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3	Free-living Bacterial Communities Are Mostly Dominated by Oligotrophs
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### 10 Abstract

11 In response to resource availability, bacteria have evolved two distinct ecological strategies. 12 Copiotrophic bacteria grow fast and are heavily favored by selection where the resource is 13 abundant. In contrast, oligotrophic bacteria grow slowly but more efficiently and are highly 14 adaptive in nutrient-poor environments (Koch, 2001). Although oligotrophs and copiotrophs 15 are ubiquitous, except for a few well-characterized environments like the open ocean and 16 animal gut, the relative abundance of oligotrophic and copiotrophic bacteria and their 17 importance in the global ecosystem are still unclear. In addition, although several studies 18 have demonstrated the impact of nutrient availability on the bacterial community structure 19 under experimental conditions (Klappenbach et al., 2000, Nemergut et al., 2016), the role of 20 nutrients in shaping the structures of bacterial communities in their natural habitats remains 21 largely unknown. Using the ribosomal RNA operon (rrn) copy number to capture the 22 bacterial ecological strategy, we analyzed 44,045 samples from two large bacterial 23 community repositories that cover 78 environmental types. Here we show that animal-24 associated microbiota are dominated by copiotrophs while plant-associated and free-living 25 bacterial communities are mostly dominated by oligotrophs. Our results suggest that nutrient 26 availability plays an important role in determining the structure and ecological strategy of 27 bacterial communities in nature. We demonstrate that the average and distribution of rrn 28 copy number are simple yet robust predictors of the ecological strategy of bacterial 29 communities that can be applied to all sequence-based microbial surveys to link the 30 community structure and function. 31

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### 34 Introduction

35 The fundamental trade-off between the growth rate and efficiency in bacteria leads to 36 genomic features that distinguish between copiotrophs and oligotrophs (Lauro et al., 2009; 37 Roller et al., 2016). These features include genome size (Roller et al., 2016), codon usage 38 bias (Vieira-Silva & Rocha, 2010), transporter gene diversity (Lauro et al., 2009), motility 39 (Lauro et al., 2009; Roller et al., 2016) and etc. Among them, the copy number of ribosomal 40 RNA operon (rrn) is the best studied genomic trait that predicts the ecological strategy of 41 bacterial species. The *rrn* copy number positively correlates with cellular ribosomal content 42 and the maximum growth rate (Roller et al., 2016; Vieira-Silva & Rocha, 2010), and deletion 43 of rrn reduces the maximum growth rate of Escherichia coli and Bacillus subtilis (Nanamiya 44 et al., 2010; Stevenson & Schmidt, 2004; Yano et al., 2013). On the other hand, the rrn copy 45 number negatively correlates with the growth efficiency. One study has shown that the 46 carbon use efficiency and protein yield (protein synthesized per unit O<sub>2</sub> consumed) 47 decreases when the *rrn* copy number increases in several strains of bacteria (Roller et al., 48 2016). As such, high *rrn* copy number indicates a copiotrophic strategy while low *rrn* copy 49 number predicts an oligotrophic one. 50 As bacteria adopt different ecological strategies, the rm copy number in their genomes 51 varies from 1 to as many as 15 (Klappenbach et al., 2000). Bacteria have the potential to 52 rapidly change their rrn copy numbers in adaptation to the growth condition, as mutant 53 strains of *B. subtilis* with only one copy of *rrn* increased their *rrn* copies through operon 54 duplication within generations (Yano et al., 2013). Despite the potential for rapid change, 55 there is clearly phylogenetic signal in bacterial rrn copy number, as closely related species 56 tend to have similar *rrn* copy numbers (Kembel et al., 2012). The existence of such 57 phylogenetic signal suggests that *rrn* copy number, albeit labile, is under strong natural 58 selection. Although rrn copy number cannot be directly measured for environmental bacteria, 59 the phylogenetic signal in *rrn* copy number makes it possible to estimate the copy number 60 from the 16S rRNA gene sequence alone (Kembel et al., 2012). And such method is

61 applicable to both 16S rRNA amplicon and shotgun metagenomic sequence data, making

62 prediction of ecological strategies of individual bacterial species and the whole community

63 relatively simple and straightforward.

64 Results

First we set out to identify an optimal *rrn* copy number that can be used to classify

66 oligotrophs and copiotrophs. We used 63 representatives of oligotrophs and copiotrophs

67 classified by Lauro et al (Lauro et al., 2009) as our training set. As expected, oligotrophs and

68 copiotrophs are well separated in their *rrn* copy number distribution (Figure 1A). The area

69 under the curve (AUC) of the receiver operating characteristic (ROC) curve is 0.86 (Figure

1B), indicating that classification by *rrn* copy number has high accuracy (Šimundić, 2008).

71 Maximizing Youden's J statistic, we found that 2 was the optimal cutoff to distinguish

oligotrophs and copiotrophs, with a true positive rate of 0.789 and a false positive rate 0.200

for identifying oligotroph. Using this cutoff, we classified bacterial species (Operational

74 Taxonomic Units or OTUs) as oligotrophs if they had 1 or 2 rrn copies or as copiotrophs if

75 they had 3 or more *rrn* copies.

76 The free-living bacterial community in open ocean is known to be dominated by oligotrophic

bacteria (Lauro et al., 2009; Yooseph et al., 2010). Therefore, we tested whether the

78 average copy number (ACN) and distribution of *rrn* captured the ecological strategy of

79 marine bacteria in the open ocean using the shotgun metagenomic sequencing data from

80 the Tara Oceans Expedition (TARA) (Pesant et al., 2015). The ACN in surface water

81 samples from TARA ranged from 1.3 to 1.9, with a median of 1.4 (Supplementary Figure

1A). As expected, the community is dominated by oligotrophic bacteria with 1 or 2 *rrn* copies.

83 Overall, 93% of the bacterial cells in the community were oligotrophic, with 66% of the

84 bacteria cells had only one copy of *rrn* and 27% of them had 2 copies (Supplementary

85 Figure 1B).

86 Next we examined the ACN of bacterial communities from a broader range of environments 87 present in the EBI Metagenomics dataset (Mitchell et al., 2018). In total, we analyzed 2,528 88 whole genome shotgun (WGS) metagenomic samples that covered 6 environmental 89 categories (freshwater, marine, non-marine saline, soil, animal and human-associated) and 90 21 environmental types (Supplementary Table 1). We observed that ACN varied 91 substantially between environmental categories (Figure 2A). Animal and human-associated 92 bacterial communities had the highest ACN (median 4.4 and 4.8 respectively), while those 93 from marine, non-marine saline and soil environments had the lowest ACN (median 1.8, 2.0 94 and 2.0, respectively). Freshwater bacteria communities had intermediate ACN (median 95 3.1).

96 As the majority of microbial surveys are based on 16S rRNA amplicon sequencing, we 97 investigated whether the ACNs estimated from amplicon sequencing were consistent with 98 those estimated from WGS sequence data, out of concern that PCR bias associated with 99 amplicon sequencing (Acinas et al., 2005) can potentially skew the ACN. From the EBI 100 Metagenomics database, we identified 275 animal and human-associated microbiomes that 101 were surveyed by both methods. We compared the ACNs estimated from the WGS and 102 amplicon sequences of these 275 samples and found that they were highly correlated 103 (Figure 3). The r-squared of the one-to-one fit between the two ACNs was 0.708 and the 104 difference in ACNs estimated by the two methods was negligible (1.2% difference on 105 average).

Because ACN estimated from 16S rRNA amplicon data was reliable, we examined the ACN of bacterial communities surveyed by amplicon sequencing in both the EBI Metagenomics and Earth Microbiome Project (EMP) database (Thompson et al., 2017). In total, we analyzed 41,517 amplicon-sequenced samples from the two databases, which were 16 times larger than the WGS metagenomic data and together covered 11 environmental categories and 74 environmental types (Supplementary Table 1). ACN from the amplicon data varied between environment categories and showed a distribution pattern similar to

113 what had been observed in the EBI WGS metagenomic data (Figure 2B) and a previous 114 report (Thompson et al., 2017). Animal and human-associated bacterial communities had 115 highest ACN (median 4.5 and 4.6 respectively), followed by air and indoor surface 116 communities with intermediate ACN (median 3.7 and 3.1 respectively). Bacterial 117 communities in the other environments (freshwater, marine, non-marine saline, soil, 118 sediment, biofilm and plant-associated) all had low ACN (median less than 2.5). 119 We then examined the distribution of the *rrn* copy number within each of the 74 120 environmental types to directly assess the relative abundance of oligotrophs and copiotrophs 121 in each type of environment. Figure 2C shows that animal and human-associated bacterial 122 communities are dominated by copiotrophs (with 3 or more rrn copies) while free-living 123 bacterial communities are dominated by oligotrophs (with 1 or 2 rrn copies). Plant-associated 124 bacterial communities show a distribution of rrn copy number similar to those of the free-125 living ones, dominated by oligotrophs. We classified bacterial communities in which the 126 relative abundance of oligotrophs was greater than 60% as oligotrophic, and communities in 127 which the relative abundance of copiotrophs was greater than 60% as copiotrophic, and the 128 rest of communities as intermediate. We found that 91.5% of 31,835 animal and human-129 associated bacterial communities were copiotrophic, while 72.6% of 6.215 free-living and 130 71.2% of 3,467 plant-associated microbiomes were oligotrophic (Figure 4).

### 131 Discussion

The *rm* copy number has been shown to be a useful predictor of the ecological strategy of individual bacterial species (Klappenbach et al., 2000; Lauro et al., 2009; Roller et al., 2016) and bacterial communities (Nemergut et al., 2016; Thompson et al., 2017). Here we extended this line of research and used the distribution of *rm* copy to reveal the existence of three types of bacterial communities in nature: those dominated by oligotrophs, copiotrophs or neither. Our analysis of a comprehensive collection of environmental samples indicates that most free-living and plant-associated bacterial communities are dominated by

139 oligotrophs while animal and human-associated communities are dominated by copiotrophs. 140 This pattern is consistent with our understanding of nutrient availability in the environment. In 141 general, soil and water are considered nutrient poor (Barber & Lynch, 1977; Kuznetsov et 142 al., 1979; Maeda et al., 2000; Ohta & Hattori, 1983) while animal-associated sites (gut, skin, 143 etc) are considered nutrient rich. Despite the vast compositional differences among the 144 communities analyzed in this study, the emergence of a global pattern in the relative 145 abundance of oligotroph and copiotroph suggests that nutrient plays a key role in shaping 146 the ecological strategy and the structure of the bacterial communities in nature. More 147 generally, our results suggest that in nature selection exerts strong influence on the 148 assembly of bacterial community compared to the other processes such as drift and 149 dispersal. Interestingly, although the vast majority of groundwater samples were dominated 150 by oligotrophs (90.9% of samples), only 9.5% of contaminated groundwater samples were 151 dominated by oligotrophs. Similarly, the relative abundance of oligotrophs also decreased in 152 the contaminated open ocean and soil. In addition, we did observe outliers in some 153 environments. For example, Antarctic soil supplemented with organic carbon (Van Horn et 154 al., 2014) had extremely high ACN (8.9) compared to untreated soil (3.1). Together, they 155 provide compelling evidence that nutrient dictates the ecological strategy and the structure of 156 the bacterial communities.

157 We have demonstrated that the average and distribution of *rrn* copy number are simple yet 158 robust predictors of the ecological strategy of bacterial communities. Unlike other imprinted 159 genomic features (Lauro et al., 2009) that require shotgun metagenomic sequences, rrn 160 copy number can be estimated from 16S rRNA gene sequences alone and therefore is 161 applicable to both shotgun metagenomic and 16S rRNA amplicon sequence data. As such, 162 the metrics of average and distribution of *rrn* copy number can be applied in all sequence-163 based microbial surveys. Additionally, as the average copy number of rrn is weighted by the 164 relative cell abundance, it will capture the overall ecological strategy of a community even at 165 relatively low sequencing depth. These features make the community rrn copy number an

166 extremely useful quantitative trait for studying the microbial ecosystem. By providing a link 167 between the community structure and function, it adds values to 16S rRNA sequence data 168 beyond simply quantifying species composition, and can help researchers generate 169 hypotheses on how communities assemble in response to nutrient availability and other 170 environmental factors. In addition, it can be used as proxies of the average growth rate and 171 carbon use efficiency of bacterial communities to improve the current global carbon cycling 172 models that have fixed values for these parameters, as has been suggested previously 173 (Roller et al., 2016).

#### 174 Methods

### 175 <u>Classifying oligotrophic and copiotrophic bacteria with the *rrn* copy number</u>

176 Using 43 genomic features, Lauro et al classified 126 strains of bacteria as either 177 oligotrophic or copiotrophic (Lauro et al., 2009). We used these strains as a training set to 178 identify an optimal *rrn* copy number cutoff for oligotroph/copiotroph classification. To reduce 179 overrepresentation of certain clades, we selected one representative species from each of 180 the 63 genera represented by the 126 strains and whose complete genome sequence was 181 available (Supplementary Table 2). To evaluate the performance of using *rrn* copy number 182 for oligotroph/copiotroph classification, we calculated the true-positive rate and the false-183 positive rate using each integer from 1 to 15 as the copy number cutoff, plotted the ROC 184 curve and calculated the AUC. To find the optimal cutoff, we computed Youden's J statistic 185 for each cutoff, and selected the copy number that had the maximum Youden's J statistic as 186 the optimal cutoff. We then classified bacterial species whose rrn copy number was greater 187 than the cutoff as copiotroph, and oligotroph otherwise.

### 188 Compilation of Data

189 We used data from the Tara Oceans Expedition (Pesant et al., 2015) to test if the *rrn* copy

190 number captured the oligotrophic ecological strategy of marine bacteria in the open ocean.

191 The TARA dataset includes 139 bacterial community profiles derived from WGS

192 metagenomic sequencing and focused specifically on the free-living bacterial communities in 193 the open ocean across the globe. We downloaded the OTU abundance table and the 194 environmental metadata from the TARA companion website (Sunagawa et al., 2015). We 195 also downloaded the SILVA 16S rRNA reference sequences that were used in the TARA 196 study to pick OTU at 97% similarity, and estimated their rrn copy numbers using the method 197 described below. We analyzed 63 samples from the surface layer. 198 To explore the global ecological strategies of bacterial communities in nature, we compiled 199 bacterial diversity survey data from two large microbial survey repositories, the EBI 200 Metagenomics (Mitchell et al., 2018) and the EMP (Thompson et al., 2017). For the EBI 201 Metagenomics dataset, we included 12,486 WGS metagenomic and 69,385 amplicon 202 sequencing runs processed by the version 2.0 or 3.0 pipeline of the EBI Metagenomics 203 (Mitchell et al., 2018). Sequencing runs of the same type (16S rRNA amplicon or WGS 204 metagenomics) that were associated with the same sample were combined together. For the 205 EMP dataset, we included 23,228 amplicon sequencing samples from its first release. We 206 downloaded OTU abundance table and the metadata associated with each sample. The 207 GreenGene 13.8 16S rRNA reference sequences were used for OTU picking at 97% 208 similarity in both repositories. The *rrn* copy numbers of GreenGene reference sequences 209 were estimated as described below. We filtered out WGS metagenomic runs whose 16S 210 rRNA reads exceeded 5% of the total sequence reads and amplicon runs if less than 95% of 211 the sequence reads were 16S rRNA reads, as these samples were likely 16S rRNA 212 amplicon sequencing data mislabeled as WGS metagenomic data or vice versa. We then 213 removed samples if less than 80% of their 16S rRNA reads could be mapped to the 214 GreenGene reference sequences or if the total mapped reads were fewer than 400. We 215 removed samples that only surveyed or were enriched for a specific bacterial clade. We also 216 removed samples whose environmental types were too general or mislabelled, and 217 environmental types that contained less than 5 samples. At the end, a total of 44,045 218 samples (2,528 WGS metagenomic and 41,517 amplicon sequencing) remained after the

219 quality filtering. Together, they cover a total of 78 environmental types. We grouped these

220 environmental types into 11 categories: air, indoor surface, sediment, freshwater, marine,

221 non-marine saline, biofilm, soil, plant-associated, animal-associated and human-associated

222 (Supplementary table 1).

### 223 Estimation of the average rrn copy number (ACN) of a community

224 The 16S rRNA gene copy number of each OTU in the SILVA and the GreenGene reference

225 databases was estimated using the phylogenetic ancestral state reconstruction method

described in Kembel et al (Kembel et al., 2012) with an updated reference tree of 1,197

227 bacterial species whose genomes were sequenced and 16S rRNA gene copy numbers were

known. The ACN of the bacterial community in a sample was weighted by the relative cell

abundance of OTUs, as shown in Equation (1),

230

$$ACN = \sum_{i} n_i C_i$$

(1)

(2)

231

where *n<sub>i</sub>* and *C<sub>i</sub>* are the relative cell abundance and 16S rRNA gene copy number,

respectively. The relative cell abundances  $n_i$  was in turn computed by Equation (2) proposed

in Kembel *et al* (Kembel et al., 2012).

235

$$n_i = \frac{\frac{g_i}{C_i}}{\sum_i \frac{g_i}{C_i}}$$

236

where  $g_i$  is the relative gene abundance of an OTU estimated by the read number.

### 238 Estimation of the distribution of *rrn* copy number in communities of an environment type

The relative cell abundance of bacteria with a certain *rrn* copy number was calculated by summing the relative cell abundance of all species in a community with that *rrn* copy number (rounded to the nearest integer). To generate the distribution plot for an environmental type (Figure 2C), we repeated the analysis for all samples (communities) of that environmental type and calculated the average relative cell abundance of bacterial species with a certain *rrn* copy number.

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# 322 Competing interests

323 The authors declare no competing financial interests.

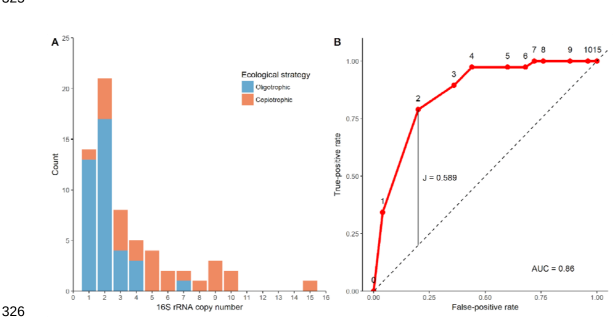
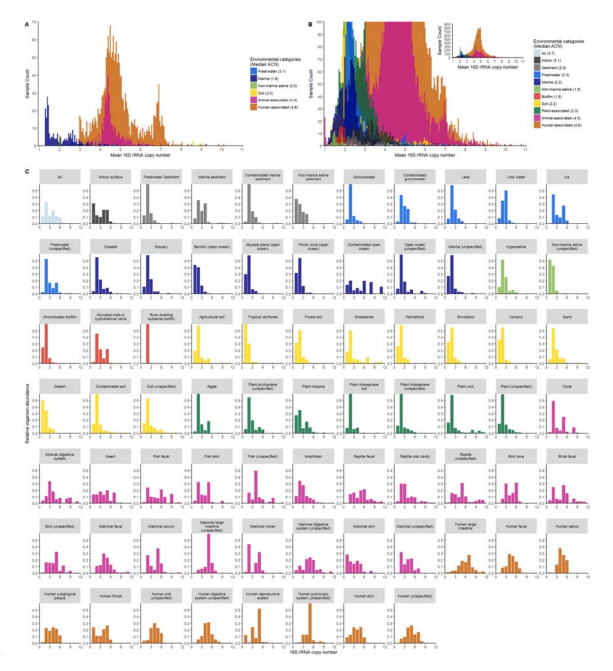




Figure 1. (A) The distribution of *rrn* copy number of the 63 representative species that were identified as either oligotrophic (blue) or copiotrophic (orange). (B) The ROC curve for using the *rrn* copy number to distinguish oligotrophic and copiotrophic bacteria, with an AUC of 0.86. The best copy number cutoff is 2 with a maximum Youden's J statistic of 0.589.

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Figure 2. Distribution of the ACN of 2,528 samples surveyed by WGS metagenomics (A) and 41,517 samples surveyed by 16S rRNA amplicon sequencing (B) in the EBI Metagenomics and the EMP databases. The relative abundance of bacterial species was plotted against their *rrn* copy number for the amplicon sequencing samples (C). Samples were colored by the environment categories.

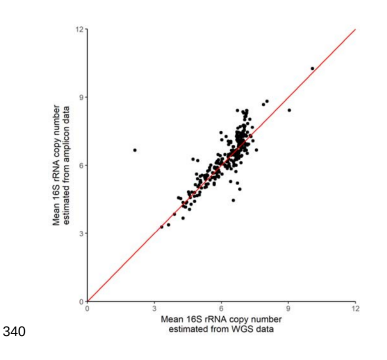
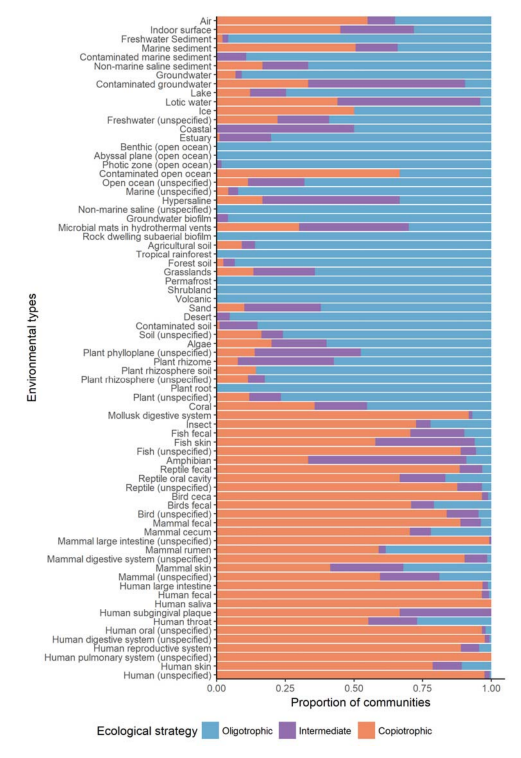




Figure 3. The ACN of 275 animal and human-associated bacterial communities surveyed by
both 16S rRNA amplicon and WGS metagenomic sequencing methods. Red line illustrates
the one-to-one fit between the ACN estimated from the two methods. The r-squared of the fit
is 0.708.





348 Figure 4. The proportion of 41,517 16S rRNA amplicon sequenced samples from 74

environmental types that are dominated (more than 60%) by oligotrophs (species with 1 or 2

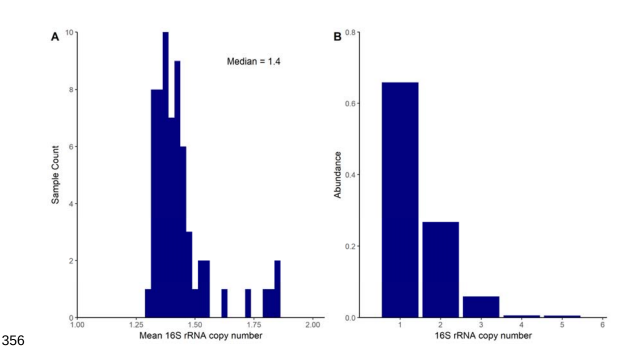
- 350 copies or rrn, orange) or by copiotrophs (species with 3 or more copies of rrn, blue), or are
- intermediate (neither group was dominant, purple).



### 353 Supplementary Data

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Supplementary Figure 1. (A) The average copy number of *rrn* copy of the 63 bacterial
communities from the surface layer in TARA. (B) The distribution of *rrn* copy number in the
bacterial communities. The abundances of species with more than 6 *rrn* copies are
negligible and not shown.

361

362 Supplementary Table 1. Environmental types covered by samples from WGS metagenomics 363 in the EBI Metagenomics (A) and 16S rRNA amplicon sequencing in the EBI Metagenomics 364 and the Earth Microbiome Project (B), and their statistics.

365 Supplementary Table 2. The *rm* copy number of 116 strains of oligotrophic or copiotrophic

366 bacteria whose complete genome sequences were available.