## 1 Mining metagenomes for natural product biosynthetic gene

## 2 clusters: unlocking new potential with ultrafast techniques

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Microorganisms produce an immense variety of natural products through 20 the expression of Biosynthetic Gene Clusters (BGCs): physically clustered 21 genes that encode the enzymes of a specialized metabolic pathway. These 22 23 natural products cover a wide range of chemical classes (e.g., 24 aminoglycosides, lantibiotics, nonribosomal peptides, oligosaccharides, polyketides, terpenes) that are highly valuable for industrial and medical 25 26 applications<sup>1</sup>. Metagenomics, as a culture-independent approach, has greatly enhanced our ability to survey the functional potential of 27 28 microorganisms and is growing in popularity for the mining of BGCs. 29 However, to effectively exploit metagenomic data to this end, it will be 30 crucial to more efficiently identify these genomic elements in highly 31 complex and ever-increasing volumes of data<sup>2</sup>. Here, we address this challenge by developing the ultrafast Biosynthetic Gene cluster 32 MEtagenomic eXploration toolbox (BiG-MEx). BiG-MEx rapidly identifies a 33 broad range of BGC protein domains, assess their diversity and novelty, 34 35 and predicts the abundance profile of natural product BGC classes in 36 metagenomic data. We show the advantages of BiG-MEx compared to

37 standard BGC-mining approaches, and use it to explore the BGC domain 38 and class composition of samples in the TARA Oceans<sup>3</sup> and Human 39 Microbiome Project datasets<sup>4</sup>. In these analyses, we demonstrate BiG-40 MEx's applicability to study the distribution, diversity, and ecological roles 41 of BGCs in metagenomic data, and guide the exploration of natural 42 products with clinical applications.

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Metagenomics offers unique opportunities to mine natural product BGCs in 44 diverse microbial assemblages from a wide range of environments<sup>5-7</sup>. However, 45 given the complexity of microbial communities found in nature, and the limitations 46 of current sequencing technologies, often only a very small fraction of the short-47 48 read sequence data can be assembled in contigs long enough to allow the identification of BGC classes. However, the annotation of individual protein 49 50 domains of BGCs, is much more straightforward, given that these have 51 comparable length to merged paired-end reads. There are several protein 52 domains known to play important functions in the BGC-encoded enzymes. 53 Specific domains or combinations thereof are commonly found in certain types of 54 BGC classes. Accordingly, these are used for the automatic identification of BGC classes in genome sequences<sup>8-10</sup> and to study the distribution and diversity of 55 particular BGC classes in the environment<sup>6,7,11–13</sup>. Although there are various BGC 56 mining tools with practical applications<sup>14</sup>, only the Natural Product Domain 57 Seeker (NaPDoS)<sup>11</sup> and the environmental Surveyor of Natural Product Diversity 58 (eSNaPD<sup>15</sup>) are dedicated to the study of BGC domains. Both of these tools 59 60 focus on nonribosomal peptides and polyketide synthases (NRPSs and PKSs, 61 respectively), and take assembled or amplicon data as input. Currently, there is 62 no technology available capable of efficiently exploiting raw metagenomic data to 63 study the composition and diversity of natural product BGC classes and domains 64 in the environment.

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66 Capitalizing on the fact that BGC domains can be readily annotated in 67 unassembled metagenomic data, and used to identify the different natural

68 product BGC classes, we developed BiG-MEx. This tool generates ultrafast BGC 69 domain annotations in short-read sequence data and applies a machine-learning 70 approach to predict the BGC class coverage-based abundances (for simplicity, 71 we will refer to these as BGC class abundance profiles). Additionally, the identified domain sequences are used to carry out a domain-based diversity 72 73 analysis. This allows BiG-MEx both to deeply exploit metagenomic data, and to 74 adapt to their ever-increasing volume. BiG-MEx consists of three interacting 75 modules that are described below and illustrated in Fig. 1:

 BGC domain identification module. We use the Ultrafast Protein domain Classification UProC<sup>16</sup> tool to identify BGC protein domains in short-read sequence data. For this purpose, we created an UProC database, which includes 150 BGC domains covering 44 BGC classes.

2. BGC domain-based diversity analysis. This module performs a domain-80 81 targeted assembly, clusters the assembled domain sequences to create Operational Domain Units (ODUs)<sup>17</sup> and computes the ODU alpha diversity. 82 83 Further, assembled domain sequences are placed onto reference 84 phylogenetic trees. The module includes pre-computed phylogenies for 48 85 BGC domains. These were selected based on domain sequences from 86 experimentally characterized biosynthetic gene clusters with enough 87 sequence information for phylogenetic analysis.

3. BGC class abundance prediction module. We created machine-learning models that predict the abundance of BGC classes based on the domain annotation. The models are class-specific and consist of a random forest (RF) classifier to predict the presence/absence of a BGC class, and a multiple linear regression (MLR) to predict its abundance. These models can be customised to target metagenomic and genomic data from different environments and taxa, respectively.

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96 To evaluate the performance of BiG-MEx, we first assessed how the UProC-97 based domain identification used in BiG-MEx improves the data processing 98 speed compared to HMMER<sup>18</sup> (i.e., the traditional approach for domain

99 annotation) for the annotation of the 150 BGC domains. This comparison showed 100 that UProC was on average 18 times faster than HMMER (Supplementary Fig. 101 1a). We then evaluated the accuracy of BiG-MEx Operational Domain Unit 102 (ODU) diversity estimation approach. We used BiG-MEx to compute the ODU 103 diversity of the NRPS adenylation (AMP-binding) and condensation domains, as 104 well as the PKS ketosynthase (PKS KS) and acyltransferase (PKS AT) domains in a simulated metagenomic dataset (Marine-TM dataset; see Materials and 105 106 Methods section 3). Additionally, we computed the ODU diversity of these 107 domains based on the domain sequences obtained from the genome sequences used to simulate the Marine-TM metagenomes. The latter estimates (henceforth, 108 109 the reference estimates) were assumed to accurately reflect the ODU diversity, 110 as they were computed using the complete domain sequences. We compared BiG-MEx ODU diversity estimates against the reference ODU diversity and 111 112 observed that these were highly correlated: PKS KS domains had a Pearson's r 113 of 0.77, while for the other domains the Pearson's r was greater than 0.9 114 (Supplementary Fig. 1b). Lastly, we evaluated BiG-MEx's BGC class abundance 115 prediction module. We point out that although we modelled the abundance of a 116 few BGC subclasses, we refer to all as BGC classes. For this analysis, we used 117 two different simulated metagenomic datasets, one for training and the other for 118 testing the BGC class abundance models (Marine-RM and Marine-TM, respectively) (see Supplementary Table 1). We predicted the BGC class 119 120 abundances in the Marine-TM metagenomes, using BiG-MEx BGC class 121 abundance prediction module, and additionally, computed the BGC class 122 abundances based on the complete genome sequences used to simulate the 123 Marine-TM metagenomes. Similarly as indicated previously, the latter abundances were taken as a reference to evaluate the accuracy of the 124 125 predictions. We observed that the predicted vs. reference abundance comparison for 20 of the 23 BGC classes we modelled (i.e., the total number of 126 127 classes detected in the Marine-RM training dataset) had a Pearson's r correlation coefficient greater than 0.5 and a median unsigned error (MUE) lower than 0.25 128 129 (Supplementary Fig. 2). Figure 2a displays the scatter plots of this comparison for

130 the NRPS, terpene, and type I and II PKS BGC classes. To benchmark BiG-MEx 131 BGC class abundance prediction module, we compared its abundance predictions against the abundance estimates derived from running antiSMASH 132 on assemblies of the Marine-TM metagenomes (hereafter referred to as the 133 "assembly approach"). The plots in Figure 2b display the Pearson correlation 134 135 coefficients and the unsigned error distributions with respect to the reference abundances comparing both approaches for the same four BGC classes 136 137 mentioned above. All BGC class abundance models included in this analysis 138 were considerably more accurate than the assembly approach (Supplementary 139 Fig. 3).

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141 To illustrate the application of BiG-MEx, we performed a Principal Coordinates Analysis (PCoA) based on BiG-MEx-derived BGC class abundance profiles of 142 143 the 139 prokaryotic metagenomes of TARA Oceans. In Figure 3a, we ordinate the first two axes of the PCoA. The first axis (PCo1: 73.5% of the total variance) 144 145 differentiated the mesopelagic (MES) from the surface (SRF) and deep 146 chlorophyll maximum (DCM) water layers (Wilcoxon rank sum test; all p-values < 147 0.0001; see Supplementary Table 2). Further, the ordination values of the 148 metagenomes along the PCo1 axis correlated with temperature (Pearson's r = -149 0.73; p-value < 0.0001). The differences in the BGC class composition between water layers were additionally confirmed with a Permutational Multivariate 150 151 Analysis of Variance (PERMANOVA) (see Supplementary Table 3). We also performed a PCoA to explore the BGC domain composition and obtained a 152 153 similar ordination of the metagenomes (Supplementary Fig. 4). These results are 154 in agreement with previous work showing the stratification of microbial communities along depth and temperature gradients<sup>19,20</sup>. In particular, a very 155 similar differentiation of the MES water layer along the first axis was also 156 observed in the PCoA performed by Sunagawa et al.,<sup>19</sup> based on the 16S mitag 157 (i.e., 16S ribosomal RNA gene tags<sup>21</sup>) composition of these same TARA Ocean 158 159 metagenomes.

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Next, we used BiG-MEx domain-based diversity module to compare the 161 162 Operational Domain Unit (ODU) diversity of the NRPS adenylation (AMP-binding) and condensation domains between the SRF, DCM and MES water layers. 163 These domains provide information about the chemical characteristics of the 164 peptides synthesized by NRPS enzymes. AMP-binding domains recruit the 165 166 amino acid monomers to be incorporated, while condensation domains catalyse the peptide bond formation<sup>22,23</sup>. In this analysis, we aimed to assess the potential 167 168 chemical diversity of the NRPS products. NRPSs are one of the most studied 169 BGC classes and are responsible for the production of many compounds with clinical applications. The results show that the ODU diversity of both domains 170 171 increased from the surface to the mesopelagic water layers and differentiated 172 significantly between water layers (pairwise Wilcoxon rank sum test; all p-values < 0.005; see Supplementary Table 2) (Fig. 3b). These results indicate that the 173 174 microbial communities inhabiting deeper water layers contain a significantly 175 higher diversity of NRPS products. The ODU diversity gradients resemble the Operational Taxonomic Unit (OTU) richness and functional diversity distributions 176 177 shown in Sunagawa et al. We found highly significant correlations between the 178 ODU diversity estimates and the taxonomic and functional richness and diversity 179 obtained by Sunagawa et al. (see Supplementary Table 4).

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181 To exemplify a more fine-grained analysis with BiG-MEx's domain-based 182 diversity module, we explored the ODU diversity of condensation domains in the 183 three TARA Oceans metagenomes obtained from the SRF, DCM, and MES 184 water layers at the sampling station TARA 085 (Antarctic Ocean). As observed 185 previously, the metagenome from the MES water layer had a higher ODU diversity (Fig. 4a). It contains many low abundance ODUs scattered throughout 186 the reference phylogeny (Fig. 4b). The phylogenetic diversity<sup>24</sup> (PD) of ODU 187 188 representative sequences of the MES metagenome, was 5.24 and 2.65 times 189 greater than the PD estimates of the SRF and DCM metagenomes, respectively. 190 Besides indicating a higher chemical diversity, this result indicates that there is 191 greater potential chemical novelty of nonribosomal peptides. Additionally, the 192 phylogenetic placement analysis revealed that the most abundant condensation 193 ODU is placed close to the reference condensation domain sequences of NRPSs 194 that produce albicidin and cystobactamide antibiotics (both topoisomerase 195 inhibitors) (Fig. 4c). As albicidin is also a phytotoxin, the dominance of such 196 ODU, which originates from the DCM layer, could be explained by the presence 197 of a large number of NRPSs that act on the photosynthetic organisms that 198 concentrate therein. The DCM layer had a notably higher chlorophyll 199 concentration than the other two layers (0.01, 0.28, and 0 mg/m3 for the SRF, 200 DCM, and MES respectively). The NRPS producing albicidin belongs to the class 201 Gammaproteobacteria and order Xanthomonadales. This is in agreement with 202 the ODU taxonomic affiliation, which was annotated as a Gammaproteobacteria 203 (lowest common ancestor). This finding is also supported by the fact that the 204 BLASTP search against the reference MIBiG database, showed that 205 condensation domains significantly similar to NRPS domains producing albicidin 206 (e-value < 1e-5), where only found in the DCM layer. We cannot exclude other 207 possible explanations of these results; however, this line of exploration might be 208 worth considering for further research. Rising ocean temperatures, as a 209 consequence of global warming, are predicted to increase the frequency of events of bacteria affecting the algae populations, which in turn can impact 210 marine ecosystems on a global scale<sup>25</sup>. Regarding potential biotechnological 211 212 applications, these results are relevant for bioprospecting, given that albicidin 213 and cystobactamide are antibiotics of interest for clinical treatments<sup>26,27</sup>.

214 We note that neither the TARA Oceans Metagenomes Assembled Genomes 215 (MAGs)<sup>28</sup>, nor the DCM assembled metagenome from TARA 085 sampling site, 216 contained albicidin or cystobactamide NRPS-like sequences. The difference 217 between our findings in comparison to standard approaches based on 218 assembled data was expected to occur, given the limitations of the latter to 219 identify BGC classes (as shown in Fig. 2). In Supplementary Figure 5, we 220 illustrate this problem by comparing the sequence length between MIBiG BGCs, 221 and the TARA Oceans MAGs, and assembled metagenomic contigs.

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223 Considering the relevance of human microbiome-derived natural product BGCs 224 in medical research, we demonstrate the applicability of BiG-MEx to explore the 225 BGC composition in the Human Microbiome Project (HMP) dataset. Our analyses 226 traversed metagenomes from the buccal mucosa, tongue dorsum, and 227 supragingival plaque body sites as well as stool samples (491 metagenomes in 228 total). We used BiG-MEx to compute the BGC domain and class abundance 229 profiles, and applied the same methodology as described for TARA Oceans, to 230 compute the domain and class-based PCoAs. In agreement with previous analyses based on the taxonomic and functional annotation<sup>4,29</sup>, we observed that 231 232 metagenomes grouped according to the body site they were sampled from in the 233 first two ordination axes (Supplementary Fig. 6a and b). We conducted a 234 PERMANOVA to test and assess the strength of the differences between body 235 sites according to their BGC class composition, which showed significant 236 differences in all body site comparisons (Supplementary Table 5). Additionally, 237 we used BiG-MEx to compare the ODU diversity of the AMP-binding and 238 condensation domains between body sites and observed that supragingival 239 plaque metagenomes contain significantly higher diversity than the other body 240 sites (pairwise Wilcoxon rank sum test; p-value < 0.0001) (Supplementary Figure 241 7 and Supplementary Table 6). This is in line with previous work showing that the 242 supragingival plaque is one of the most functionally and taxonomically diverse 243 body sites in the HMP dataset<sup>4</sup>.

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245 Besides the mining analyses, BiG-MEx BGC class profiling can be used for the 246 screening and prioritization of (meta)genomic samples. BGC class abundance 247 profiles derived from shallow sequencing depth (meta)genomic data can be used 248 for the identification of strains or environments with high biosynthetic potential, 249 before investing in deep sequencing or long read sequencing technologies. As a 250 proof-of-concept for this application, in Supplementary Fig. 8 we show a 251 comparison of the BGC class abundance predictions computed in metagenomes of 100 and 5 million reads. 252

In our example applications, we processed 630 metagenomes, which sum to more than 85 billion paired-end reads. The analyses showed that BiG-MEx ultrafast domain and class profiling, and ODU diversity estimates provide biologically meaningful information, which can be used to mine BGCs in metagenomic data and as a basis from which to assess the ecological roles of their products in specific environments.

BiG-MEx extends BGC-based research and exploitation into large environmental datasets. It can be used to study the biogeography, distribution, and diversity of natural product BGCs either at the class, domain or ODU levels. Such analyses have the potential to accelerate the discovery of new bioactive products.

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### 264 Materials and Methods

### 265 **1. Data acquisition, pre-processing and annotation**

266 We retrieved the 139 prokaryotic metagenomes of the TARA Oceans dataset from the European Nucleotide Archive<sup>30</sup> (ENA:PRJEB1787, filter size: 0.22-1.6 267 268 and 0.22-3). To pre-process the metagenomic short-read data, we clipped the 269 adapter sequences (obtained from Shinichi Sunagawa personal communication, 270 July 21, 2015) using the BBDuk tool from the BBMap 35.00 suite (https://sourceforge.net/projects/bbmap/) with a maximum Hamming distance of 271 272 one (hdist=1). We then merged the paired-end reads using VSEARCH  $2.3.4^{31}$ , 273 quality trimmed all reads at Q20 and filtered out sequences shorter than 45bp 274 using BBDuk, and de-replicated the resulting quality-controlled sequences with 275 VSEARCH. We annotated the BGC domains by first predicting the Open Reading Frames (ORFs) in the pre-processed data with FragGeneScan-plus<sup>32</sup> and then 276 277 running BiG-MEx on the predicted ORF's amino acid sequences.

278 We downloaded 491 human microbiome metagenomes from the Data Analysis 279 and Coordination Center (DACC) for the Human Microbiome Project (HMP) 280 (https://www.hmpdacc.org/hmp/HMASM/). included Our dataset the 281 metagenomes of the supragingival plaque (118), tongue dorsum (128), buccal 282 mucosa (107), and the stool (138) body sites. These metagenomes have been 283 already pre-processed as described in The Human Microbiome Project 284 Consortium 2012<sup>33</sup>. The additional pre-processing tasks we performed consisted 285 of merging the metagenomic reads with VSEARCH, quality trimming all reads at 286 Q20 and filtering out sequences shorter than 45 bp with BBduk. To annotate the 287 BGC domains, we predicted the ORFs with FragGeneScan-plus and ran BiG-288 MEx BGC domain identification module on the ORF's amino acid sequences 289 (Supplementary Table 7).

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### 291 **2. Exploratory analysis performed on TARA Oceans and HMP datasets**

292 The domain abundance profiles of the TARA Oceans and HMP metagenomes were used to predict the BGC class abundance profiles with BiG-MEx BGC class 293 294 abundance prediction module. The models used to generate the predictions for 295 the TARA Oceans, and the oral and stool HMP metagenomes, were trained with 296 the Marine-RM, Human-Oral and Human-Stool simulated metagenomic datasets, 297 respectively. For each dataset, we performed a Principal Coordinate Analysis 298 (PCoA) as follows: 1) We applied a total sum scaling standardization to both the 299 domain and class abundance matrices; 2) We used the standardized matrices to 300 compute the domain and class Bray-Curtis dissimilarity matrices; 3) We 301 performed the PCoAs on the dissimilarity matrices with vegan R package utilizing the function capscale<sup>34</sup>. 302

303 We applied a Permutational Multivariate Analysis of Variance (PERMANOVA)<sup>35</sup> 304 to quantify the strength and test the differences between water layers and body 305 sites according to their BGC class composition. For these analyses, we selected 306 a balanced subset of metagenomes from the TARA Oceans and HMP datasets 307 (63 and 216 metagenomes, respectively; see below). We performed a 308 PERMANOVA on the Bray-Curtis dissimilarity matrix, computed for the TARA 309 Oceans and HMP metagenome subsets as described above, to test the 310 differentiation between all groups simultaneously. Subsequently, we tested each 311 pair of groups independently, applying the Bonferroni correction for multiple 312 comparisons. To perform the PERMANOVA, we employed the adonis function of 313 the vegan R package, with the permutation parameter set to 999.

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To compare the domain ODU diversity of the NRPS adenylation (AMP-binding) 315 316 and condensation domains between the surface (SRF), deep chlorophyll 317 maximum (CDM) and mesopelagic (MES) water layers, we used a subset of 63 318 TARA Oceans metagenomes, representing the three water layers in 21 sampling 319 stations. We computed the ODU Shannon diversity in these metagenomes, using 320 routines implemented in the BiG-MEx domain-based diversity module. 321 Additionally, we used the same BiG-MEx module to examine the diversity of the 322 condensation domains in the metagenomes representing the three water layers 323 at sampling station TARA 085. To perform the ODU taxonomy annotation, we MMsegs2 taxonomy assignment function<sup>36</sup> based on UniRef100<sup>37</sup> 324 used 325 sequences (release-2018 08), with the e-value and sensitivity parameters set to 326 0.75 and 0.01, respectively. To compare the AMP-binding and condensation 327 ODU diversity between body sites, we applied a similar approach as described 328 above. We selected a subset of 216 metagenomes, 54 from each of the 329 supragingival plague, tongue dorsum, buccal mucosa, and stool body sites. This 330 subset includes only the metagenomes obtained from individuals for whom the 331 four body sites were sampled. We applied BiG-MEx domain-based diversity 332 module to compute the ODU Shannon diversity estimates.

The Wilcoxon rank-sum tests (two-sided) to assess the significance of the differentiations between metagenomes from different groups (i.e., water layers or body sites), were performed with the wilcox.test function from the R package stats<sup>38</sup>.

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### 338 **3. Data simulation, pre-processing and annotation**

### 339 **3.1 Construction of simulated metagenomic datasets**

We created four simulated metagenomic datasets: Two of these approximate the taxonomic composition found in marine environments (Marine-RM and Marine-TM), and the other two, the taxonomic composition found in the human oral cavity and stool body sites (Human-Oral and Human-Stool, respectively). Each dataset is composed of 150 metagenomes, all of which have a size of two million paired-end reads. To simulate a metagenomic dataset, we first created a dataset of reference genome sequences and the genome
abundance profiles to specify the metagenomes' taxonomic composition. That
is, we defined a hypothetical microbial community from which a metagenome
is simulated by specifying which reference genomes and the number of times
each genome occurs in the community.

To create the Marine-RM (Marine Reference Microbiome) genome dataset, we downloaded all genomes belonging to the Ocean Microbial Reference Gene Catalogue (OM-RGC)<sup>19</sup> having an assembly status of "Complete genome" from RefSeq<sup>39</sup> (on December 7<sup>th</sup>, 2017). If a given species did not have a complete genome sequence available, we randomly selected another species of the same genus. In total, we obtained 378 genomes corresponding to 363 species.

358 We applied a similar methodology to create the Marine-TM (Marine TARA 359 Microbiome) genome dataset. To determine the taxonomic composition, we 360 used the genus affiliation of TARA Oceans Operational Taxonomic Units 361 (OTUs)<sup>19</sup>. We only included 30 shared genera (randomly selected) between 362 TARA OTU and the Marine-TM genome dataset. This latter filtering was 363 necessary to reduce the taxonomic overlap, given that we used the Marine-TM 364 dataset to evaluate the performance of the BGC class abundance models 365 trained with the Marine-RM dataset (see section 4.3). For the remaining 366 genera for which there was at least one representative completely sequenced 367 genome, we downloaded a maximum of three genomes per genus from 368 RefSeq, irrespective of their species affiliation. This resulted in a database 369 composed of 344 genomes from 308 species.

To create the genome datasets for the Human-Oral and Human-Stool metagenomic datasets, we used the genomes sequenced by the HMP derived from samples of the oral cavity and stool body sites. Given that few of these genomes were completely sequenced, we also included partially complete sequenced genomes. We downloaded all genomes with an assembly status of "Complete genome" or "Chromosome" or "Scaffold" generated by the HMP from the GenBank database<sup>40</sup> (on March 15th, 2018). In the cases where a

genome (sequenced by the HMP) had an assembly status lower than
"Scaffold", we downloaded another genome with the same species affiliation
and an assembly status of "Complete genome" or "Chromosome". The
Human-Oral and Human-Stool reference genome datasets contain 209, and
479 genomes representing 140 and 338 species, respectively.

382 To create the community abundance profile of a metagenomic dataset, we 383 randomly selected between 20 and 80 genomes from its genome reference 384 dataset and defined the number of times each genome occurs by sampling 385 from a lognormal distribution with mean 1 and standard deviation of 0.5. 386 Lastly, we simulated the metagenomes with MetaSim v0.9.5<sup>41</sup>. MetaSim was 387 set to generate paired-end reads with a length of 101bp, and a substitution rate increasing constantly along each read from 1×10<sup>-4</sup> to 9.9×10<sup>-2</sup>. With this 388 389 data, we aimed to simulate the short-read sequences generated by an Illumina 390 HiSeq 2000 platform.

391 Dataset statistics are shown in Supplementary Table 1. The assembly 392 accessions, organism names, taxids and NCBI FTP paths of the genome 393 sequences used to create the genome databases are found in the 394 Supplementary File 1. The workflow used to create the simulated 395 metagenomic datasets can be found at <u>https://github.com/pereiramemo/BiG-</u> 396 MEx/wiki/Data-simulation

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### 398 **3.2 Annotation of the simulated metagenomes**

399 To estimate the reference BGC class abundances in a simulated 400 metagenome, we annotated the BGC classes in its reference genome 401 sequences with antiSMASH 3.0, mapped the paired-end reads to the identified 402 BGC sequences with BWA-MEM 0.7.12<sup>42</sup>, and filtered out read alignments 403 with a quality score lower than 10. Next, we removed read duplicates with 404 Picard tools v1.133 (http://broadinstitute.github.io/picard), and computed the mean coverage with BEDtools v2.23<sup>43</sup>. The coverage estimates were assumed 405 406 to accurately reflect the BGC class coverage-based abundances, as they were 407 computed using complete BGC sequences, obtained from the genome sequences used to simulate the metagenomes. Additionally, we merged the
paired-end reads of the simulated metagenomes with VSEARCH 2.3.4,
predicted the ORFs with FragGeneScan-plus, and used BiG-MEx domain
identification module to annotate the BGC domains in the ORF's amino acid
sequences. The workflow to annotate the synthetic metagenomes can be
found at <a href="https://github.com/pereiramemo/BiG-MEx/wiki/Data-simulation#7-bgc-domain-annotation">https://github.com/pereiramemo/BiG-MEx/wiki/Data-simulation#7-bgc-domain-annotation</a>

## 416 **4. Performance evaluation**

## 417 **4.1 BGC domain identification module**

418 We compared the running time (wall-clock) of UProC (i.e., uproc-prot) against 419 a typical search using hmmsearch from the HMMER3 package<sup>18</sup>, for the 420 identification of the 150 BGC domains included in BiG-MEx, in nine prokaryotic 421 metagenomes of the TARA Oceans dataset (Supplementary Table 8). To run 422 hmmsearch, we used the domain HMM profiles of antiSMASH. We annotated 423 the nine metagenomes with both these tools in four independent rounds, each 424 round using a different thread number (i.e., 4, 8, 16 and 32 threads). All 425 parameters of uproc-prot and hmmsearch were set to default. The annotations 426 were carried out on a workstation with Intel(R) Xeon(R) CPU E7-4820 v4 427 2.00GHz processors.

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## 429 **4.2 BGC domain-based diversity analysis module**

430 We evaluated BiG-MEx Operation Domain Unit (ODU) diversity estimation 431 approach using NRPS adenylation (AMP-binding) and condensation, and PKS 432 ketosynthase and acyltransferase domains (PKS KS and PKS AT. 433 respectively). In this analysis, we used the BGC domain-based diversity 434 analysis module to compute the ODU diversity in the Marine-TM dataset, and 435 compared these estimates with the ODU diversity computed using the 436 complete domain sequences. To obtain the latter ODU diversity, we applied the workflow implemented in BiG-MEx, with the exception that instead of 437 438 assembling the domain sequences, we extracted these from the complete

genome sequences used to simulate the Marine-TM metagenomes. We
annotated the four domains in the complete genome sequences with
hmmsearch using the antiSMASH HMM profiles.

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### 443 **4.3 BGC class abundance predictions**

444 We used the BGC class models trained with the Marine-RM metagenomic dataset to predict the BGC class abundances in the Marine-TM metagenomic 445 dataset. We applied the methodology described in section 3.2 to compute the 446 447 BGC class abundances in the Marine-TM metagenomes based on the complete genome sequences (i.e., reference abundance). To predict the BGC 448 449 class abundances using machine-learning models, we annotated the Marine-450 TM metagenomes with the BiG-MEx domain identification module and used the domain abundance profiles as an input for the BiG-MEx BGC class 451 452 abundance prediction module. The evaluation consisted of computing the 453 Pearson correlation and median unsigned squared error (MUE) between the predicted and reference BGC class abundances. The MUE was computed as 454  $|\hat{A} - A|/A$ , where  $\hat{A}$  and A are the predicted and reference abundance, 455 456 respectively. To benchmark the machine-learning models, we compared the 457 BGC class abundance predictions against the abundance estimates based on 458 the assembly of 50 metagenomes of the Marine-TM dataset (assembly 459 approach). The assembly approach consisted of assembling the 460 metagenomes with MEGAHIT (default parameters), running BiG-MEx domain 461 identification module to select the contigs with potential BGC sequences, 462 annotating the selected contigs with antiSMASH 3.0, and estimating the BGC 463 class abundance following the same approach as described in section 3.2 464 (Supplementary Table 9). We computed the unsigned error, and the Pearson 465 correlation coefficient of BGC class abundance estimates obtained by the 466 assembly approach and predicted by BiG-MEx, with respect to the reference BGC class abundances. The analysis performed to evaluate the accuracy of 467 468 the models can be reproduced here: https://rawgit.com/pereiramemo/BiG-469 MEx/master/machine leaRning/bgcpred workflow.html

#### 470

# 471 **4.4 Evaluation of the BGC class abundance predictions in shallow** 472 metagenomes

We selected 30 merged pre-processed TARA Oceans metagenomes and randomly subsampled these to generate two sets of metagenomes, one with 100 million and the other with 5 million reads, using the seqtk v1.0 tool (https:// github.com/lh3/seqtk). We then annotated the BGC domains and predicted the BGC class abundances in this data using BiG-MEx (as described in sections 1 and 2), and compared the BGC class abundance predictions between the two sets of metagenomes.

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### 481 **5. BiG-MEx implementation**

### 482 **5.1 BGC domain identification module**

BiG-MEx BGC domain identification module uses the UProC 1.2.0<sup>16</sup> software 483 484 to classify short-read sequences using BGC domain references. To train 485 UProC for this purpose, we manually curated all amino acid sequences matching 150 antiSMASH hidden Markov model profiles (HMMs)<sup>10</sup>. In this 486 487 task, we removed sequences shorter than 25 amino acids and checked for the 488 presence of overlaps between sequences of different HMM profiles. In 489 addition, we categorized multi-domain proteins into multiple families. For the 490 training process, we included a set of negative control profiles to assess the 491 ratio of false positive hits. Namely, we used the t2fas, fabH, bt1fas, ft1fas 492 profiles negative controls for the PKS KS, t2ks, t2ks2, t2clf, as 493 Chal sti synt N, Chal sti synt C, hglD and hglE profiles. Once we curated 494 the amino acid sequence data, we applied the SEG(mentation) low complexity filter from the NCBI Blast+ 2.2 Suite<sup>44</sup> and created the UProC database. This 495 496 UProC database downloaded from be can 497 https://github.com/pereiramemo/BiG-MEx. Based on the identified reads 498 containing a BGC domain sequence, the module computes a count-based abundance profile of BGC domains. 499

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### 501 **5.2 BGC domain-based diversity analysis module**

502 This module performs two different analyses: Operational Domain Unit (ODU) 503 diversity estimation and phylogenetic placement of domain sequences. The 504 pipeline to estimate the ODU diversity, analyses each domain independently, 505 and consists of the following steps: 1) Short-read sequences, where the 506 domain being studied was identified, are recruited to perform a targeted 507 assembly metaSPAdes 3.11<sup>45</sup> with default parameters; 2) The Open Reading Frames (ORFs) in the resulting contigs are predicted with FragGeneScan-508 509 Plus; 3) Domain sequences are identified within the ORF amino acid 510 sequences with *hmmsearch* from HMMER v3 and extracted: 4) Domain amino acid sequences are clustered into ODUs using MMseqs2<sup>46</sup> with the cascaded 511 512 clustering option and the sensitivity parameter set to 7.5; 5) Annotated unassembled reads are mapped to the domain nucleotide sequences with 513 514 BWA-MEM 0.7.12, and the mean depth coverage is calculated using BEDtools 515 v2.23; 6) Based on this information, the coverage-based abundance of the 516 ODUs is computed and used to estimate an ODU alpha Shannon diversity. To 517 allow a comparison of the ODU diversity estimates between samples with 518 different sequencing depth, we include an option to estimate the diversity for 519 rarefied subsamples.

520 To perform the phylogenetic placement of domain sequences, we applied an approach similar to NaPDoS<sup>11</sup>. However, we extended the phylogenetic 521 522 placement analysis to 48 domains and included more comprehensive 523 reference trees, which are critical for the analysis of large metagenomic 524 samples. In detail, the phylogenetic placement consists of aligning the target 525 domain sequences to their corresponding reference multiple sequence 526 alignment (MSA) with MAFFT<sup>47</sup> (using --add option). Subsequently, the 527 extended MSA together with its reference tree are used as the input to run 528 pplacer<sup>48</sup> (with parameters: --keep-at-most 10 and --discard-nonoverlapped; all 529 other parameters set to default). pplacer performs the phylogenetic placement 530 using the maximum-likelihood criteria and outputs the extended tree in Newick and jplace formats<sup>49</sup>, and a table with statistics and information about the 531

532 placement of each sequence (i.e., likelihood, posterior probability, expected 533 distance between placement locations (EDPL), pendant length, and edge 534 number). To visualise the phylogenetic placement, a tree figure is generated using the ggtree R package<sup>50</sup>, where the coverage of the placed sequences is 535 536 mapped on their tree tips and used to scale a bubble representation. Besides 537 the phylogenetic placement, we included in this module an option to perform a 538 BLASTP search of the assembled domain sequences against the reference 539 domain sequences.

540 To construct the reference phylogenies, we first downloaded all the BGC amino acid sequences from the MIBiG database<sup>51</sup>. We identified the domain 541 542 sequences with hmmsearch using the BGC domain HMM profiles from 543 antiSMASH. Subsequently, we extracted and clustered these sequences with MMseqs2 to create a non-redundant dataset of amino acid sequences for 544 545 each domain. If the number of reference sequences identified in the MIBiG 546 database was greater than 500, we used a clustering threshold of 0.7 identity 547 at the amino acid level; otherwise, the threshold was set to 0.9; all other 548 parameters of MMseqs2 were set as specified previously. All domains with 549 less than 20 representative sequences were discarded. This resulted in a 550 subset of 48 domains that were considered for the phylogenetic 551 reconstructions. For each set of domain representative sequences, we 552 generated an MSA with MAFFT using the E-INS-I algorithm, removed sequence outliers with OD-seq<sup>52</sup> and constructed a phylogenetic tree with 553 RAXML<sup>53</sup>. To select the protein evolutionary model for the phylogenetic 554 555 reconstruction, we used the automatic model selection implemented in RAxML 556 with the maximum likelihood criterion. We used the GAMMA model of rate 557 heterogeneity and searched the tree space using the rapid hill-climbing 558 algorithm<sup>54</sup>, starting from a maximum parsimony tree. For the sake of 559 reproducibility, we specified a random seed number (i.e., -p 12345). Finally, we used RAxML to root the trees and compute the SH-like support scores<sup>55</sup>. In 560 561 Supplementary File 2, we provide for each domain phylogeny the number of 562 sequences and amino acid substitution model used, the mean, standard

563 deviation, maximum and minimum cophenetic distances between sequences,

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Faith's phylogenetic diversity<sup>24</sup> and the name of its corresponding BGC class.

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## 566 **5.3 BGC class abundance prediction module**

567 BiG-MEx uses machine-learning models to predict the abundance of the BGC 568 classes, based on the counts of annotated domains in unassembled 569 metagenomes. Each model is class-specific and was trained using the 570 abundance of the BGC class and its corresponding protein domains, as the 571 response and predictor variables, respectively. We used the classification 572 rules defined in antiSMASH for the annotation of BGC classes, to determine 573 the protein domains used as predictor variables in each model. To model the 574 abundance of a given BGC class, we implemented a two-step zero-inflated process. First, the presence or absence of the target BGC class is predicted 575 using a random forest (RF) binary classifier<sup>56</sup>. Second, a multiple linear 576 577 regression (MLR) is applied to predict the class abundance, but only if the 578 class was previously predicted as present. In the cases where the number of 579 zero values was lower than 10 or non-existent, we directly applied an MLR. 580 We trained the models using simulated metagenomic data (i.e., Marine-RM, 581 Human-Oral and Human-Stool datasets). The models predict a coverage-582 based abundance since this was the response variable used in the training 583 process. The RF binary classification models were created with the 584 randomForest function of the randomForest R package<sup>57</sup>, with the parameters 585 ntree set to 1000 (number of trees grown), nodesize set to 10 (minimum size 586 of terminal nodes), and mtry set to 1 (number of variables randomly sampled 587 as candidates at each split). For the MLR, we used the Im function of the stats 588 R package (https://www.R-project.org/) with default parameters.

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## 590 Code availability

591 BiG-MEx is freely distributed using Docker container technology 592 (www.docker.com), under the GNU General Public License v3.0. It can be 593 downloaded from https://github.com/pereiramemo/BiG-MEx, where we also 594 provide thorough documentation. Currently, we provide BGC class abundance 595 models targeting the marine environment, four different human body sites, and 596 the genus Streptomyces. To help users create their own BGC class abundance 597 models and compute the predictions, we developed the R package bgcpred: 598 https://github.com/pereiramemo/bgcpred. bgcpred is integrated in BiG-MEx, and 599 is used to generate the BGC class abundance predictions.

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### 601 **Data availability**

In Supplementary file 1, we provide the GenBank and RefSeq assembly accessions for the genomes used to generate the simulated metagenomic datasets. We provide the BGC class and domain abundance tables, obtained from the simulated data, at https://github.com/pereiramemo/BiG-MEx/.

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## 620 **Competing interests**

621 The authors declare no competing interests.

622

## 623 Author contributions

624 EP-F, AF-G, PLB, and MHM conceived BiG-MEx's algorithms. EP-F developed

the tools, and analysed the data, and wrote the paper with contributions from all

authors. All authors reviewed and approved the manuscript.

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### 779 Figures

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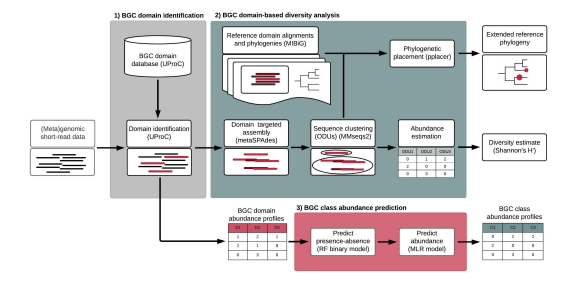
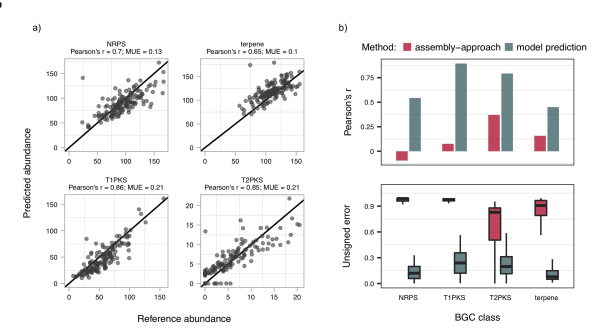


Fig. 1 | BiG-MEx analysis workflow. 1) BGC domain identification module. To annotate the BGC domains with UProC, we created an UProC database including 150 domains, which originate from 44 different BGC classes. This database was generated based on the amino acid sequences of antiSMASH hidden Markov model (HMM) profiles<sup>10</sup>. Using UProC output, this module generates a count-based abundance profile of BGC domains; 2) BGC domain-based diversity analysis module. Using the previously identified domains, this module performs a targeted assembly with metaSPAdes<sup>45</sup> to reconstruct the domain sequences. Assembled domain sequences are clustered into Operational Domain Units, and the number of ODUs and the coverage of the domain sequences within each ODU (used to approximate the abundance of the ODU) are used to compute the ODU alpha diversity. The environmental reconstructed domain sequences are placed onto reference phylogenetic trees with pplacer<sup>48</sup> (maximum likelihood criteria). In this module, we include pre-computed phylogenies for 48 domains, which are based on sequence data contained in the Minimum Information about a Biosynthetic Gene cluster (MIBiG)<sup>51</sup> database, allowing us to identify the relationships of query sequences with domains from pathways of known function; 3) BGC class abundance prediction module. The domain abundance profiles are used to predict the BGC class coverage-based abundance profiles using class-specific machine-learning models. These models consist of a two-step process: First, the presence/absence of the BGC class is predicted using a random forest (RF) classifier; Secondly, the abundance is predicted with a multiple linear regression (MLR) only if the class was previously predicted as present.

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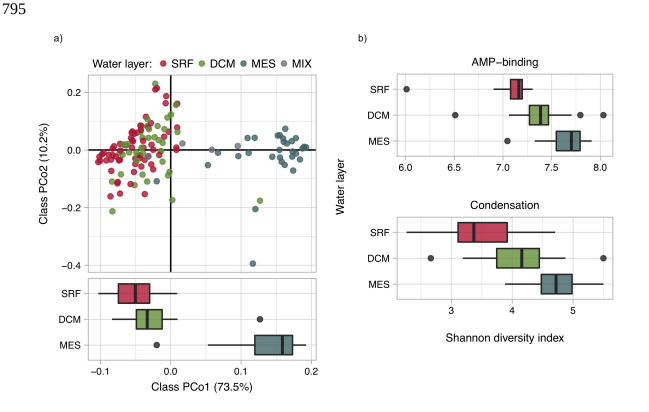
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**Fig. 2** | **Evaluating and benchmarking the BGC abundance prediction models.** (a) Scatter plots comparing the reference and predicted abundances of the NRPS, terpene, T1PKS and T2PKS BGC classes. MUE: Median Unsigned Error. The black, solid line represents the one-to-one relationship between the reference and predicted BGC class abundances. The BGC class abundance models were trained with the Marine-RM metagenomes and used to predict the abundances in the Marine-TM metagenomes. (b) Plots of the Pearson correlation coefficients (upper panel) and the unsigned error distributions (lower panel) of the BGC class abundances predicted by the models and estimated by the assembly approach, with respect to the reference abundances. In this comparison, we used 50 Marine-TM metagenomes. For the sake of clarity, 12 outlying unsigned error values (3% of the total comparisons) were excluded from the plot. The assembly approach consisted of the following tasks: 1) Assembling the metagenomes of the Marine-TM dataset; 2) Selecting the contigs with potential BGC sequences using BiG-MEx domain identification module; 3) Annotating the contigs with antiSMASH; 4) Mapping the short-read sequences to the identified BGC sequences; 5) Estimating the BGC class abundances.

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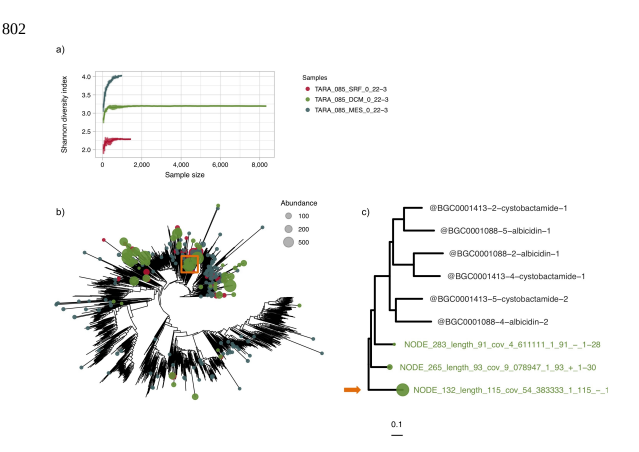
**Fig. 3 | BiG-MEx BGC class composition and domain-based diversity analysis in the TARA Oceans dataset. (a)** Principal Coordinates Analysis (PCoA) performed on a Bray-Curtis dissimilarity matrix of BGC class relative abundance profiles of the 139 prokaryotic metagenomes of TARA Oceans. BGC class abundance profiles were generated with BiG-MEx BGC class abundance module, using machine-learning models trained with the simulated Marine-RM metagenomic dataset. The abbreviations SRF, DCM, MES, and MIX correspond to surface, deep chlorophyll maximum, mesopelagic, and mixed epipelagic water layers, respectively. The boxplot in the bottom section of the panel shows the PCo1 value distributions for the metagenomes from the SRF, DCM and MES water layers. The PCo1 axis differentiated the MES water layer from the other two layers (Wilcoxon rank sum test; all p-values < 0.0001). **(b)** Bar plots showing the distribution of the ODU Shannon alpha diversity indices for the AMP-binding and condensation domains (NRPSs). The ODU diversity was computed for a match subset of 63 TARA Oceans metagenomes representing SRF, DCM, and MES water layers in 21 sampling stations. The AMP-binding and Condensation ODU diversity estimates were significantly different between the three water layers (pairwise Wilcoxon rank sum test; all p-values < 0.0001).

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**Fig. 4** | **BiG-MEx diversity analysis of condensation domains in three metagenomes from TARA Oceans sampling station TARA\_085 (a)** Rarefaction curves of the Shannon alpha diversity indices generated by BiG-MEx domain-based diversity analysis module, comparing the diversity of condensation ODUs in the metagenomes of the SRF, DCM, and MES water layers. Condensation domain sequences were clustered into ODUs using a 75% amino acid identity threshold. The diversity was computed using the number and abundance of distinct condensation ODUs. (b) Phylogenetic placement of the condensation ODU representative sequences, as performed by the BiG-MEx domain-based diversity analysis module. The SRF, DCM and MES had a phylogenetic diversity (Faith's PD)<sup>24</sup> of 58.15, 114.98 and 304.88, respectively. The size and colour of the bubbles on the leaves represent the ODU abundance and sample origin, respectively. (c) Detail of the clade contained in the orange, hollow square in (c), including the most abundant ODU (obtained in the TARA\_085\_DCM\_0\_22-3 sample; indicated with an orange arrow).

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