak Shape ③. Features are identified as isotopic peak ②, and adducts are annotated ④ using a dynamic rule table. Optionally, the annotation can be verified ⑤ with LC/MS data acquired in opposite ion mode.

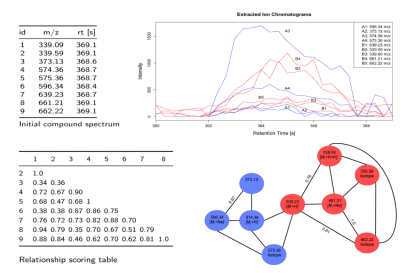


Figure 2. Schematic clustering of low intensity features initially grouped by retention time into a single compound spectrum. Top left: the features, initially grouped by retention time. Top right: the EICs of all features. The labels A and B correspond to the result after graph clustering. Bottom left: the scoring matrix, used as edge weights in the graph. Bottom right: the relationship graph, where edges indicate an above-threshhold score. The node labels include the ion species annotation, and the node color shows the graph separation after refinement with the *LPC* algorithm (A=blue, B=red).

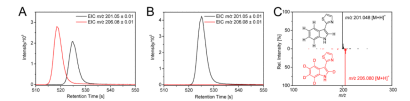


Figure 3. Identification of the phytoalexin camalexin as Trp-derived metabolite in silver nitrate-treated /Arabidopsis thaliana/ leaves using [ring- D_5]-Trp as isotope-labeled tracer. Extracted ion chromatograms (EICs) corresponding to the protonoted molecular ions of camalexin (black) and D_5 -camalexin (red) obtained from UPLC/ESI(+)-QTOFMS analyses of extracts of [ring- D_5]-Trp-fed leaves (A) and control leaves (B). Extracted compound spectra of camalexin and its isotopologue are shown in the right picture (C).

Table 1

Calculation of molecular mass for the MM39 compound mixture analyzed by UPLC/ESI-QTOFMS in positive and negative ion mode, either in pure solvent or spiked at different concentrations into a *Arabidopsis thaliana* leaf extract. The number of annotatable compounds is shown in brackets. In the combined case the annotations of the positive mode are verified and augmented with the negative mode data.

	in solvent	spiked into leaf extract				Overall	
	20 μΜ	20 μΜ	5 μΜ	1 μΜ	0.2 μΜ		
ESI(+)	32 (35)	29 (32)	24 (28)	18 (21)	10 (10)	113 (126)	89.7%
ESI(-)	15 (19)	18 (18)	15 (16)	6 (6)	2 (3)	56 (62)	90.3%
ESI(+/-)	36	35	28	23	15	137	

 Table 2

 Five phenolic choline esters found and annotated in all twelve brassicaceous seeds.

m/z	t_s [s]	NL	elemental comp.	annotation	
280.15	275	-59	C ₁₅ H ₂₂ NO ₄ ⁺	FC	
310.16	279	-59	$C_{16}H_{24}NO_5^+$	SC	
458.21	403	-59	$C_{25}H_{32}NO_7{^+}\\$	FC(5-8')G	
472.21	221	-221	$C_{22}H_{34}NO_{10}^{+}$	SC 4-O-Hex	
476.23	303	-59	$C_{25}H_{34}NO_{8}{^{+}} \\$	FC(4-O-8')G	

NL, neutral loss; FC, ferulolycholine; SC, sinapoylcholine; G, guaiacyl; Hex, hexose