### Meta-Mass Shift Chemical (MeMSChem) profiling of metabolomes from coral reefs

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42 Abstract

43 Untargeted metabolomics of environmental samples routinely detects thousands 44 of small molecules, the vast majority of which cannot be identified. Meta-Mass 45 Shift Chemical (MeMSChem) profiling was developed to identify mass 46 differences between related molecules using metabolomic networks. This 47 approach illuminates metabolome-wide relationships between molecules and the 48 putative chemical groups that differentiate them (e.g.,  $H_2$ ,  $CH_2$ ,  $COCH_2$ ). 49 MeMSChem was used to analyze a publicly-available metabolomic dataset of 50 coral, algal, and fungal mat holobionts (i.e., the host and its associated microbes 51 and viruses) sampled from some of Earth's most remote and pristine coral reefs. 52 Each type of holobiont had unique mass shift profiles, even when the analysis 53 was restricted to parent molecules found in all samples. This result suggests that 54 holobionts modify the same molecules in different ways and offers insights into 55 the generation of molecular diversity. Three genera of stony corals had distinct patterns of molecular relatedness despite their high degree of taxonomic 56 57 relatedness. MeMSChem profiles also partially differentiated between individuals, 58 suggesting that every coral reef holobiont is a potential source of novel chemical 59 diversity.

60

#### 61 Significance Statement

62 Coral reef taxa produce a diverse array of molecules, some of which are
 63 important pharmaceuticals. To better understand how molecular diversity is

generated on coral reefs, metabolomes were analyzed using a novel approach
called Meta-Mass Shift Chemical (MeMSChem) profiling. MeMSChem uses the
mass differences between molecules in networks to determine how molecules
are related. Interestingly, the same molecules gain and lose chemical groups in
different ways depending on the taxa it came from, offering a partial explanation
for high molecular diversity on coral reefs.

70 \body

71 Untargeted tandem mass spectrometry is a powerful tool for wide-scale 72 analysis of small molecules. The resulting metabolomes are potential treasure 73 troves of new molecules and chemistries, but a major problem in realizing this 74 potential is that most detected molecules cannot be identified (1-5). One possible 75 solution is to use spectral fragmentation similarity to identify relatives of known molecules to generate novel annotations (6-8). These approaches have rapidly 76 77 expanded reference databases, but they remain inherently limited by the number 78 of known molecules. Therefore, there is a need for analyses that do not rely upon 79 molecular reference libraries (9).

80 The online platform Global Natural Products Social Molecular Networking 81 (GNPS: 5) uses spectral fragmentation patterns to compare tens of thousands of 82 molecular features and create networks of structurally-similar molecules. Here 83 we expand the analysis of GNPS networks to identify chemical differences between related molecules (Figure 1). This approach is called Meta-Mass Shift 84 85 Chemical (MeMSChem) profiling and it uses the mass differences (or mass 86 shifts) between related molecules to identify and annotate known chemical 87 groups such as H<sub>2</sub>, CH<sub>2</sub>, COCH<sub>2</sub>, etc. Annotating molecules based on their mass 88 shifts facilitates correlations between metabolomics, biochemistry and genomics, 89 which could help pinpoint sites of molecular modifications resulting from known 90 and unknown enzymatic activities.

91	Coral reefs are noted sources of novel, commercially-useful compounds
92	(10). Reef holobionts (e.g., corals, sponges, and algae with their associated viral
93	and microbial communities; 11) have unique metabolomes, with a high degree of
94	within-holobiont similarity (12, 13). The positive relationship between taxonomic
95	and molecular diversity is evident at the ecosystem level, but mechanisms
96	explaining how high molecular diversity is generated remain missing. To address
97	this question, MeMSChem was applied to an existing dataset (12) comprised of
98	seven coral reef holobiont types collected in the Line Islands, which are some of
99	the most remote and pristine coral reefs in the world (14, 15). MeMSChem
100	profiling showed that molecular mass shift patterns differ significantly between
101	holobionts, offering new insights into why high molecular diversity is found on
102	coral reefs.

103

#### 104 **Results**

105 Identifying redundant mass shifts in metabolomes in coral reef holobionts: 106 The dataset used as the basis for creating MeMSChem was previously published 107 in Quinn et al. 2016 (12) and can be found on the Mass spectrometry Interactive 108 Virtual Environment (MassIVE) at https://massive.ucsd.edu/ with the accession number: MSV000078598. This dataset was derived from an LC-MS/MS analysis 109 110 of three genera of scleractinian coral (Montipora spp., Pocillopora spp., Porites 111 spp.), two coralline algae (crustose coralline algae [CCA] and Halimeda sp.), two non-calcifying algae (macroalgae and turf algae) and a fungal mat. 112

113 The online platform Global Natural Products Social Molecular Networking 114 (GNPS, gnps.ucsd.edu; Figure S1; 5) was used to cluster identical MS/MS 115 spectra into nodes and identify the degree to which each node was structurally 116 similar (i.e., related) to other nodes (henceforth referred to as molecular features) 117 based on a cosine score of spectral similarity. All pairs of molecular features 118 receiving a cosine score above a threshold of 0.6 were considered to be related and connected in the network (see SI Materials and Methods for more details 119 120 regarding the cosine score threshold). Each mass shift between network 121 connections was then mined for multiple (i.e., redundant) occurrences (Figure 122 1B). When five or more molecular feature pairs differed by a mass of  $m/z \pm 1$ 123 0.001, the mass shift was counted. All molecular features comprising the pairs 124 with this mass shift were assigned to a bin (Figure 1C-E, see the SI Materials 125 and Methods for more details). 126 MeMSChem profiling identified 62 mass shifts that passed the filter of m/z127  $\pm$  0.001 and greater than 5 occurrences (Table S1). Among these mass shifts, 10 128 were annotated to known adducts and artifacts and were removed prior to further 129 analyses (Table S1 and S2). The remaining mass shifts were annotated to 130 known chemical groups involving hydrogen, carbon, and oxygen where possible, 131 leading to the annotation of 13 of the 62 mass shifts identified (Table S1). This 132 represents a conservative list of annotations and the additional mass shifts

133 identified here may be annotatable in future investigations.

134	Mass shifts of 0 were abundant in the networks and may represent
135	isomers. These mass shifts were removed due to the likelihood that two isomers
136	were merged into a single molecular feature or that the same molecular feature
137	was split into two molecules during networking, due to the high degree or
138	spectral similarity or differences in the number of observable fragments,
139	respectively. An approach using retention time differences or chiral separation
140	columns should be employed to separate isomers in future applications of
141	MeMSChem.
142	Quantifying mass shifts in holobionts: MS/MS-based molecular features
143	associated with the redundant mass shifts were quantified from the MS scan of
144	the parent molecule using the Optimus software
145	(https://github.com/MolecularCartography/Optimus; Figure 1C). A molecular
146	feature filter was applied to remove features that were not detected in all
147	samples. Consequently, only the features present in all samples were quantified.
148	This filter allowed us to determine whether holobionts exhibit different mass shifts
149	associated with the same molecules (c.f., different mass shifts associated with
150	molecules that are only found that holobiont; Figure 1E-F).
151	Three forms of metabolome-wide data were generated for each sample
152	(Figure 1A-C). First, all instances where a redundant mass shift was observed in
153	the network was tabulated for each sample. These 'counts' data provided a
154	metric of the commonness and rarity of each mass shift in each sample (Figure
155	1D). Second, the abundance of every molecular feature was summed by mass

156 shift regardless of whether that feature was the higher or lower mass feature in a 157 network pair. These 'summed abundance' data provided a metric for the overall 158 occurrence of each mass shift throughout each sample (Figure 1E; see the SI 159 Materials and Methods for equations). Third, for each network pair, the difference 160 in abundance between the more and less massive feature was calculated, then 161 these values were summed by mass shift for each sample (Figure 1F, see the SI 162 Materials and Methods for equations). These 'difference in abundance' data 163 reflected whether, metabolome-wide, molecules were more likely to gain or lose 164 a given mass, potentially reflecting active interconversion or branching of largely 165 shared biosynthetic pathways. All resultant data are provided in the Supporting 166 Information 'Processed Data' file. Among the redundant mass shifts, seven of the 167 ten most redundant mass shifts were putatively annotated to known chemical 168 groups, constituting nearly 50% of the network pairs isolated from the networks. 169 These mass shifts included *m/z* 2.016, 14.016, 28.032, 56.064, 26.016, 18.010, 170 12.000, which were putatively annotated as H<sub>2</sub>, CH<sub>2</sub>, C<sub>2</sub>H<sub>4</sub>, C<sub>4</sub>H<sub>8</sub>, C<sub>2</sub>H<sub>2</sub>, H<sub>2</sub>O, and 171 C, respectively.

Examining known mass shifts associated with library-identified molecular features: Instances in which known mass shifts were associated with identified molecules provided conformational evidence that mass shifts were correctly annotated. Four examples are highlighted in Figure 2B: 1) A feature identified as phenatro-furanone with a mass shift of *m/z* 18.014 (H<sub>2</sub>O; Figure 2B Example 1; Figure S2). 2) A subnetwork with three forms of Lyso-PAF and related

178	compounds (Figure 2B Example 2; Figure S3). The identification of one
179	molecular feature, Lyso-PAF C-16, in these sample was previously confirmed
180	using a reference standard by Quinn et al. 2016 (12). This subnetwork is
181	particularly informative because the three identified compounds were networked
182	to one another, showing that the mass shifts truly correspond to a de-saturation
183	and elongation of a fatty acid chain, $m/z$ 2.018 (H <sub>2</sub> ) and $m/z$ 28.032 (C <sub>2</sub> H <sub>4</sub> ). 3) A
184	subnetwork of ceramide-related compounds (Figure 2B Example 3; Figure S4)
185	with mass shifts of $m/z$ 2.015 (H <sub>2</sub> ), $m/z$ 14.015 (CH <sub>2</sub> ), and 165.057 (C <sub>6</sub> H <sub>10</sub> O <sub>5</sub> ,
186	glycosylation). A coral-associated ceramide was recently identified (16) with one
187	additional desaturation relative to the ceramide identified here, and this newly-
188	identified ceramide has an extremely similar mass ( $m/z$ 536.504) to the unknown
189	feature ( $m/z$ 536.508) networked to the ceramide here. The newly-identified
190	ceramide also differs in mass from the identified ceramide by $m/z$ 2.015,
191	consistent with one fewer saturation. 4) A subnetwork of three unidentified
192	molecules with mass shifts of $m/z$ 28.032 (C <sub>2</sub> H <sub>4</sub> ), $m/z$ 28.033 (C <sub>2</sub> H <sub>4</sub> ), and $m/z$
193	56.065 (C <sub>4</sub> H <sub>8</sub> ; Figure S5).
194	Differences in mass shift profiles between types of holobionts: To
195	determine how well MeMSChem profiling resolved each holobiont type, Random
196	Forests classification (17) was used to generate an out-of-bag error (henceforth

- 197 referred to as 'model error'), which reflects the extent to which the model
- 198 correctly categorized every sample (i.e., whether *Halimeda* sp. samples were
- 199 correctly placed into the model's Halimeda group). Random Forests offers

exceptional classification performance and is robust to non-normally distributed
data and correlated predictors (18), both of which were present in this dataset
(see the Supporting Information 'Processed Data' file).

203 The usefulness of re-categorizing molecules by their mass shifts was first 204 evaluated based on the number of times that each mass shift was observed 205 (counts data described above). The model error of the Random Forests 206 classifying holobiont types using the counts data was 0.44, which indicates that 207 44% of the time samples were assigned to the incorrect holobiont type. The 208 resolution gained from the observed counts data (i.e., actual data) was compared 209 to that from 1000 permutations of the data in which pairs were randomly binned 210 and counted while keeping the original proportions consistent. The observed 211 counts data outperformed 95% of the randomly-generated datasets, suggesting 212 that the counts of redundant mass shifts aided in differentiating between 213 holobiont types despite the relatively high model error (Figure 3A).

214 Molecular abundance data were then incorporated into the analysis and 215 compared against the holobiont resolution gained from the counts data. When 216 the summed abundances of each mass shift among molecules present in all 217 holobionts were considered, the model error from the abundance data was 0.36 218 (Figure 3B). Therefore, incorporating feature abundance data improved the 219 accuracy of the model by 8% when resolving between holobiont types. The value 220 of summing feature abundances by mass shift was also tested by comparing its 221 accuracy to the models of 1000 permutations of the data in which network pairs

222 were randomly binned and summed while keeping the original proportions 223 consistent (as was done for the counts data above). Among only the molecular 224 features present in all holobionts, summing of feature abundances by mass shift 225 resolved holobiont types better than 90% of the datasets generated from random 226 summing of feature abundances (Figure 3B). Thus, binning abundance data by 227 redundant mass shifts categorizes molecules in a non-random manner. 228 Molecular abundances binned by mass shifts also reflected differences among 229 holobiont types better than when holobionts were compared with data that lacks 230 any feature abundance information (i.e., counts of the number of mass shifts). 231 To determine whether mass shifts may reflect active sites of molecular 232 interconversions, as would be expected if a molecular modification had occurred, 233 the summed abundances were compared to the differences in abundance 234 between molecular pairs by mass shift. This is akin to one molecule being the 235 reactant and the other the product. The model error of the difference in 236 abundance data was 0.34, demonstrating that organizing the data by the 237 differences in abundances slightly outperformed the summed abundance data 238 (model error = 34% and 36%, respectively; Figure 3C). When compared to 1000 239 random permutations of the actual data, the difference in abundance data 240 outperformed 86% of the random models.

Classification was further improved by incorporating the full molecular dataset, and thus the molecules that were present in all holobionts and the molecules that were only found in one or a few holobionts. When these

244 molecules were included, the model error was 0.02. This reflects a 32% 245 decrease in the model error relative to when only molecules found in all 246 holobionts were considered and was nearly perfect in assigning samples to their 247 correct holobiont type. The real data outperformed 92% of randomly generated 248 datasets (Figure 3D and summarized in Figure 3E). These results suggest that 249 the highest level of holobiont resolution is achieved when: 1) molecular features 250 were binned by the redundant mass shifts they exhibit, 2) molecular abundances 251 were included as the difference in abundance between molecules in a network 252 pair, and 3) molecules/pairs that are only found in certain holobionts were 253 included in addition to those molecules present in all holobionts.

254 Mass shifts that best distinguish each holobiont type: Among the 255 molecular features present in all holobionts, coral genera were best differentiated 256 from one another by mass shifts corresponding to two carbons that were either 257 saturated (m/z 28.032, C<sub>2</sub>H<sub>4</sub> or 2\*CH<sub>2</sub>) or unsaturated (m/z 26.016 C<sub>2</sub>H<sub>2</sub>; p < 0.01 258 for both; Figure 4A). The three coral genera exhibited distinct patterns between 259 these two mass shifts: molecular features of *Montipora* exhibited the addition of 260  $C_2H_4$  and loss of  $C_2H_2$ , while *Pocillopora* exhibited the opposite pattern. *Porites*-261 associated molecules did not gain or lose either mass shift. Putative CH<sub>2</sub> and 262  $CH_2OOH$  mass shifts best differentiated the non-coral holobionts (p < 0.01 for 263 both; Figure 4B). Halimeda features predominantly gained CH<sub>2</sub>, as did turf algae, 264 the fungal mat, and all of the corals, though to a lesser degree than Halimeda. 265 Additions of CH<sub>2</sub>OOH were unique to Halimeda. Corals were best differentiated

from non-corals based on larger losses of CO and H<sub>2</sub>, the latter suggesting a
 dehydrogenated state perhaps due to higher concentrations of unsaturated lipids.

269 **Discussion** 

270 MeMSChem profiling provides an approach to identify mass shifts 271 between related molecules and assign them to known chemical groups in 272 complex metabolomes. Seven coral reef holobiont types were well resolved by 273 MeMSChem profiling. Even among molecular features detected in all holobionts, 274 mass shift profiles differed among holobiont types, suggesting that each type of 275 holobiont is modifying the same molecules in different ways. The chemical 276 differences between holobionts was much more apparent when all molecules 277 were considered (i.e., molecules only produced by certain holobionts were also 278 incorporated), suggesting that disparate mass shift patterns play a role in 279 generating molecular diversity in this ecosystem. Shifts in the abundance of 280 molecules exhibiting each mass shift better resolved holobiont types than the 281 number of times each mass shift was detected. Together, these findings suggest 282 that holobionts differ more in their patterns of molecular abundance changes 283 (akin to gene expression) than in the diversity of mass shifts they can carry out 284 (akin to genomic diversity).

285 *Mass shifts associated with holobionts reflect differences in molecular* 286 *diversity:* By focusing on the differences in mass shift profiles between related 287 molecules, MeMSChem profiling expands metabolomic analysis beyond

288 molecular matches in reference libraries to systemic insights into holobiont 289 biochemistry. If annotated mass shifts reflect single types or classes of molecular 290 modifications catalyzed by enzymes, then disparate mass shift patterns among 291 holobionts may arise for multiple reasons. Holobionts for which the hosts have 292 large genomic differences, such as stony corals and turf algae, may merely 293 possess different biochemical pathways. Among closely-related holobiont types 294 such as the three stony coral genera, the distinct patterns of molecular 295 relatedness may arise from differential expression of shared genes. Yet, the 296 largest disparity among coral holobionts was found by including the mass shifts 297 of molecules that are unique to each holobiont. This suggests that the mass 298 shifts of holobiont-specific molecules largely generate each coral holobiont's 299 unique biochemical profile despite the high degree of taxonomic relatedness 300 among these corals.

301 The mass shifts that differed among holobiont types included differences 302 putatively assigned to single and double-bonded carbon and oxygen, likely 303 among phospholipids and their derivatives based upon the molecules identified in 304 this dataset previously (12) and in the current analyses. These data show the 305 expected lower saturation state of corals relative to algae (19,20) based on the 306 mass shift of m/z 2.016 putatively assigned to H<sub>2</sub>. Greater fatty acid saturation 307 flexibility can mitigate the damage of elevated temperatures that lead to 308 bleaching in corals (21), suggesting that corals benefit from a higher degree of 309 saturation flexibility and homeoviscous adaptation potential relative to the non-

310 corals studied here. While desaturations in coral molecules generate double-311 bonds between carbons, the shift towards gaining H<sub>2</sub>O in coral samples suggest 312 these double bonds may be replaced by hydroxyl groups, either directly or 313 through shifts in the relative abundances of molecules. Hydration of 314 phospholipids can lead to conformational changes that alter membrane surface 315 potential and signaling activity (22), suggesting that the higher abundance of 316 hydroxyl groups in corals reflects systemic changes in cell-cell interactions and 317 cellular signaling pathways.

318 Applications of MeMSChem profiling: MeMSChem offers a new way to 319 analyze existing LC-MS/MS datasets and provides a novel approach for 320 identifying system-wide changes in small molecules across metabolomes. Here 321 we analyzed a dataset collected from one of the most remote places in the world. 322 Like this dataset, other researchers may have LC-MS/MS datasets that cannot 323 be re-sampled or recreated. Therefore, offering a way to gain novel information in 324 *silico* is an attractive proposition for many working with untargeted metabolomic 325 data.

While MeMSChem was applied here to uncover similarities and differences among types of holobionts, it opens doors to answering many more questions. Rather than comparing known groups, MeMSChem may be used to uncover clusters in seemingly homogenous populations (e.g., individuals of a coral species in a common environment, human patients suffering from the same disease, cohorts of offspring growing in a shared location). Annotated mass shifts

can also be searched for and quantified, which may be particularly useful whenlooking for a ubiquitous process such as antioxidant activity.

334 If molecules of interest are identified, the mass shifts around them may be 335 used to detect putative sites of known modifications or novel biochemistries. 336 Annotated and unknown mass shifts will require further verification with targeted 337 analyses, such as spiking samples with authentic standards, networking, and examining the mass shifts associated with these standards. Once putative 338 339 modifications are identified, genetics and molecular biology approaches can be 340 used to confirm or identify the responsible enzyme(s). Such an approach may be 341 particularly useful for tracking molecular changes in time-series samples, a 342 primary need for clinicians (23). Future applications of MeMSChem may also 343 employ a more precise binning approach, taking into account the smaller relative 344 variance at higher masses, changes in MS accuracy across parent masses, and 345 precursor differences. Through this process, the continued application of 346 MeMSChem and the novel form of data it generates will produce a wealth of 347 information from data-rich untargeted metabolomics datasets. 348 *Conclusions:* Untargeted metabolomics continues to grow as a tool to 349 examine the complex physiologies of life on Earth. We applied an approach that

analyzes untargeted metabolomic data in a new way, based on the chemical
relationships between molecules. An analysis of seven coral reef holobionts
demonstrated that the relationships between molecules are diverse and distinct
between holobiont types. That different types of holobionts had unique

MeMSChem profiles despite high genomic similarity suggests that each possesses physiological capabilities that are not easily identified through traditional genomic approaches. The distinct molecular repertoires identified in each holobiont, coupled with the wide diversity of holobiont types on coral reefs, offers new insights into why this ecosystem is an abundant source of chemical diversity.

360

## 361 Materials and Methods

362 LC-MS/MS data collection and molecular networking: Samples of seven 363 types of holobionts (hosts and associated viral and microbial communities) 364 including corals, algae, and a fungal mat were extracted in 70% methanol and 365 analyzed with LC-MS/MS (as described in Quinn et al. 2016 [12]; see the SI 366 Materials and Methods for data acquisition details). Files were submitted for 367 molecular network analysis using the online workflow in GNPS (Figure S1; 5), 368 which compares spectral fragmentation patterns and networks related molecules 369 (Fig. S1). Molecular spectra were also compared against reference libraries of 370 known molecules in GNPS. Details of the networking parameters can be found in 371 the SI Materials and Methods.

*Identifying aggregations of mass shifts in network pairs:* Across all pairs,
the difference in mass between each pair of networked molecular features
(referred to as 'network pair mass shifts') was searched for aggregations around
precise masses. Criteria for identifying aggregations (i.e., redundancies) were

376 established using the similar masses of CO and  $C_2H_4$  (*m/z* 27.995 and *m/z* 377 28.031, respectively; Figure S6; see the SI Materials and Methods for details). 378 The network pairs involved in aggregations were binned by mass shift and 379 counted per sample ('Counts' dataset in the SI 'Processed Data' file). All 380 molecular features involved in redundant mass shifts were then quantified using 381 the Optimus workflow (https://github.com/MolecularCartography/Optimus). 382 Optimus was used to quantify features involved in redundant mass shifts that 383 were present in all files/holobionts, features involved in redundant mass shifts 384 that were present in each holobiont type, and all molecular features including 385 those that were not involved in redundant mass shifts (for normalization of the 386 two former datasets). Molecular abundance data were then used to quantify the 387 molecules exhibiting each mass shift and to quantify the prevailing direction of 388 each mass shift (gaining or losing) in each sample (see the Results and SI 389 Materials and Methods for more details).

390 Data analysis using Random Forests: MeMSChem data were analyzed 391 using the ensemble machine learning algorithm Random Forests (16). The seven 392 holobiont types were used as classifiers and MeMSChem data were used as 393 predictors. The out-of-bag error (referred to as 'model error') indicated how well 394 each holobiont type was resolved by the Random Forests model. Permutation 395 tests were used to determine how well the MeMSChem data differentiated the 396 seven holobiont types. These tests were carried out by comparing the model 397 error of the actual data to a distribution of model errors generated from 1000

- 398 randomizations of the data (see the SI Materials and Methods for more details).
- 399 The relative importance of each mass shift in differentiating between holobiont
- 400 types was determined using the Random Forests mean decrease accuracy score
- 401 and feature importance score (for each holobiont type).

#### 402 Supporting Information

- 403 Figures S1–S6, Table S1–S2, SI Materials and Methods, R code for the
- 404 Random Forests permutation test, and the processed data for the four datasets
- 405 represented in Figure 3 can be found in the Supporting Information.
- 406

#### 407 Author Contributions

- 408 A.H. developed the method, analyzed data, prepared figures, and wrote the
- 409 manuscript, D.P. analyzed data and prepared figures, R.Q. analyzed data, I.P.
- 410 analyzed data and contributed analytical tools, F.A. analyzed data and
- 411 contributed coding and statistical approaches, E.R. analyzed data, G.W. and B.B.
- 412 contributed statistical approaches, M.V. collected samples, T.A contributed tools
- 413 for the mass spectrometry analysis, P.D. contributed tools for the mass
- 414 spectrometry analysis and interpretation, F.R. developed the method. All authors
- 415 discussed the results and commented on the manuscript.
- 416

#### 417 **Competing Financial Interests**

- 418 The authors declare no competing financial interests.
- 419

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#### 508 **Figure Captions**

#### 509 Figure 1. Data processing and generation based on a simplified molecular

510 **network and two redundant mass shifts.** (A) GNPS used MS/MS

- 511 fragmentation spectra to elucidate molecular similarities and network similar
- 512 molecules (i.e. related molecules). (B) Redundant mass shifts between related
- 513 molecules were identified and annotated to known chemical groups when
- 514 possible. (C) Molecular features that differed by a redundant mass shift were
- 515 quantified based on MS. Data were generated for (*D*) the number of times each
- 516 redundant mass shift was observed across all networks, (E) the summed

abundance of all molecules exhibiting each redundant mass shift, and (F) the

sum of the differences in abundances between the more massive and less

519 massive molecules for all pairs of molecules connected by a mass shift.

520

# 521 **Figure 2. Molecular network of the coral MS/MS dataset.** (*A*) The global

molecular networks of MS/MS spectra from the coral reef holobiont metabolomic dataset. Each node represents a unique or set of identical spectra (i.e., molecular feature). Lines connecting the nodes represent their spectral similarity, creating subnetworks that can be considered to be molecular families. Circles indicate zoomed-in regions of selected subnetworks shown in (*B*). Node labels represent parent masses and the line labels between the connected nodes represents the mass shift between related molecular features. Nodes with diamond shapes had

a spectrum library match, (e.g., Lyso-PAF, as identified by Quinn et al. 2016; 12)

and are further labeled with the molecular names. The size of the nodes

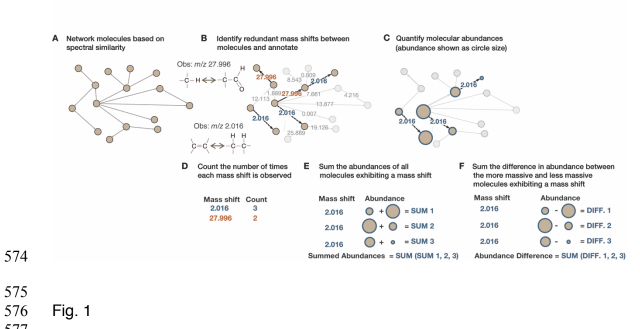
531 indicates the sample frequency in which the spectra were found.

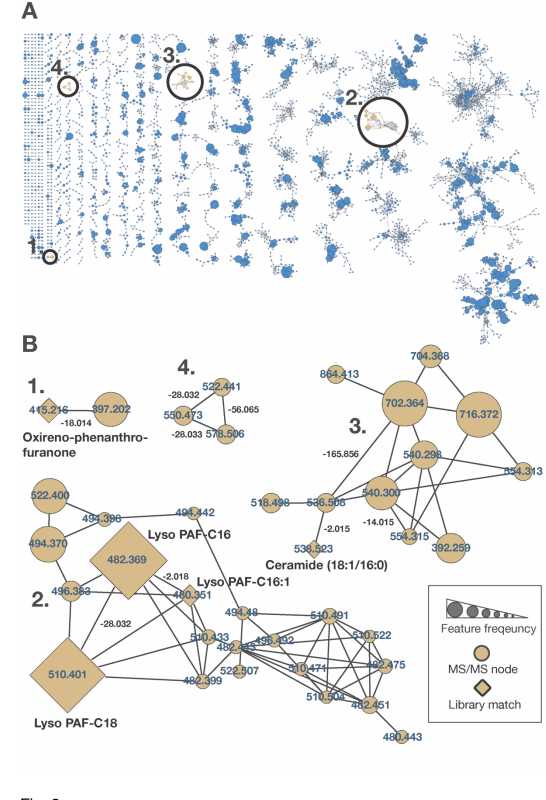
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533 Figure 3. Results of tests measuring the extent to which holobionts were 534 resolved by MeMSChem profiling. (A) A visualization of the first two 535 dimensions of a Random Forests proximity matrix of the number of times that 536 each redundant mass shift was identified (counts data). The color of the filled 537 circle represents the holobiont type of the sample while the color of the halo 538 around each circle corresponds to the holobiont type it was placed in by the 539 Random Forests model (i.e., if the circle and halo are different colors the model 540 incorrectly categorized the sample). (B) An analogous representation of (A) for 541 the summed abundances of molecules grouped by the mass shifts they exhibit 542 among only the molecular features present in all holobionts. (C) An analogous 543 representation of (A) using the difference in abundances of molecules 'gaining' 544 minus 'losing' a mass, summed by the mass shift they exhibit among only the 545 molecular features present in all holobionts. (D) An analogous representation of 546 (A) using the difference in abundances of molecules 'gaining' minus 'losing' a 547 mass, summed by the mass shift they exhibit among all the molecules in the 548 dataset. (E) A histogram of the permutation test from randomly generated 549 datasets to determine how well MeMSChem profiling resolves each holobiont 550 type based on the model error. Letters above each line correspond to the model 551 error of the actual data in the figure panel matching that letter. The histograms

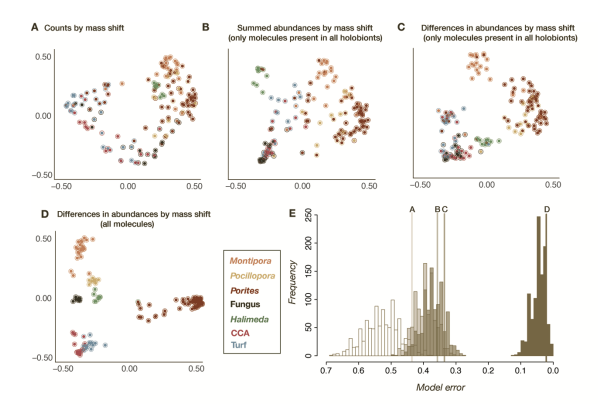
reflect the model errors of 1000 permutations of the actual data in which pairs were randomly binned while keeping the original proportions consistent. This was repeated for the data in (A-D), the distributions for which are shown in order and darkening color of counts, summed abundances, differences in abundances in molecules present in all holobionts, and differences in abundances in the entire molecular dataset.

559	Figure 4. The annotated mass shifts that best differentiated each holobiont
560	type. (A) The annotated mass shifts that best distinguish between coral genera
561	based on the mean decrease accuracy of a supervised Random Forests. (B) The
562	annotated mass shifts that best distinguish between the non-coral holobiont
563	types. ( $C$ ) The annotated mass shifts that best distinguish the coral holobionts
564	from the non-coral holobionts.
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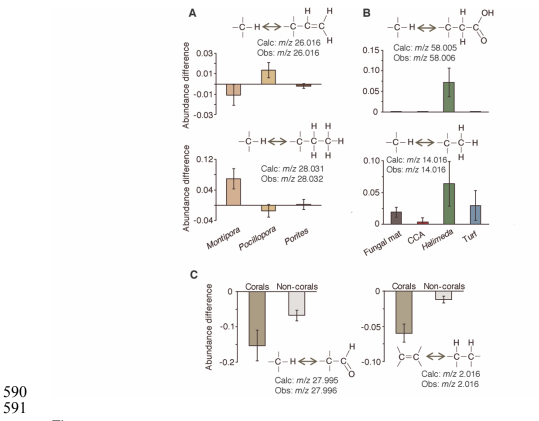




- 582 Fig. 2







592 Fig. 4