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# Chemically informed analyses of metabolomics mass spectrometry data with Qemistree

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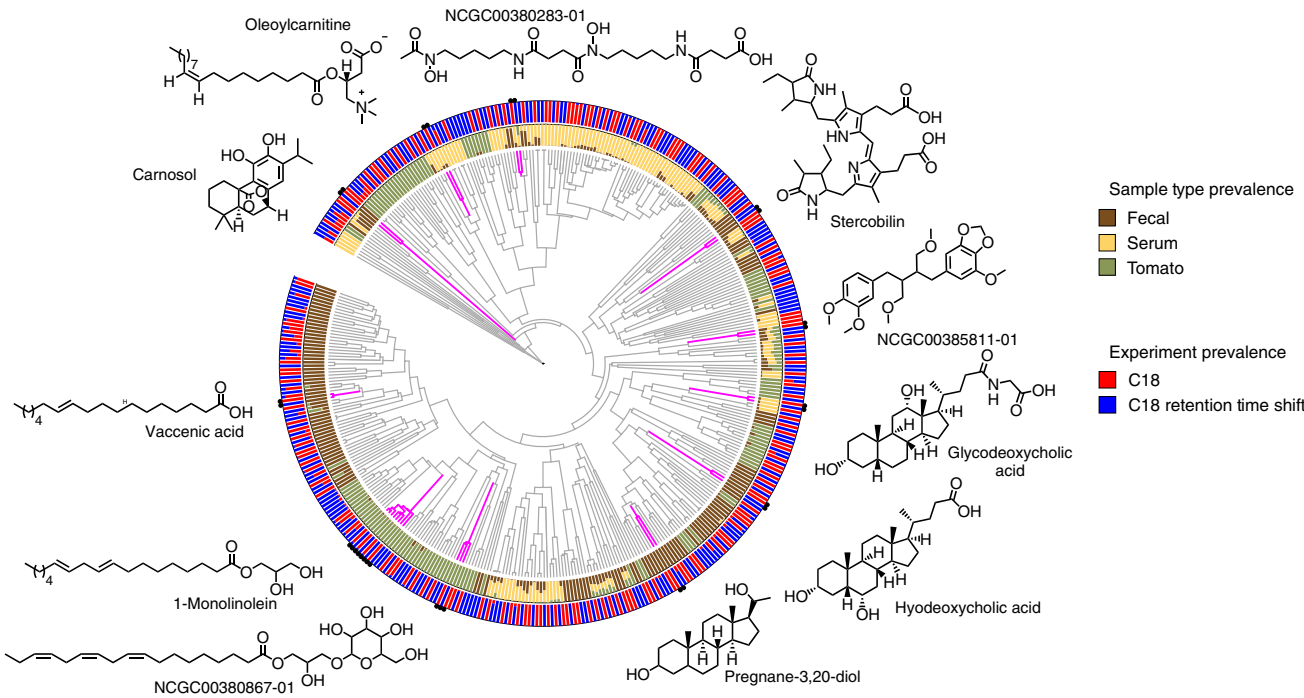
**Untargeted mass spectrometry is employed to detect small molecules in complex biospecimens, generating data that are difficult to interpret. We developed Qemistree, a data exploration strategy based on the hierarchical organization of molecular fingerprints predicted from fragmentation spectra. Qemistree allows mass spectrometry data to be represented in the context of sample metadata and chemical ontologies. By expressing molecular relationships as a tree, we can apply ecological tools that are designed to analyze and visualize the relatedness of DNA sequences to metabolomics data. Here we demonstrate the use of tree-guided data exploration tools to compare metabolomics samples across different experimental conditions such as chromatographic shifts. Additionally, we leverage a tree representation to visualize chemical diversity in a heterogeneous collection of samples. The Qemistree software pipeline is freely available to the microbiome and metabolomics communities in the form of a QIIME2 plugin, and a global natural products social molecular networking workflow.**

Molecular networking<sup>1</sup>, introduced in 2012, was one of the first data organization approaches to visualize the relationships between tandem mass spectrometry (MS/MS) fragmentation spectra. In molecular networking, relationships between similar MS/MS spectra are visualized as edges. As MS/MS spectral similarity indicates chemical structural similarity<sup>1</sup>, chemical structural information can thus be represented as a network and chemical relationships can be visualized. This approach forms the basis for the web-based MS infrastructure, global natural products social molecular networking<sup>2</sup> (GNPS) (<https://gnps.ucsd.edu/>), which sees ~200,000 new accessions per month. Molecular networking has successfully been used for a range of applications<sup>3</sup> in drug discovery, natural products research, environmental monitoring, medicine and agriculture. To tap into the chemistry of complex samples through metabolomics, a subset of MS/MS spectra can be annotated by spectral library matching or by using *in silico* approaches. While molecular networking facilitates the visualization of closely related molecules in molecular families, the inference of chemical relationships at a dataset-wide level and in the context of diverse sample metadata requires complementary representation strategies. To address this need, we developed an approach that uses fragmentation trees<sup>4</sup> and machine learning<sup>5</sup> to calculate all pairwise chemical relationships. These chemical relationships are represented as a chemical tree that can be visualized in the context of sample metadata and molecular annotations obtained from spectral

matching and *in silico* annotation tools. We show that such a chemical tree representation enables the application of various tree-based tools, originally developed for analyzing DNA sequencing data<sup>6–9</sup>, for exploring mass spectrometry data.

Here, we introduce Qemistree (pronounced ‘chemis-tree’) software that constructs a chemical tree based on predicted molecular fingerprints from MS/MS fragmentation spectra<sup>10</sup>. Molecular fingerprints are vectors where each position encodes a substructural property of the molecule, and recent methods allow us to predict molecular fingerprints from tandem mass spectra<sup>11–15</sup>. In Qemistree, we use SIRIUS<sup>16</sup> and CSI:FingerID<sup>13</sup> to obtain predicted molecular fingerprints. Users can first perform feature detection<sup>17,18</sup> to generate a list of observed ions with associated peak areas and MS/MS fragmentation spectra, referred to as chemical features henceforth, to be analyzed by Qemistree (Extended Data Fig. 1). Only chemical features with MS/MS data are included; features with only MS1 (precursor mass) are not considered. SIRIUS then determines the molecular formula of each feature using the isotope and fragmentation patterns and estimates the best fragmentation tree explaining the fragmentation spectrum. Subsequently, CSI:FingerID operates on the fragmentation trees using kernel support vector machines to predict molecular properties (2,936 properties, Supplementary Dataset 1). We use these molecular fingerprints to calculate pairwise distances between chemical features and hierarchically cluster the fingerprint vectors to generate a tree representing their chemical

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**Fig. 1 | Qemistree mitigates aspects of technical artifacts by coclustering structurally similar molecules across MS runs.** A chemical tree based on predicted molecular fingerprints representing the structural relationships between compounds detected in the evaluation dataset. The outer ring shows the relative prevalence of molecules stratified by the MS run; the inner ring shows the same stratified by fecal, serum and tomato samples in the evaluation dataset. All structures shown are spectral reference library matches obtained from feature-based molecular networking<sup>7,18</sup> in GNPS (level 2 or 3 according to the 2007 Metabolomics Standards Initiative<sup>40</sup>). Note that untargeted MS is blind to stereochemistry and often regiochemistry (for example, double bonds in a fatty acid); therefore, molecules could be related isomers of the illustrated structures.

structural relationships. Although alternative approaches to hierarchically cluster features based on cosine similarity of fragmentation spectra exist<sup>19–21</sup>, we use molecular fingerprints predicted by CSI:FingerID for this. Previous work has shown that CSI:FingerID outperforms other tools for automatic in silico structural annotation<sup>22</sup>. Therefore, we leverage it to search molecular structural databases to provide complementary insights into structures when no match is obtained against spectral libraries. Subsequently, we use ClassyFire<sup>23</sup> to assign a five-level chemical taxonomy (chemical kingdom, superclass, class, subclass and direct parent ontology) to all molecules annotated via spectral library matching and in silico prediction (Supplementary Tables 1 and 2 include an assessment of improved annotation rates as a result of in silico annotations).

Phylogenetic tools such as iTOL<sup>24</sup> can be used to visualize Qemistree trees interactively in the context of sample information and feature annotations for easy data exploration. The outputs of Qemistree can also be plugged into other workflows in QIIME2 (ref. <sup>25</sup>) (many of which were originally developed for microbiome sequence analysis) or in R, Python and so on for system-wide metabolomic data analyses<sup>6,7,9,26</sup>. In this study, we apply Qemistree to perform chemically informed comparisons of samples in the presence of technical variation such as chromatographic shifts that commonly affect MS data analysis. Additionally, we exemplify the use of a tree-based representation to visualize and explore chemical diversity using a heterogeneous collection of food products. Qemistree can be used iteratively to incorporate multiple datasets without the need for cumbersome reprocessing (such as repeated feature detection or retention time alignment), allowing for large-scale dataset comparisons. Qemistree is available to the microbiome community as a QIIME2 plugin (<https://github.com/biocore/q2-qemistree>) and the metabolomics community as a workflow on GNPS<sup>2</sup> (<https://ccms-ucsd.github.io/GNPSDocumentation/qemistree/>).

The chemical tree from the GNPS workflow can be explored interactively using the Qemistree-GNPS dashboard (<https://qemistree.ucsd.edu/>; see Methods).

## Results

**Resolving technical variation using chemical relationships.** To verify that molecular fingerprint-based trees correctly capture the chemical relationships between molecules, we designed an evaluation dataset using four distinct biological specimens: two human fecal samples, a tomato seedling sample and a human serum sample. Samples were prepared by combining them in binary, tertiary and quaternary mixtures in various proportions to generate a set of diverse but related metabolite profiles (Supplementary Table 3). Untargeted MS/MS was used to analyze the chemical composition of these samples and obtain fragmentation spectra. The MS experiments were performed twice using different chromatographic elution gradients, causing a retention time shift between the two runs (Extended Data Figs. 2 and 3). Processing the data of these two experiments with traditional LC–MS-based pipelines leads to the same molecules being detected as different chemical features in downstream analysis. Figure 1 shows the analysis of three different sample types to demonstrate this. In Extended Data Fig. 4, we highlight how these technical variations make the same samples appear chemically disjointed.

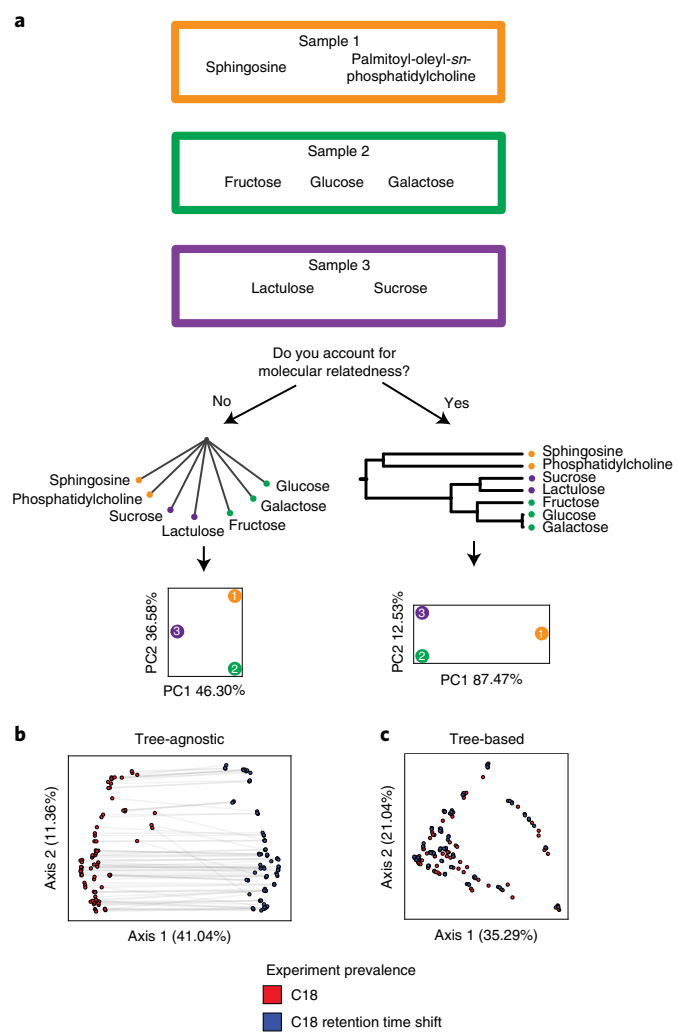
Using Qemistree, we mapped each of the spectra in the two chromatographic conditions (batches) to a molecular fingerprint, and organized these in a tree structure (Fig. 1). Because molecular fingerprints are independent of retention time shifts, spectra are clustered based on their chemical features with spectral library matches (typically 1–20% of all features, depending on how well the sample type has been investigated) or other forms of

annotation (for example, substructure Mass2Motifs<sup>27</sup>) could also be used to compare the chemical composition of samples across different MS runs. Qemistree improves on this by enabling the use of all MS/MS spectra with molecular fingerprints (86.90% in these data at the present time, Supplementary Table 1) for downstream comparative analyses, by not constraining analysis to the chemical features with spectral matches only. This tree structure can be decorated using sample type descriptions, chromatographic conditions, spectral matches obtained from molecular networking in GNPS (when available) and any other chemical annotations<sup>23,27</sup>. Figure 1 shows that similar chemical features were detected exclusively in one of the two batches. However, based on the molecular fingerprints, these chemical features were arranged as neighboring tips in the tree regardless of the retention time shifts. This result shows how Qemistree can reconcile and facilitate the comparison of datasets acquired on different chromatographic gradients.

**Tree-guided system-wide comparisons in metabolomics.** Having demonstrated Qemistree's practical use on biologically inspired synthetic datasets, we now turned to a conceptual example illustrating the general principle. We demonstrated an application of a chemical hierarchy in performing chemically informed comparisons of metabolomics profiles. In standard metabolomic statistical analyses, each molecule is assumed unrelated to the other molecules in the dataset. Some of the pitfalls of this assumption are highlighted in Fig. 2a. Consider a scenario where we want to compare samples 1–3. An analysis schema that does not account for the chemical relationships among the molecules in these samples (Fig. 2a, left), will assume that the sugars in samples 2 and 3 are as chemically related to the lipids in sample 1 as they are to each other. This would lead to the naive conclusion that samples 1 and 2, and samples 2 and 3 are equally distinct, yet from a chemical perspective they are not. On the other hand, if we account for the fact that sugar molecules are more chemically related to one another than they are to lipids, we can obtain a chemically informed sample-to-sample comparison.

The chemical structural compositional similarity (CSCS)<sup>28</sup> was developed to compute pairwise sample-to-sample comparison by considering cosine similarity of MS/MS spectra from molecular networking. Here, we use a tree-based approach to account for chemical relationships, which allows us to adopt phylogeny-based tools for metabolomics analyses (Supplementary Table 4). Specifically, we first constructed a tree of chemical similarities by hierarchical clustering molecular fingerprints from CSI:FingerID (using pairwise Euclidean distance between fingerprint vectors, see Methods). This tree is analogous to phylogenetic trees used in ecology, such that the tips of the tree are molecules (instead of species). We then computed weighted UniFrac<sup>9</sup> distances (a tree-based metric that has widely been used in microbial ecology to compare microbiomes) to compare metabolomic profiles. In Fig. 2a, we show that by using a tree of chemical relationships between molecules in samples 1–3, we can visualize that sample 1 is chemically very distinct (along PC1 in a principal component analysis) from samples 2 and 3.

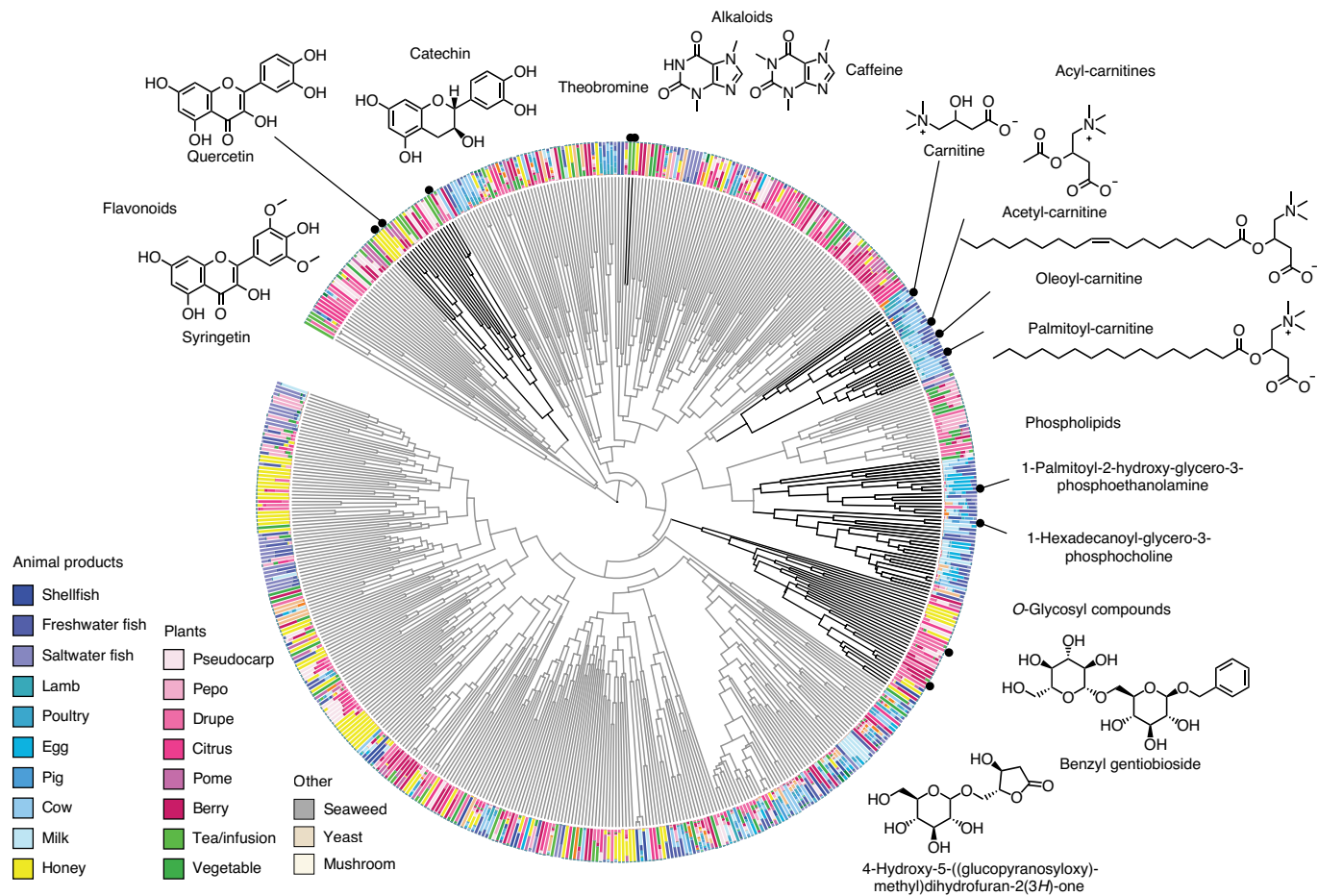
Returning to our evaluation dataset, we can highlight the importance of comparing samples by accounting for their molecular relatedness. Principal coordinates analysis (PCoA) of the evaluation dataset (including both pure samples and sample mixtures,  $N=162$ ) that ignores the tree structure (Fig. 2b) performs far worse than the Qemistree PCoA that uses the tree (Fig. 2c). With the structural context provided by Qemistree, the differences between replicates across batches are comparable to the within-batch differences (Extended Data Fig. 5). The retention time shift in this dataset leads to a strong signal due to chromatography conditions that obscures the biological relationships among the samples (permutational analysis of variance (ANOVA); tree-agnostic<sup>29</sup> pseudo  $F=120.75$ ,  $P=0.001$  versus tree-informed<sup>9</sup> pseudo  $F=18.2239$ ,  $P=0.001$ ).



**Fig. 2 | The pitfalls of assuming equal relatedness of molecules and the advantages of a chemical tree for sample comparison.** **a**, A scenario where the goal is to compare the chemical composition in three samples, and the consequences of accounting for or ignoring molecular relatedness. **b,c**, PCoA of all samples ( $N=162$ ) in the evaluation dataset colored by chromatography conditions. PCoA plot using tree-agnostic (Bray-Curtis<sup>29</sup>) distances that do not account for the chemical relationship between features detected across chromatography conditions (**b**) and tree-based (Weighted UniFrac<sup>9</sup>) distances, which are based on the hierarchical relationships between molecules in the evaluation dataset (**c**).

We observed and remediated a similar pattern originating from plate-to-plate variation in a recently published study investigating the metabolome and microbiome of captive cheetahs<sup>30</sup> (Extended Data Fig. 6). In this study, placing the molecules in a tree using Qemistree reduced the observed technical variation (Extended Data Fig. 6a,c), and highlighted the dietary effect that was expected (Extended Data Fig. 6b,d). These results show how systematic and spurious molecular differences can be mitigated in an unsupervised manner using chemically informed distance measures based on a tree structure.

**Visualizing chemical prevalence in heterogeneous datasets.** As a case study demonstrating the use of Qemistree on a set of biological specimens, we used the platform to explore chemical diversity in food samples collected in the Global FoodOmics initiative (<http://globalfoodomics.org>). Understanding the chemical relationships between different foods is challenging because most molecules



**Fig. 3 | A chemical hierarchy of food-derived compounds based on predicted molecular fingerprints.** A chemical tree based on molecular fingerprints representing the structural relationships between chemical features (tree tips) detected in food products (single ingredient, that is, simple foods  $N=119$ ). The tree is pruned to only keep tips that were assigned a structural annotation (SMILES) by either a MS/MS spectral library matching or in silico using CSI:FingerID. All structures shown are spectral reference library matches obtained from feature-based molecular networking in GNPS. The outer ring shows the relative abundance of each compound across a diverse range of food sources. We highlight clusters of compounds that are characteristic of specific food sources.

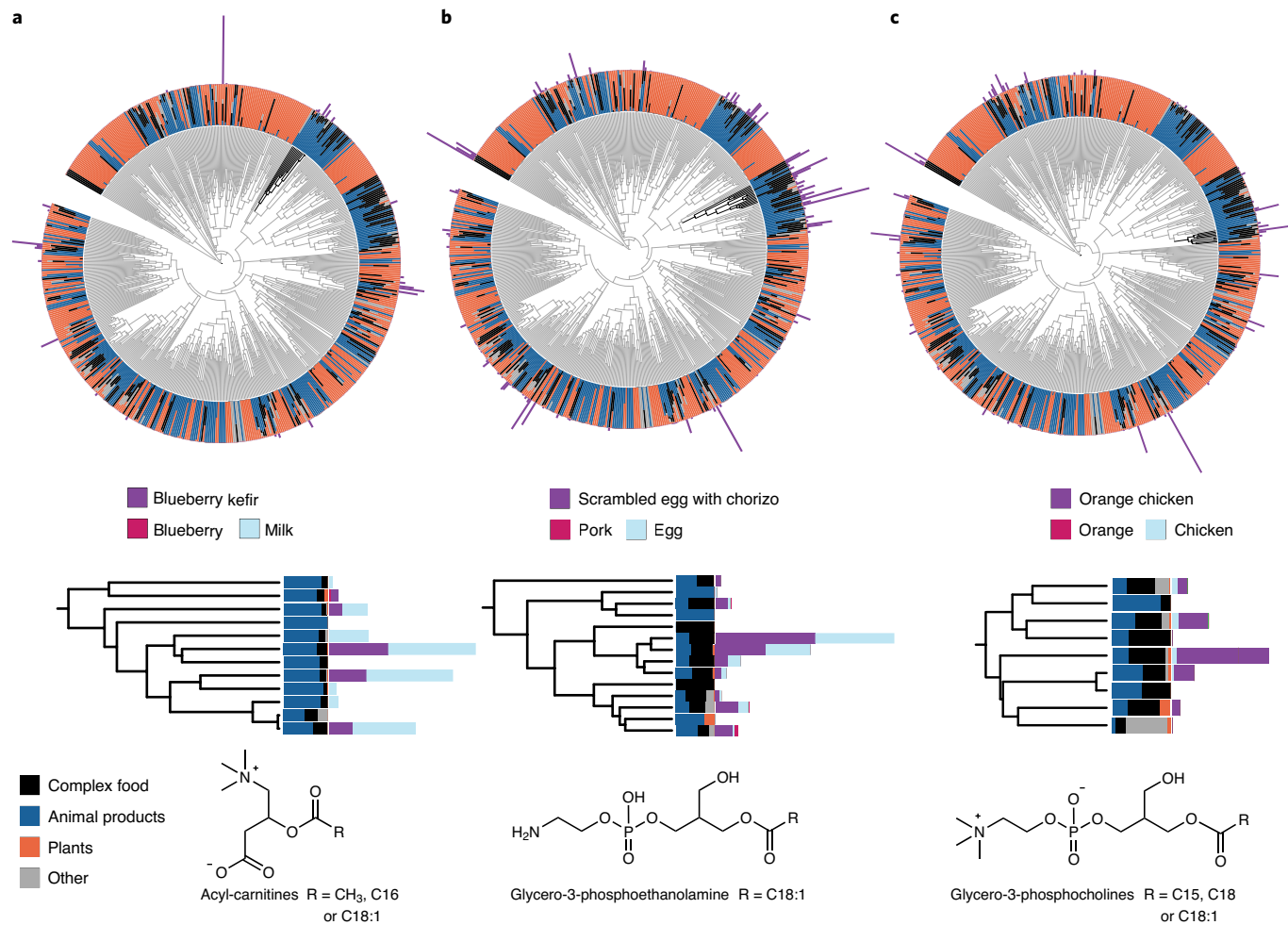
within foods are unannotated. We selected a diverse range of food ingredients to represent animal, plant and fungal groupings<sup>31</sup>. We first performed feature-based molecular networking using MZmine<sup>17,18</sup> to obtain spectral library matches for a subset of the chemical features (~20% annotated with cosine cutoff  $>0.7$ ). Using Qemistree, we collated GNPS spectral library matches and in silico predictions from CSI:FingerID to annotate ~91% of the chemical fingerprints (total 663 after quality filtering; Supplementary Table 1) with molecular structures. We also retrieved chemical taxonomy assignments for structures that were classified by ClassyFire<sup>23</sup> (~92% of all structures at the time of submission); the remaining were in the queue to be processed on the ClassyFire server for taxonomy assignment upon submission of the paper (see Methods). Labeling annotations allowed us to retrieve subtrees of distinct chemical classes (Fig. 3a) such as flavonoids, alkaloids, phospholipids, acyl-carnitines and O-glycosyl compounds in food products. We propagated ClassyFire annotations of chemical features (tree tips) to each internal node of the tree and labeled the nodes by pie charts depicting the distribution in chemical superclasses (Extended Data Fig. 7) and classes (Extended Data Fig. 8) of its tips. The molecular fingerprint-based hierarchy of chemical features agreed well with ClassyFire taxonomy assignment, further demonstrating that molecular fingerprints can meaningfully capture structural relationships among molecules in a hierarchical manner. Furthermore,

Qemistree coupled the chemical tree to sample metadata, revealing distinct chemical classes expected for each sample type. Branches representing acyl-carnitines were exclusively found in animal products (Fig. 3a). In contrast, honey, although categorized as an animal product, shared most of its chemical space with plant products, reflective of the plant nectar and pollen-based diet of honey bees. We observed a clade of flavonoids in both plant products and honey (Fig. 3 and Extended Data Fig. 8), but no other animal-based foods.

While it is expected that a complex food such as blueberry kefir contains molecules from blueberries, dairy, bacteria, and yeast we can now visualize how individual ingredients and food preparation contribute to the chemical composition of complex foods. We noted that metabolite signatures that stem directly from particular ingredients, such as phosphoethanolamine from eggs, are present in scrambled egg (Fig. 4b), but not in the other two foods highlighted (Fig. 4a,c). We can also observe the addition of ingredients in foods that were not listed as present in the initial set of ingredients. We were able to retrieve that there is black pepper in the scrambled egg with chorizo and orange chicken, but that this signal is absent from the blueberry kefir (Extended Data Fig. 9).

## Discussion

We show that our tree-based approach coherently captures chemical ontologies and relationships among molecules and samples



**Fig. 4 | A hierarchy of the compounds observed in simple foods and seven complex samples.** **a–c**, Hierarchies of the compounds observed in simple foods and seven complex samples: two meals of orange chicken, a cooked cucumber and the sauce from a meal (schmorgurken), sour cream, blueberry kefir and scrambled eggs with chorizo ( $N=126$  samples). Top, the inner rings show the relative abundance of each compound across simple animal products, plant products, fungi and algae (other) and the seven complex foods (black). In the outer rings, the absolute abundances of compounds in blueberry kefir (**a**), scrambled eggs with chorizo (**b**) and orange chicken (**c**) are overlaid on the trees to illustrate the shared and unique chemistry of complex foods. Below, compound subtrees for representative compounds from each meal are highlighted. Note that untargeted MS is blind to stereochemistry and oftentimes regiochemistry (for example, double bonds in a fatty acid); the structures shown are based on the spectral annotation of the reference library. This is equal to level 2 or 3 according to the 2007 metabolomics standards initiative<sup>40</sup>.

in various publicly available datasets. Qemistree depends on representing chemical features as molecular fingerprints, and does share limitations with the underlying fingerprint prediction tool CSI:FingerID. For example, fingerprint prediction depends on the quality and coverage of MS/MS spectral databases available for training the predictive models, and these will improve as databases are enriched with more compound classes. Nevertheless, the use of CSI:FingerID-predicted molecular fingerprints is highly advantageous. While annotations from spectral matches may be more accurate, their coverage is too low to adequately summarize the chemical content of complex samples. Qemistree is also applicable in negative ionization mode; however, fewer molecular fingerprints can be confidently predicted due to fewer publicly available reference spectra, resulting in less-extensive trees.

A key contribution of this work is to introduce the concept of building chemical hierarchies that can be used to leverage phylogeny-based tools (which have been highly advantageous for DNA sequencing analysis), for metabolomics data exploration. Hierarchical relationships have provided a powerful framework to understand the relatedness of organisms. These techniques form a

cornerstone for the interpretation of genomics data with phylogenetics and phylogenomics, and even taxonomy. The suite of tools and algorithms that have been developed over the past few decades in these fields, which use hierarchical structures, potentially have general relevance to the investigation of MS data. Using Qemistree we can begin to explore the applicability of other methods, such as Faith's Phylogenetic Diversity<sup>7</sup> to understand within-sample complexity, or phylogenetic-independent contrasts<sup>32</sup> with a metabolomics-inspired topology as these representations enter normal use.

We showed that a hierarchical representation could be used to infer chemically informed relationships between samples (Fig. 2). While we used molecular fingerprints predicted by CSI:FingerID to build chemical hierarchies here, this approach can be extended to incorporate other strategies to compare molecules for building chemical trees. For example, chemical relationships based on assigned chemical classes<sup>33</sup>, spectral motifs<sup>37</sup>, shared biosynthetic origin<sup>33</sup> or other structural comparison methods<sup>34</sup> could also be used as a basis for such a tree. These approaches will result in different tree topologies capturing complementary chemical information

for subsequent analyses. Ultimately, a broader benchmarking effort would be needed to understand when each approach should be used, similar to benchmarking efforts in the environmental DNA sequencing community<sup>35</sup>.

In addition to providing a framework for chemically informed sample comparisons within a dataset, Qemistree also provides a framework for comparing independently processed datasets. In the Qemistree workflow, we represent chemical features as their molecular fingerprints; this representation is largely independent of the technical variation such as chromatography shifts across MS experiments. Therefore, the chemical content of samples from different experiments can be compared by using a fingerprint-based representation without the need to repeat feature detection and feature alignment. This workflow is similar to how large-scale sample comparisons are made possible in sequence-based analyses<sup>36</sup>, where datasets are processed upfront, and rapidly coanalyzed according to the users' requirements. Extending these applications to MS data would allow metabolomics investigations of the scale of the Earth Microbiome Project<sup>31</sup> and the American Gut Project<sup>37</sup> to find global biochemical patterns. However, there is a need to benchmark experimental protocol comparability, as well as establish community-adopted standards that facilitate the global reuse of data. While these problems are substantial, we have seen examples of communities coming together to solve these issues for systematic and global data comparability<sup>31,38,39</sup>.

In summary, we introduce a new tree-based approach for computing and representing chemical features detected in tandem MS-based untargeted metabolomics studies. A hierarchy enables us to leverage existing tree-based tools, and can be augmented with structural and environmental annotations, greatly facilitating analysis and interpretation. We anticipate that Qemistree, as a data organization and comparison strategy, will be broadly applicable across fields that perform global chemical analysis, from medicine to environmental microbiology to food science and well beyond the examples shown here.

## Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41589-020-00677-3>.

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## Methods

**Qemistree algorithm.** The Qemistree workflow uses MS1-based feature tables and MS1, MS2 fragment ion information (MGF file format) as inputs (Extended Data Fig. 1). These inputs can be generated by processing untargeted MS data using MZmine<sup>17</sup> following the feature-based molecular networking method<sup>18</sup> (an example batch file that can be used to perform feature detection and generate the inputs for Qemistree can be found at the accession number [MSV000085226](#)). The files exported from MZmine with the Export/Submit to GNPS and SIRIUS Export module, and are then imported into QIIME2 (ref. <sup>25</sup>) as the following semantic types: FeatureTable(Frequency) (for the feature table) and MassSpectrometryFeatures (for the ion information).

### #PREPROCESSING:

- Use mzXML files from the instrument
- Perform feature detection using MZmine2
- Export sirius MGF and feature table (row m/z, row ID, feature area under the curve per sample)
- Convert the feature table to FeatureTable[Frequency] for QIIME2
- Create a FeatureData[Molecules] file for QIIME2 using 'row ID' and 'row m/z'
- Import the MGF file as MassSpectrometryFeatures for QIIME2

We use SIRIUS (v.4.0.1), ZODIAC<sup>41</sup> and CSI:FingerID to predict molecular substructures within MS features in the MGF files imported as MassSpectrometryFeatures. SIRIUS computes fragmentation trees for each molecular formula candidate of a feature (using the PubChem database by default) and ranks these by score. SIRIUS uses MS1 spectrum in the MGF file to determine the candidate ion adduct(s) to be used for the fragmentation tree computation of each feature. ZODIAC takes the top SIRIUS candidates as input and reranks molecular formula candidates considering reciprocal compound similarities in the dataset to increase correct molecular formula assignments. Subsequently, CSI:FingerID predicts molecular fingerprints for each feature based on the molecular formula with the highest ZODIAC score.

Note that all spectra provided to the Qemistree pipeline do not necessarily produce a fingerprint. SIRIUS does not compute fragmentation trees for multiply charged compounds and CSI:FingerID does not predict molecular fingerprints from spectra with fewer than three explained peaks. To ensure that high confidence molecular formulas are used in Qemistree, we only consider small molecules ( $m/z < 600$  Da) with a ZODIAC score above 0.98 (ref. <sup>41</sup>).

### #SUBSTRUCTURE PREDICTION:

For each feature with MS2 spectra in the MGF file:

- Compute fragmentation trees (using SIRIUS)
- Re-rank molecular formula candidates on the complete dataset (using ZODIAC)
- Predict fingerprints based on best molecular formula assignment (using CSI:FingerID)

A dataset  $M$  (that is, a set of exports from MZmine) is a matrix of size  $n$  rows by  $l$  columns. Each row represents a molecule ( $m_1, m_2, \dots, m_n$ ), and each column represents a molecular substructure feature. As such, each molecule  $m_i$  is composed of a vector (with length  $l$ ) of predicted probability values (one for each SIRIUS-generated molecular substructure). We remove from our analyses the features without a corresponding vector  $m_i$ . In our tests, we have observed that for each dataset 10–15% of the input features are discarded.

For indexing purposes, we relabel each molecule  $m_i$  with the MD5-checksum of the predicted fingerprint vector. The motivation to apply the MD5 hashing function is to assign a unique identifier to each feature, which is particularly useful when comparing datasets independently processed using MZmine. If two distinct molecules ( $i, j$ ) have identical checksums, that is  $md5(m_i) = md5(m_j)$ , then we aggregate those two vectors such that all rows in  $M$  are unique. This operation is also propagated down to the table of molecular intensities, in that context intensities are added together.

To coanalyze multiple datasets  $M_1, M_2, \dots, M_k$ , we combine the matrices into a new dataset  $M'$ . For any two repeated molecules  $m_i$  and  $m_j$  in  $M'$  we merge their intensities and values as described before. Last, we create a hierarchy of chemical relationships  $T$  using a distance matrix  $D$  measuring the distance between all pairs of molecules in  $M'$ . For qualitative substructure comparisons, we use the Jaccard distance metric and a threshold of 0.5. Otherwise, we use the Euclidean distance with the original probability vectors. By default, our implementation relies on the Euclidean distance so that a threshold value is not needed. In practice, we noted different metrics at this stage have only small effects on the downstream analyses. With  $D$ , we cluster the molecules in a hierarchical fashion using the unweighted pair group method with arithmetic mean. The tips in the resulting tree  $T$  have a one-to-one correspondence with all the molecules  $m_i$  in  $M'$ .

### #HIERARCHY CREATION (meta-analysis)

For each fingerprint, feature table in DATASETS:

- Collate fingerprints into a matrix of features by fingerprints
- Match the tuple to have the exact same features and same order
- Merge all the fingerprints and feature tables

- (use MD5 hash of fingerprint vectors to merge identical fingerprints)

Compute a distance matrix between fingerprints

- If the probability vectors are binarized use a qualitative metric (Jaccard) otherwise use a quantitative metric (Euclidean)
- Build a hierarchical tree based on the distance matrix

Qemistree analysis can be performed either through a command-line interface using q2-qemistree qiime2 plugin (<https://github.com/biocore/q2-qemistree>) or as a web-based workflow on GNPS (<https://ccms-ucsd.github.io/GNPSDocumentation/qemistree/>). We have created a dashboard at <https://qemistree.ucsd.edu> for GNPS users to interactively explore Qemistree tree visualization. It requires the Qemistree task ID to import Qemistree results from GNPS, and allows users to modify the chemical tree visualization by changing parameters such as filtering features based on ClassyFire taxonomy level, label of the tips and sample metadata column for plotting abundance bar plots. We provide step-by-step instructions on how to use this dashboard at <https://ccms-ucsd.github.io/GNPSDocumentation/qemistree/>.

We note that molecular similarity profiling, as represented here, may underemphasize the large biological effects of small differences among molecules (for instance, a methyl group can have a large impact on the activity of a drug, but will have a small impact on the Qemistree profile). Whether to emphasize or attenuate small differences among related features is an ongoing discussion in other related fields, such as DNA sequencing, and the best approach depends on application<sup>42</sup>.

Qemistree leverages CSI:FingerID to increase chemical annotations in MS data (Extended Data Table 2). CSI:FingerID has been shown to outperform all other in silico methods for molecular formula identification in blind critical assessment of small molecule identification contests<sup>32,43</sup>. Representing molecules as CSI:FingerID fingerprints allows us to query rich structural databases (for example, >100 million compounds in PubChem) instead of spectral libraries that are sparser (~160,000 reference spectra only covering tens of thousands of compounds).

Using Qemistree, we collate GNPS spectral library matches and in silico predictions from CSI:FingerID and run ClassyFire<sup>25</sup> to assign a five-level chemical taxonomy (kingdom, superclass, class, subclass and direct parent) to all molecules annotated via spectral library matching and in silico prediction (Extended Data Table 3).

Note that we have developed the infrastructure such that when users first run ClassyFire through Qemistree, they get taxonomic assignments for all the structures that have previously been classified by ClassyFire and are retrievable by InchiKey through a GNPS API service (<https://ccms-ucsd.github.io/GNPSDocumentation/api/>). The remaining structures are queued on the ClassyFire server for automatic and continuous taxonomy assignment. We provide users with a table of structures that were unclassified at the time of query; this can be used to retrieve additional taxonomic assignments using the Qemistree module get-classifyfire-taxonomy downstream of the initial query (<https://github.com/biocore/q2-qemistree>). As more and more classifications are recorded on GNPS, the users can retrieve more taxonomic assignments using Qemistree.

**Evaluation dataset.** *Sample preparation and extraction.* Four samples were used in the gradient benchmarking dataset: (1) the 'serum' sample consists of the NIST SRM 1950 reference sample made of human serum spiked with compounds<sup>44</sup>, (2) Two human fecal samples from the American Gut Project<sup>37</sup> obtained from a single male individual with a 35-d interval (Sample fecal-1, 10 November 2013, and fecal-2, 14 December 2013) and (3) the 'tomato' seedling sample (*Solanum lycopersicum* plant) was prepared using 3-weeks post-germination specimens (fresh whole seedlings were used). Note that the participant had stool samples collected by consent under the Human Research Protection Program (HRPP) 150275 protocol (Evaluating the Human Microbiome). The protocol was approved by the HRPP of the University of California, San Diego. Written informed consent obtained from the patient concerning dissemination and scientific publication of the results is also included in the approved protocols. The NIST SRM 1950 sample (1 ml), two fecal samples (210 mg of fresh material each) and the tomato seedlings (800 mg of fresh material) were dissolved in 1 ml of 7:3 methanol:water in a 1-ml polypropylene round-bottom tube (QIAGEN) and homogenized in a tissue lyser (Tissue Lyser II, Qiagen) at 25 Hz for 5 min. The tubes were then centrifuged at 15,000 r.p.m. for 15 min, and 600  $\mu$ l of the supernatant was collected and loaded on solid-phase extraction cartridges (Oasis HLB, Waters) made of hydrophilic-lipophilic balance stationary phase (30 mg and 30  $\mu$ m particle size), that were first activated with 100% methanol and 100% water (1 ml each). After loading the supernatants on the cartridges, washing elution was carried out with 95:5 methanol:water (1 ml), and the samples were eluted with 7:3 methanol:water (2 ml), followed by 100% methanol (1 ml). The samples were dried down with a vacuum concentrator (CentriVap, Labconco) and resuspended in 2.5 ml of 7:3 methanol:water containing 0.5  $\mu$ M of amitriptyline as an internal standard. Samples were prepared by mixing the four different samples in various proportions. The resulting extracts were analyzed by MS along with binary, and quaternary mixtures of these samples in different proportions (Extended Data Table 3). For example, the serum and tomato samples were mixed in the following ratios: 100:0, 75:25, 50:50, 25:75 and 0:100.

**LC-MS experiments.** Samples were analyzed using ultra high-performance liquid chromatography (Vanquish, Thermo Scientific) coupled to a quadrupole-Orbitrap mass spectrometer (Q Exactive, Thermo Scientific). The quadrupole-Orbitrap

mass spectrometer (Q Exactive, Thermo Scientific) was fitted with an electrospray source (HESI-II) operating in positive ionization mode. The source used the following parameters: spray voltage, +3,500 V; heater temperature, 437.5 °C; capillary temperature, 268.75 °C; S-lens RF, 50 arbitrary units (a.u.); sheath gas flow rate, 52.5 a.u. and auxiliary gas flow rate, 13.75 a.u. The samples were acquired in nontargeted MS2 acquisition mode, with up to four MS2 scans of the most abundant ions per MS1 scan. The spectra were recorded from 0.48 to 17 min. The following parameters were used for full MS scan: resolution (35,000), Automatic Gain Control target ( $1.0 \times 10^6$ ), maximum injection time (125 ms) and scan range (150–1,500  $m/z$ ). For the data-dependent in MS2, the following parameters were used: resolution (17,500), AGC target ( $2.5 \times 10^6$ ), maximum injection time (125 ms), loop count (4), isolation window (1.5  $m/z$ ) fixed first mass (70  $m/z$ ) (70–1,500  $m/z$ ) and up to four MS/MS scans of the most abundant ions per duty cycle. Higher-energy collision induced dissociation was performed with a normalized collision energy of 30 (20, 35, 50). The data-dependent settings were set as follows: minimum AGC ( $1.25 \times 10^4$  (intensity threshold  $1.0 \times 10^5$ )), apex trigger 3 to 15 s, charge exclusion 3–8 and >8, exclude isotopes (on), dynamic exclusion (14.0 s).

Two different chromatographic conditions were used for the mass spectrometer (named C18, C18-RTshift). In each case, a Phenomenex Kinetex C18 1.7- $\mu$ m column (100 A)  $100 \times 2.1$  was used. The column was equipped with a C18 guard cartridge (Phenomenex). The mobile phases consisted of A (100% water + 0.1% formic acid) and B (100% acetonitrile + 0.1% formic acid), and the flow rate was set to 500  $\mu$ l min $^{-1}$  throughout the experiment, and the column maintained at 40 °C. The chromatographic elution method was set as follows. For the C18: 0–0.25 min, 20% B; 0.25–4 min, 50% B; 4–15 min, 100% B; 15–15.90 min, 100% B; 16–18 min, 20% B. For the C18-RTshift: 0–0.25 min, 20% B; 0.25–4 min, 50% B; 4–13 min, 100% B; 15–15.90 min, 100% B and 16–18 min, 20% B. Each sample was analyzed in triplicate, and the injection sequence was randomized. A 'QC mix' made of the four samples was used to optimize the experiment parameters and injected them periodically throughout the sequence. No carry over was observed. Successful injections had a relative standard deviation of no more than 15% for replicates and QC mix samples, and the retention time deviation for the internal standards (amitriptyline  $m/z$  278.190 and 3.57 min) was observed below 1 s for replicates and QC (quality control) mix, and not more than 2–3 s for replicates and QC mix samples (see feature  $m/z$  485.366 at 11.0 min). For most ions shifts of 1–2 min are observed. The difference between LC–MS/MS profiles for a pooled sample analyzed in the chromatographic conditions C18 and C18-RTshift are presented as 2D maps in Extended Data Figs. 2 and 3.

**MS data processing.** Thermo MS (.RAW) were converted to  $m/z$  extensible markup language (mzML)<sup>45</sup> in centroid mode using MSConvert ProteoWizard<sup>46</sup> (release 201812). The mzML files were processed with MZmine toolbox<sup>17</sup> (v.2.38) on Ubuntu 18.04 LTS 64-bits workstation (intel Xeon 5E-2637, 3.5 GHz, eight cores, 64 GB of RAM) following the feature-based molecular networking method<sup>48</sup>.

**Global FoodOmics dataset. Sample preparation and extraction.** Samples were collected, extracted and MS data were acquired as a part of the Global FoodOmics project according to the sampling and data acquisition protocols described in Gauglitz et al.<sup>47</sup> Briefly, 126 food samples were selected from the Global FoodOmics dataset. One hundred and nineteen simple food samples (simple in contrast to complex and defined as a single-ingredient food) were selected to cover a broad spectrum of fruits, vegetables, meat and fungi. Each food was represented in at least triplicate in the data subset. Additionally, seven complex samples were selected that contained simple foods from the simple food subset in their ingredient lists. The complex foods were from two separate meals of orange chicken, a cooked cucumber and the sauce from a meal (schmorgurken; in a tomato and sour cream sauce), sour cream, blueberry kefir and scrambled egg with chorizo. Sample metadata describes the food samples based on a food hierarchy beginning with plant versus animal versus fungus (sample\_type\_group1) and increasing in detail down to persian cucumber versus cherry tomato and so on (sample\_type\_group6).

Briefly, samples were extracted in 95% LC–MS grade ethanol; 5% LC–MS grade water. Samples were analyzed using the same LC–MS/MS setup and software as described above for the maXis II QTOF mass spectrometer (Bruker Daltonics), using a Phenomenex Kinetex C18 1.7  $\mu$ m (100 A)  $100 \times 2.1$  column equipped with a guard cartridge (Phenomenex). The instrument tuning and internal calibrant remained the same as described above. MS spectra were acquired in a positive ion mode in the range  $m/z$  50–1,500. The mobile phases consisted of A (100% water + 0.1% formic acid) and B (100% acetonitrile + 0.1% formic acid), and the flow rate was set to 0.5  $\mu$ l min $^{-1}$  throughout the experiment and the column maintained at 40 °C.

**MS data processing.** The MS data (.d) were converted to .mzXML with lock mass calibration applied using CompassXport batch mode in Data Analysis v.4.4 software (Bruker Daltonics) running on a Windows 10 PC. The MS data was processed with MZmine toolbox<sup>17</sup> (v.2.38) using the parameters outlined in an XML batch file (see Data availability).

**Multivariate comparisons.** To evaluate the benefits of using a tree for multivariate analysis, we generated pairwise sample distances using Bray–Curtis<sup>29</sup> (agnostic of chemical relationships) and Weighted UniFrac<sup>9</sup> (chemical relationship

tree-informed). Both of these metrics compare samples quantitatively; that is, using the abundances of each feature. Notably, UniFrac weights the distances based on the shared branches of the tree used for computation. The distances within- and between-sample groupings were compared using a one-sided permutational ANOVA test.

**Comparison to cosine-score-based clustering.** We compared the clustering of samples using Weighted UniFrac on molecular fingerprint-based hierarchy to Bray–Curtis metric (which does not account for chemical relationships) and two MS/MS cosine similarity informed methods: CSCS distance metric<sup>38</sup> and Weighted UniFrac on MS/MS cosine-score-based hierarchy. We include a direct comparison of the three approaches in performing chemically meaningful clustering of samples in the Global FoodOmics dataset ( $N = 126$ ; Extended Data Table 4). Food ontology level 1 corresponds to animal, plant and fungal samples in Earth Microbiome Project Ontology<sup>31</sup> and levels 2 to 4 represent progressively more detailed food categories. We note that both cosine-based and fingerprint-based pipelines cluster sample groups reasonably well, with molecular fingerprint-based hierarchy leading to improved sample clustering in this dataset.

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

## Data availability

The MS data, metadata and methods for the evaluation dataset have been deposited on the GNPS/MassIVE public repository<sup>2,32</sup> under the accession number **MSV000083306**. Source data for the figures are available as Supplementary Datasets 2–5. The parameters used for molecular networking are available on GNPS at <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=fda476c72724b29a91693a108fa5a9d>. The chemical hierarchy generated by Qemistree (v.2020.1.2) is available on iTOL<sup>24</sup> at <https://itol.embl.de/tree/709513416494381587432576>. The MS data, metadata and methods for Global FoodOmics dataset have been deposited on the GNPS/MassIVE public repository<sup>2,32</sup> under the accession number **MSV000085226**. The parameters used for molecular networking are available on GNPS at <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=ceb2a199d6b4f4fb0f8490d9c96d631>. The chemical hierarchy generated by Qemistree (v.2020.1.2) is available on iTOL<sup>24</sup> at <https://itol.embl.de/tree/13711034118313741584046018>. The MS data, metadata and methods for Cheetah fecal dataset have been deposited on the GNPS/MassIVE public repository<sup>2,32</sup> under the accession number **MSV000082969**. The parameters used for molecular networking are available on GNPS at <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=093798dfde2448239410c3d465ef9fea>.

## Code availability

All source code is publicly available under BSD-2-Clause on GitHub at <https://github.com/biocore/q2-qemistree>. Qemistree is also available as an advanced analysis workflow on GNPS at <https://ccms-ucsd.github.io/GNPSDocumentation/qemistree/>. All analyses are documented in Jupyter Notebooks available at <https://github.com/knightlab-analyses/qemistree-analyses>.

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### Author contributions

A.T. and P.C.D. conceived the concept and managed the project. A.T. and Y.V.B. developed the algorithm and wrote the code for Qemistree. A.T. and Y.V.B. contributed equally to the work. L.F.N., R.K. and P.C.D. supervised method implementation. K.D., M.W., J.J.v.d.H., M.E., D.M. and A.G. tested and provided suggestions on how to improve the method. M.W. managed the deployment of Qemistree on GNPS. A.T. and M.W. developed the GNPS-Qemistree Dashboard. D.A. and A.T. wrote the documentation for the GNPS-Qemistree workflow. Y.V.B., Q.Z. and A.T. developed Qemistree-iTOL visualization. L.F.N. and M.N.E. performed the MS for the evaluation dataset. A.T., Y.V.B. and L.F.N. analyzed and interpreted the evaluation data. J.M.G. performed MS of the Global FoodOmics samples. A.T. and J.M.G. analyzed and interpreted the Global FoodOmics data. A.D.B. made the comparisons to CSCS. K.D., M.F., M.L. and S.B. supported the integration of SIRIUS, ZODIAC and CSI:FingerID.

A.T., Y.V.B., R.K. and P.C.D. wrote the manuscript. L.F.N., J.M.G., M.N.E., J.J.v.d.H., M.E., K.D., Q.Z., D.M., A.D.B., A.G., J.H., M.F., M.L. and S.B. improved the manuscript.

### Competing interests

M.W. is a founder of Ometa Laboratories LLC. P.C.D. is a scientific advisor for Sirenas, Cybele and Galileo PCD is also a scientific advisor and founder of Enveda and Ometa Laboratories LLC with approval by University of California San Diego. LLC. K.D., M.L., M.F. and S.B. are founders of Bright Giant GmbH.

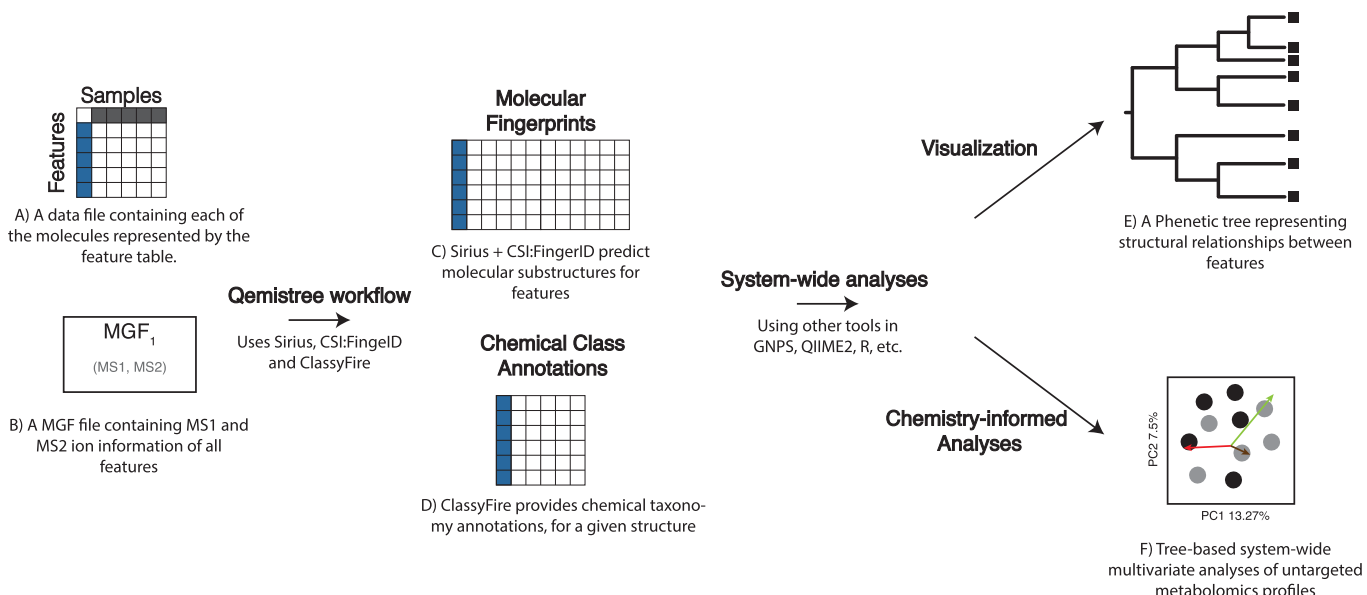
### Additional information

**Extended data** is available for this paper at <https://doi.org/10.1038/s41589-020-00677-3>.

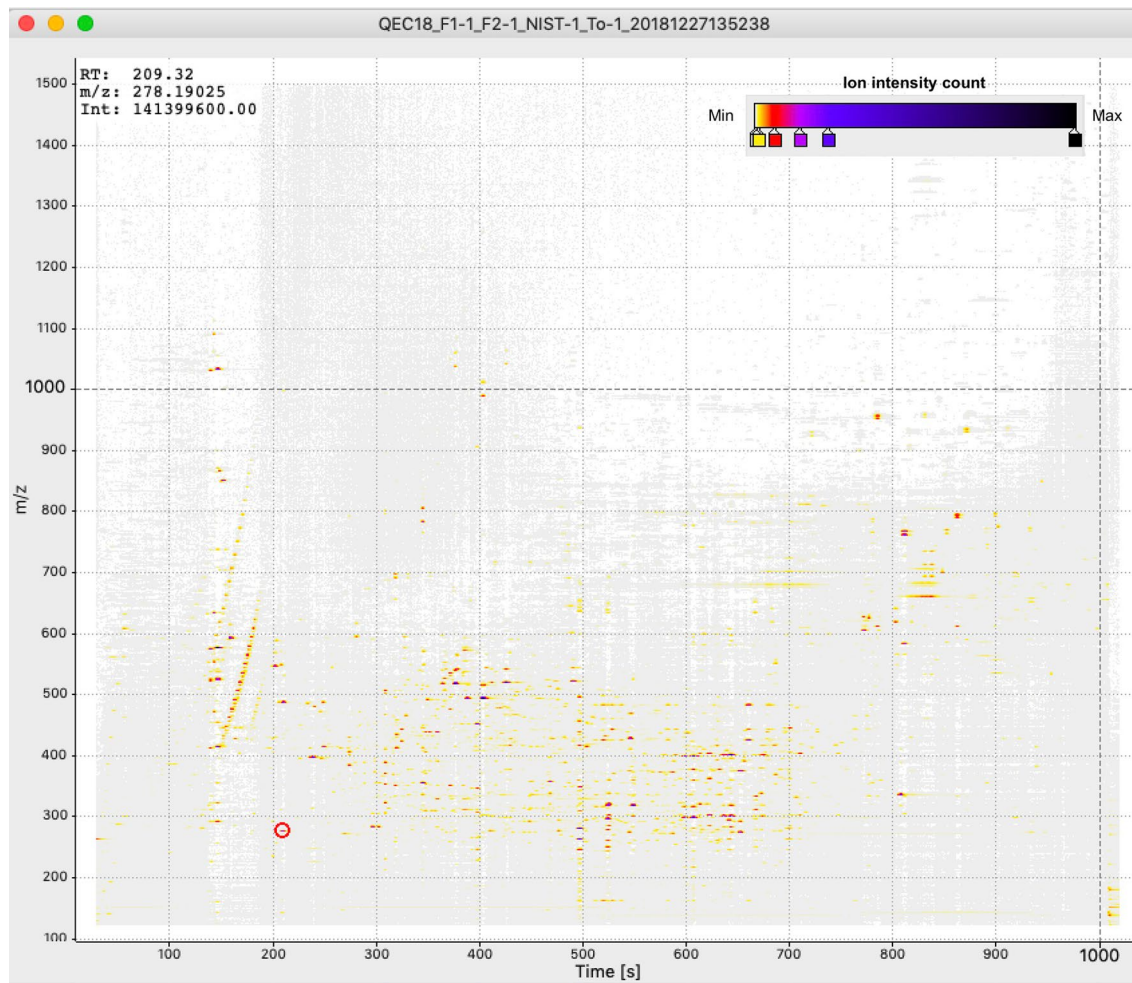
**Supplementary information** is available for this paper at <https://doi.org/10.1038/s41589-020-00677-3>.

**Correspondence and requests for materials** should be addressed to P.C.D.

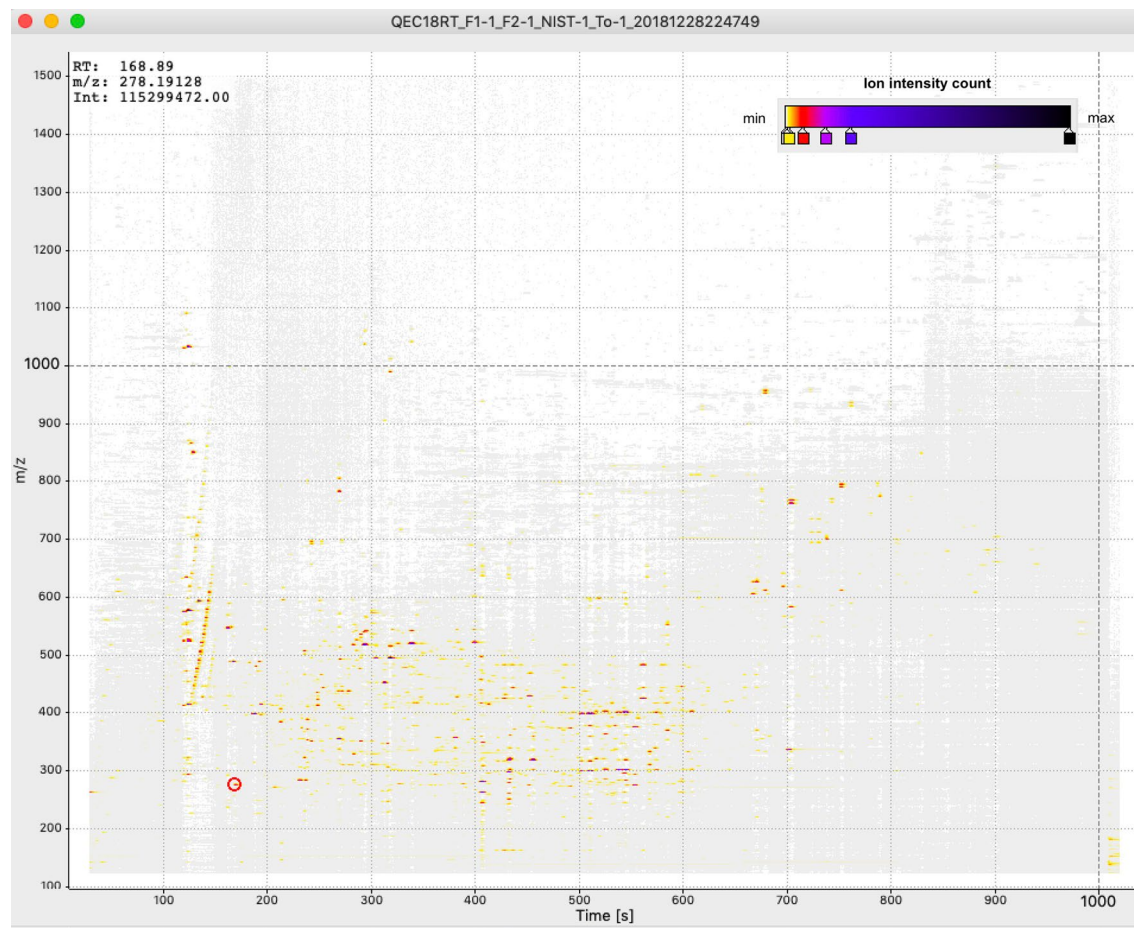
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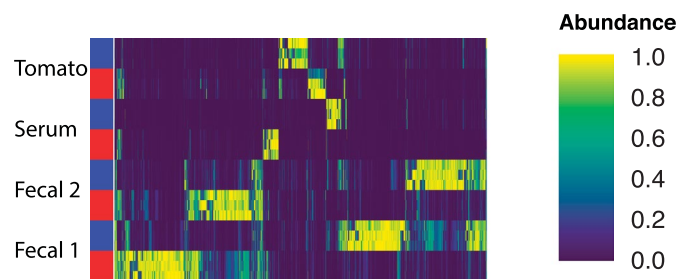
**Extended Data Fig. 1 | End-to-end Qemistree analysis using GNPS and QIIME2.** Qemistree analysis can be performed using two required input files: 1) A table of molecule (or chemical feature) abundances per sample and 2) an MGF file with MS1 and MS2 ion information. These inputs can be generated by processing mass spectrometry files (.mzXML) through MZmine for feature detection. In Qemistree, these input files are processed through SIRIUS and CSI:FingerID to generate molecular fingerprints and *in silico* structural annotations (SMILES) per MS feature. We use the predicted molecular fingerprints to generate a phenetic tree of relationships between MS features based on sub-structural similarity. This tree can be visualized in iTOL for further data exploration. If the user inputs a sample metadata file, they can also visualize the abundances of each MS feature stratified by sample grouping of interest. Additionally, the Qemistree queries ClassyFire to classify the structural annotations into chemical ‘kingdom’, ‘superclass’, ‘class’, ‘subclass’ and ‘direct parent’. We further allow the users to input a file with MS/MS spectral library matches (optional) into the workflow such that these library matches (typically, 2–20% of all MS features), instead of *in silico* annotation, are used for ClassyFire queries whenever available. All the outputs of the Qemistree workflow can be analyzed further using QIIME 2 tools (such as tree-based alpha and beta diversity, mmvec: <https://github.com/biocore/mmvec>, songbird: <https://github.com/biocore/songbird>) or explored in Python, R etc. as needed.



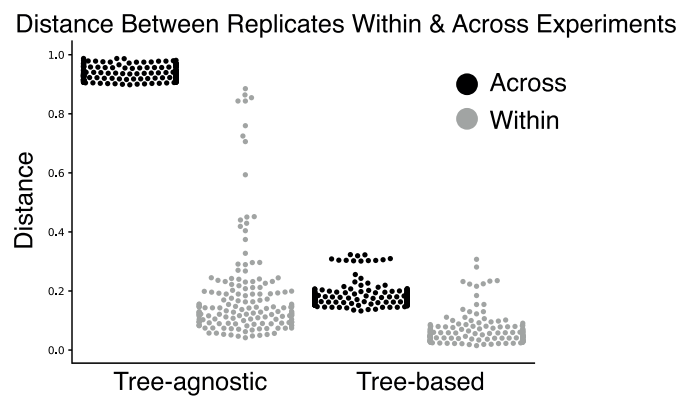
**Extended Data Fig. 2 |** 2D map of the LC-MS/MS data of the pooled sample for the C18 chromatographic conditions.



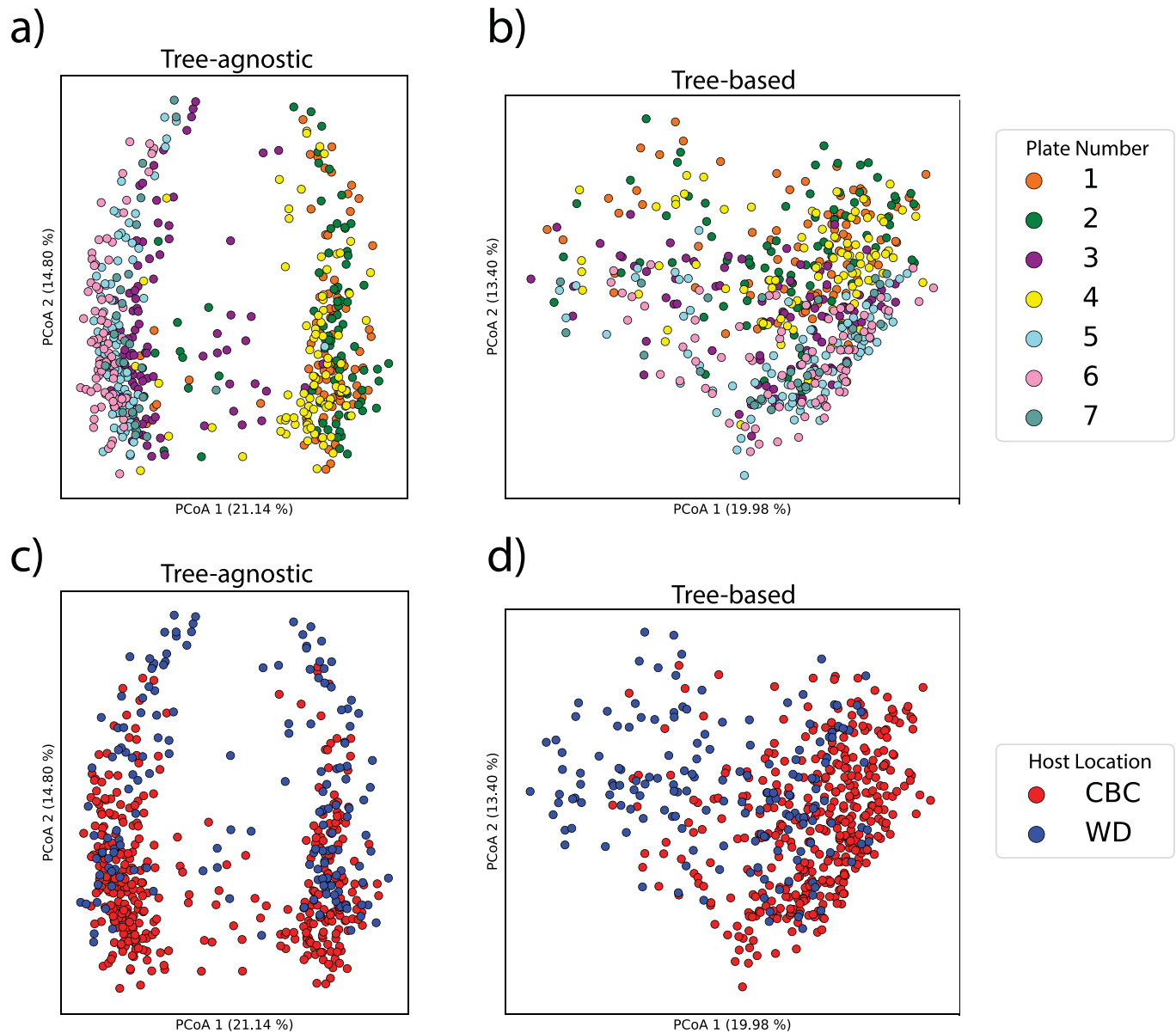
**Extended Data Fig. 3 |** 2D map of the LC-MS/MS data of the pooled sample for the C18-RTshift chromatographic conditions.



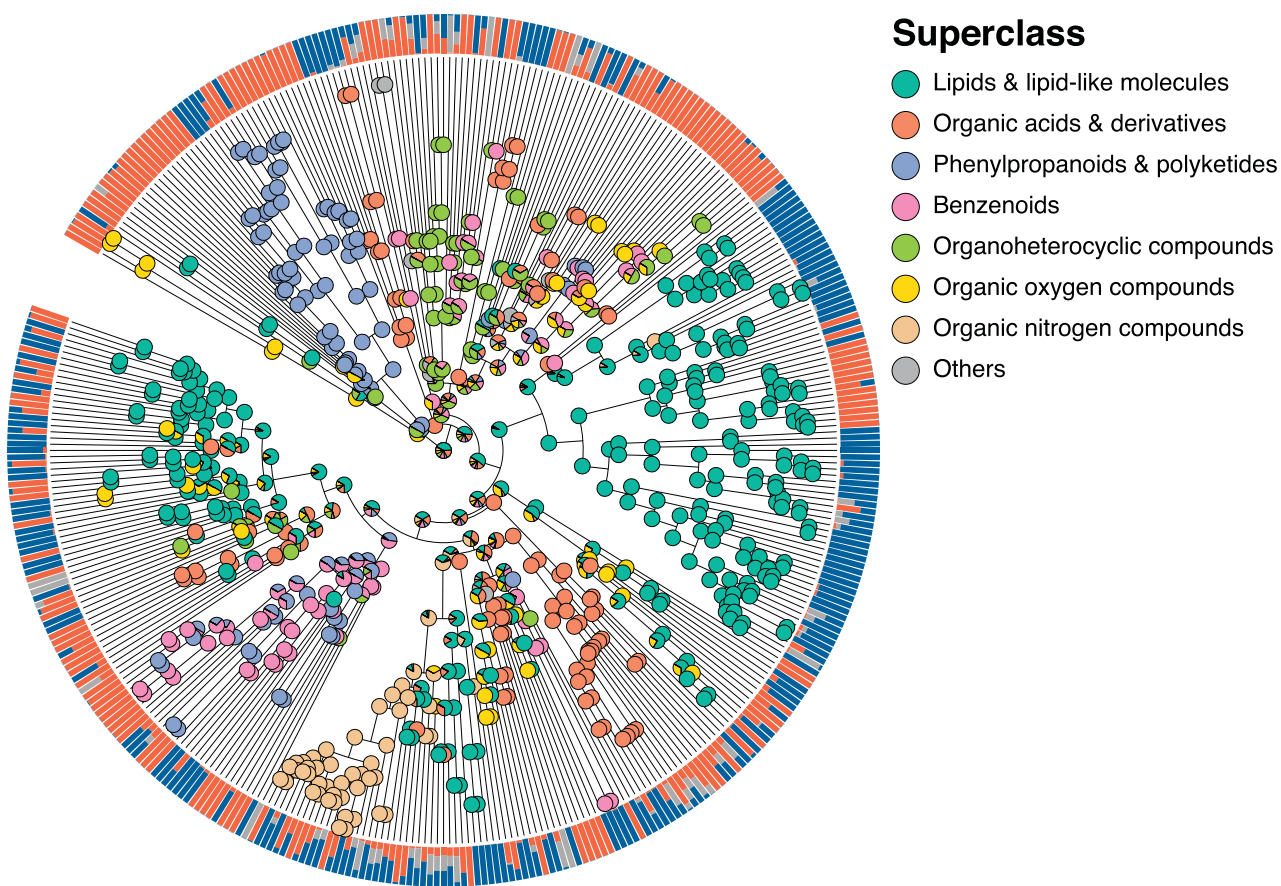
**Extended Data Fig. 4 | Technical variation in mass-spectrometry due to chromatographic shifts.** Sample (y-axis) by molecule (x-axis) heatmap of 2 fecal samples, tomato seedling samples, and serum samples in the evaluation dataset grouped by chromatography conditions.



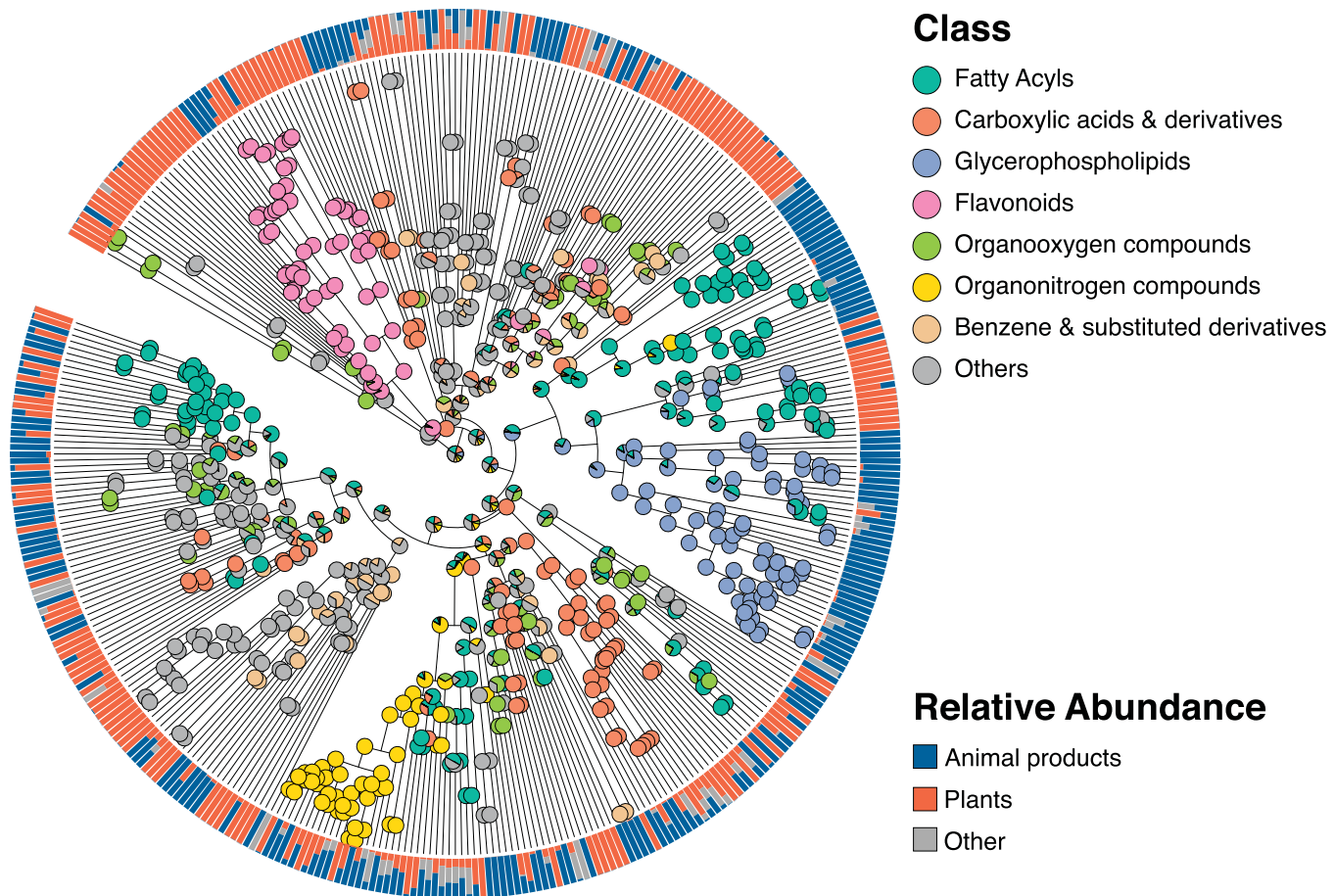
**Extended Data Fig. 5 | Qemistree reduces the differences between biological replicates across mass-spectrometry runs.** A comparison of distances between sample replicates within and across chromatography gradients when using tree-agnostic (Bray-Curtis) distances and tree-based (Weighted UniFrac) distances.



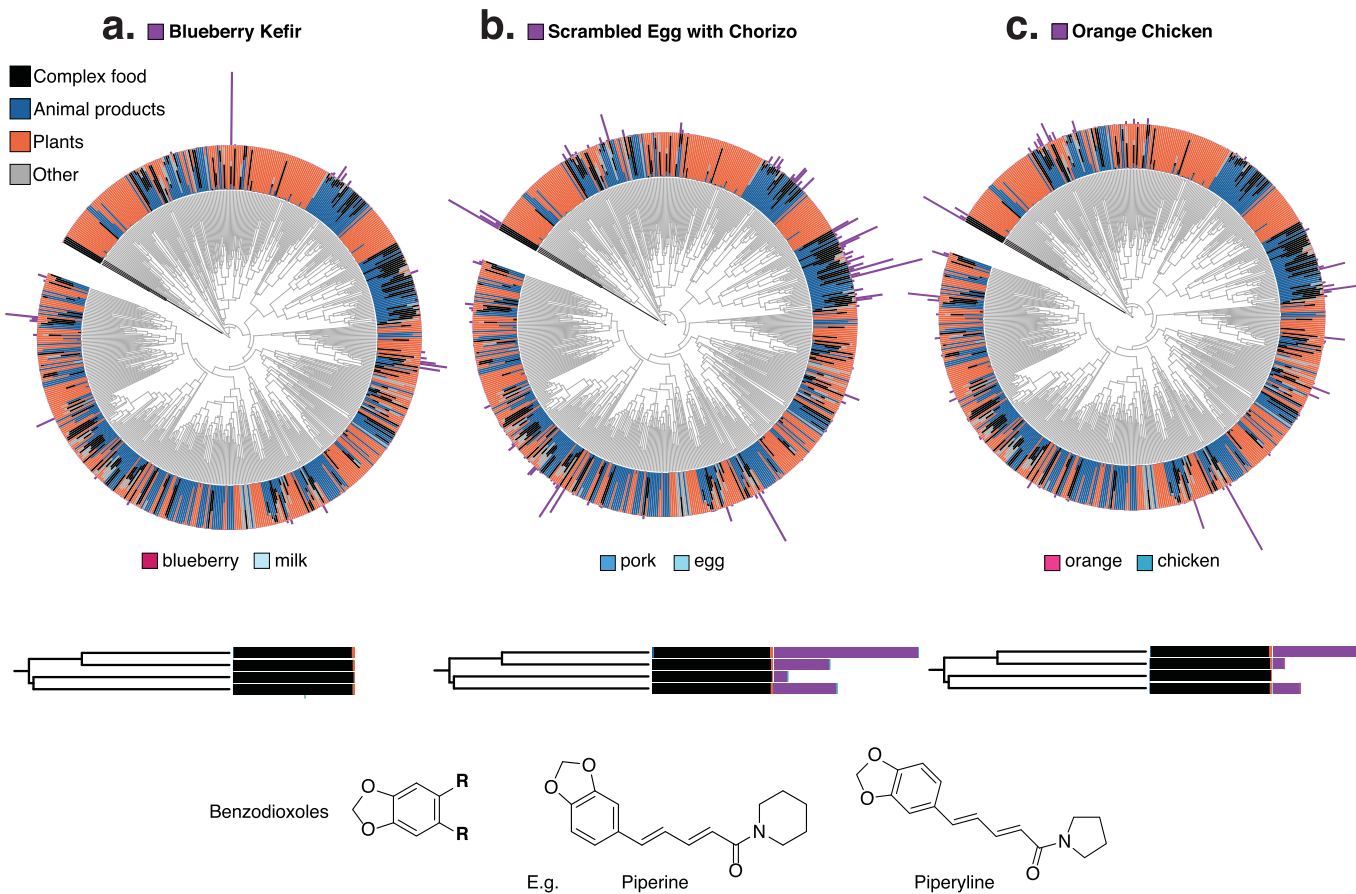
**Extended Data Fig. 6 | Qemistree mitigates plate-to-plate variation in fecal metabolomics study to highlight a biologically-relevant effect.** **a**) Principal coordinate analysis (PCoA) of tree-agnostic distances (Bray-Curtis) colored by plate number (pseudo- $F$ =32.39,  $p=0.001$ ). **b**) PCoA of tree-informed distances (Weighted UniFrac) colored by plate number (pseudo- $F$ =15.67,  $p=0.001$ ). The same PCoA of **(c)** Bray-Curtis distances (pseudo- $F$ =33.50,  $p=0.001$ ) and **(d)** Weighted UniFrac distances (pseudo- $F$ =48.42,  $p=0.001$ ) colored by cheetah location which governed the diet of cheetahs. CBC: Cheetah Breeding Center; WD: Wildlife Discoveries.



**Extended Data Fig. 7 | Chemical taxonomy of food-derived compounds at chemical superclass level.** Chemical hierarchy of compounds (tree tips) detected in simple food products (single ingredient foods, N=119). Internal nodes are labeled by pie charts of the superclass level taxonomy of children tips. Outer ring shows the relative abundance of each compound across simple animal products, plant products, and other (fungi and algae). The chemical hierarchy iTOL link: <https://itol.embl.de/tree/709513416412858158733337>.



**Extended Data Fig. 8 | Chemical taxonomy of food-derived compounds at chemical class level.** Chemical hierarchy of compounds (tree tips) detected in simple food products (single ingredient foods, N = 119). Internal nodes are labeled by pie charts of the class level taxonomy of children tips. Outer ring shows the relative abundance of each compound across simple animal products, plant products, and other (fungi and algae). The chemical hierarchy iTOL link: <https://itol.embl.de/tree/7095134164128581587333337>.



**Extended Data Fig. 9 | Chemical hierarchy of the compounds observed in simple foods and seven complex samples. a,b,c)** 2 meals of orange chicken, a cooked cucumber and the sauce from a meal (schmorgurken), sour cream, blueberry kefir, and egg scramble with chorizo ( $N=126$  samples). The inner ring shows the relative abundance of each compound across simple animal products, plant products, fungi and algae (other) and complex foods. The absolute abundances of compounds in blueberry kefir (a), scrambled eggs with chorizo (b), and orange chicken (c) (outer bars) are overlaid on the tree to illustrate the shared and unique chemistry of complex foods. We highlight a classifier subtree annotated as benzodioxoles, compounds found in black pepper (in black) that are almost exclusively detected in complex foods. Note that untargeted mass-spectrometry is blind to stereochemistry and oftentimes regiochemistry (for example double bonds in a fatty acid); the structures shown are based on the spectral annotation of the reference library.

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### Software and code

Policy information about [availability of computer code](#)

## Data collection

Evaluation dataset: Thermo mass spectrometry data (.RAW) were converted to m/z extensible markup language (mzML) in centroid mode using MSConvert ProteoWizard (release 201812).

Global Foodomics dataset: The mass spectrometry data (.d) were converted to .mzXML with lock mass calibration applied using CompassXport batch mode in Data Analysis 4.4 software (Bruker Daltonics, Bremen, Germany).

The version of Qemistree we used relies on Python 3.6, QIIME 2019.7 and R 3.5.1. A complete list of dependencies, and their versions can be found here:

<https://github.com/knightlab-analyses/qemistree-analyses/blob/master/environment.yml>

## Data analysis

Evaluation dataset: The mzML files were processed with MZmine toolbox 17 (version 2.38) on Ubuntu 18.04 LTS 64-bits workstation (intel Xeon 5E-2637, 3.5 GHz, 8 cores, 64 Go of RAM) following the Feature-Based Molecular Networking method.

Global Foodomics dataset: The mass spectrometry data was processed with MZmine toolbox 17 (version 2.38) on a Windows 10 PC using the parameters outlined in an XML batch file provided in Data availability section.

The code used to generate the figures can be found here:  
<https://github.com/knightlab-analyses/qemistree-analyses/>

The source code for Qemistree can be found here:  
<https://github.com/biocore/q2-qemistree>

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## Data

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[Metabolome evaluation dataset (accesion code - MSV000083306.)]

The mass spectrometry data, metadata, and methods for the evaluation dataset have been deposited on the GNPS/MassIVE public repository under the accession number MSV000083306. The parameters used for molecular networking are available on GNPS: <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=efda476c72724b29a91693a108fa5a9d>. The chemical hierarchy generated by Qemistree (version 2020.1.2) is available on iTOL: <https://itol.embl.de/tree/709513416494381587432576>.

[Global foodomics dataset (accession code - MSV000085226.)]

The mass spectrometry data, metadata, and methods for Global Foodomics dataset have been deposited on the GNPS/MassIVE public repository under the accession number MSV000085226. The parameters used for molecular networking are available on GNPS: <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=ceb28a199d6b4f4fb08490d9c96d631>. The chemical hierarchy generated by Qemistree (version 2020.1.2) is available on iTOL: <https://itol.embl.de/tree/13711034118313741584046018>.

[Cheetah fecal dataset (accession code - MSV000082969)]

The mass spectrometry data, metadata, and methods for Global Foodomics dataset have been deposited on the GNPS/MassIVE public repository under the accession number MSV000082969. The parameters used for molecular networking are available on GNPS: <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=093798dff2448239410c3d465ef9fea>. The chemical hierarchy generated by Qemistree (version 2020.1.2)

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## Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

### Sample size

Sample size calculation is not applicable to this manuscript. Qemistree is a data visualization tool, therefore we report all results as visual improvements in data representation.

For evaluation dataset, we used pure fecal, tomato and serum samples mixed in gradually increasing proportions and data were acquired on Q-Exactive. The number of samples was determined by the resulting combinations from binary, tertiary and quaternary mixtures (n=162). In order to account for within-experiment variation and to compare this to variation across chromatography conditions we collected mass spectrometry data in triplicates.

For Global FoodOmics data, samples were selected to represent animal, plant, and fungal groupings of the Earth Microbiome Project ontology such that each food was represented in at least triplicate in the data (n=126). This number of samples is able to maintain balanced classes in the dataset, as well as a broad representation of the food types collected in this study. For the purposes of this study this criteria was sufficient since it enabled us to visualize a diverse chemical space in a study size representative of typical biological experiments.

### Data exclusions

No data from the above mentioned sample set was excluded in our report.

### Replication

For the evaluation dataset we ran triplicates for each sample mixture, and experimental condition. In all cases 100% of samples passed quality control assessments and were used for downstream analyses.

### Randomization

For evaluation dataset, mass-spectrometry data on the same set of samples was acquired using two different chromatography conditions; sample groupings were assigned based on chromatography conditions used. For Global FoodOmics data, samples were grouped based on Earth Microbiome Project ontology (animal, plant, fungal).

### Blinding

Blinding is not relevant to our study. Sample collection, data acquisition and data analysis was standardized for all groups and all data is visualized without corrections or exclusions.

## Reporting for specific materials, systems and methods

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<input checked="" type="checkbox"/>	Animals and other organisms
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<input checked="" type="checkbox"/>	Dual use research of concern

**Methods**

n/a	Involved in the study
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<input checked="" type="checkbox"/>	Flow cytometry
<input checked="" type="checkbox"/>	MRI-based neuroimaging

**Human research participants**Policy information about [studies involving human research participants](#)

## Population characteristics

We used two fecal samples from a single participant (gender = male; age = 64) collected on 11-10-2013 and 12-14-2013 (also stated in the Methods section)

## Recruitment

The participant had stool samples collected by consent under the following protocol: HRPP 150275 (Evaluating the Human Microbiome). The Protocol was approved by the Human Research Protection Program (HRPP) of the University of California, San Diego.  
The participant was self-recruited (self-selection bias). The participant suffers from Crohn's disease, therefore the fecal metabolome data could include disease- and medication-related chemical signatures.

## Ethics oversight

Written informed consent obtained from the patient concerning dissemination and scientific publication of the results is also included in the approved protocols.

Note that full information on the approval of the study protocol must also be provided in the manuscript.