1	Iroki: automatic customization for phylogenetic trees
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#### 19 Abstract

#### 20 Background

21 Phylogenetic trees are an important analytical tool for examining species and community 22 diversity, and the evolutionary history of species. In the case of microorganisms, decreasing 23 sequencing costs have enabled researchers to generate ever-larger sequence datasets, which 24 in turn have begun to fill gaps in the evolutionary history of microbial groups. However, 25 phylogenetic analyses of large sequence datasets present challenges to extracting 26 meaningful trends from complex trees. Scientific inferences made by visual inspection of 27 phylogenetic trees can be simplified and enhanced by customizing various parts of the tree, 28 including label color, branch color, and other features. Yet, manual customization is time-29 consuming and error prone, and programs designed to assist in batch tree customization 30 often require programming experience. To address these limitations, we developed Iroki, a 31 program for fast, automatic customization of phylogenetic trees. Iroki allows the user to 32 incorporate information on a broad range of metadata for each experimental unit 33 represented in the tree.

34

#### 35 Results

36 Iroki was applied to four existing microbial sequence datasets to demonstrate its utility in 37 data exploration and presentation. Specifically, we used Iroki to highlight connections 38 between viral phylogeny and host taxonomy, explore the abundance of microbial groups 39 associated with Shiga toxin-producing *Escherichia coli* (STEC) in cattle, examine short-

- 40 term temporal dynamics of virioplankton communities, and to search for trends in the
- 41 biogeography of Zetaproteobacteria.
- 42
- 43 Conclusions
- 44 Iroki is an easy-to-use application having both command line and web-browser
- 45 implementations for fast, automatic customization of phylogenetic trees based on user-
- 46 provided categorical or continuous metadata. Iroki enables hypothesis testing through
- 47 improved visualization of phylogenetic trees, streamlining the process of biological sequence
- 48 data exploration and presentation.
- 49
- 50 Availability
- 51 Iroki can be accessed through a web browser application or via installation through
- 52 RubyGems, from source, or through the Iroki Docker image. All source code and
- 53 documentation is available under the GPLv3 license at https://github.com/mooreryan/iroki.
- 54 The Iroki web-app is accessible at www.iroki.net or through the VIROME portal
- 55 (http://virome.dbi.udel.edu), and its source code is released under GPLv3 license at
- 56 https://github.com/mooreryan/iroki\_web. The Docker image can be found here:
- 57 https://hub.docker.com/r/mooreryan/iroki.
- 58

## 59 Keywords

- 60 Phylogeny, visualization, sequence analysis, bioinformatics, metagenomics
- 61

## 62 Iroki: automatic customization for phylogenetic trees

63

#### 64 Background

65 Community and population ecological studies often use phylogenetic trees as a means for 66 assessing the diversity and evolutionary history of organisms. In the case of 67 microorganisms, the declining cost of sequencing has enabled researchers to gather ever-68 larger sequence datasets from unknown microbial populations within environmental 69 samples. While large sequence datasets have begun to fill in the gaps in the evolutionary 70 history of microbial groups [1–5]; they have also posed new analytical challenges as 71 extracting meaningful trends within such highly dimensional datasets can be cumbersome. 72 In particular, scientific inferences made by visual inspection of phylogenetic trees can be 73 simplified and enhanced by customizing various parts of the tree including label and branch 74 color, branch width, and other features. Though several tree visualization packages allow 75 for manual modifications [6–9], the process can be time consuming and error prone 76 especially when the tree contains many nodes. Moreover, these packages are typically not 77 capable of batch customization without prior computer programming experience [10-13]. 78

Iroki, a program for fast, automatic customization of phylogenetic trees, was developed to address these limitations and enable users to incorporate a broad array of metadata information for each experimental unit represented in the tree. Iroki is available for use through a web browser interface at <u>www.iroki.net</u>, through the VIROME portal (http://virome.dbi.udel.edu), and through a UNIX command line tool. Results are saved in

84 the widely used Nexus format with color metadata tailored for use with FigTree [8] (a freely
85 available and efficient tree viewer).

86

### 87 **Implementation**

88 Iroki enhances visualization of phylogenetic trees by coloring node labels and branches

89 according to categorical metadata criteria or numerical data such as abundance

90 information. Iroki can also rename nodes in a batch process according to user specifications

91 so that node names are more descriptive. A tree file in Newick format containing a

92 phylogenetic tree is always required. Additional required input files depend on the

93 operation(s) desired. Coloring functions require a color map or a biom [14] file. Node

94 renaming functions require a name map. The color map, name map, and biom files are

95 created by the user and, along with the Newick file, form the inputs for Iroki.

96

#### 97 Explicit tree coloring

98 Iroki's principle functionality involves coloring node labels and/or branches based on 99 information provided by the user in the color map. The color map text file contains either 100 two or three tab-delimited columns depending on how branches and labels are to be 101 colored. Two columns, pattern and color, are used when labels and branches are to have 102 the same color. Three columns, pattern, label color, and branch color, are used when 103 branches and labels are to have different colors. Patterns are searched against node labels 104 either as regular expressions or exact string matches.

106 Entries in the color column can be any of the 657 named colors in the R programming 107 language [15] (e.g., skyblue, tomato, goldenrod2, lightgray, black) or any valid hexadecimal 108 color code (e.g., #FF78F6). In addition, Iroki provides a 19 color palette with 109 complementary colors based on Kelly's color scheme for maximum contrast [16]. Nodes on 110 the tree that are not in the color map will remain black. 111 112 Depending on user-specified options, a pattern match to node label(s) will trigger coloring of 113 the label and/or the branch directly connected to that label. Inner branches will be colored 114 to match their descendent branches if all descendants are the same color, allowing quick 115 identification of common ancestors and clades that share common metadata. 116 117 Tree coloring based on numerical data 118 Iroki provides the ability to generate color gradients based on numerical data, such as 119 absolute or relative abundance, from a tab-delimited biom format file [14]. Single-color 120 gradients use color saturation to illustrate numerical differences, with nodes at a higher level 121 being more saturated than those at a lower level. For example, highly abundant nodes will 122 be represented by more highly saturated colors. Two-color gradients show numerical 123 differences through both color mixing and luminosity. Additionally, the biom file may 124 specify numerical information for one group (e.g., abundance in a particular sample) or for 125 two groups (e.g., abundance in the treatment group vs. abundance in the control group). 126 For biom files with one group, single- or two-color gradients may be used. However, biom

127 files specifying two-group metadata may only use the two-color gradient.

128

## 129 Renaming nodes

130	Some packages	for generating phylogenetic t	trees restrict the use of special characters and

- 131 spaces or require node names to be shorter than a specified length or (RAxML [17],
- 132 PHYLIP [18], etc.). Name restrictions present challenges to scientific interpretation of
- 133 phylogenetic trees. Iroki's renaming function uses a two-column, tab-delimited name map
- 134 to associate current node names, exactly matching those in the tree file, with new names.
- 135 The new name column has no restrictions on name length or character type. Iroki ensures
- 136 name uniqueness by appending integers to the ends of names, if necessary.

137

138 Combining the color map, name map, and biom files

139 Iroki can be used to make complex combinations of customizations by combining the color

140 map, name map, and biom files. For example, a biom file can be used to apply a color

141 gradient based on numerical data to the labels of a tree, a color map can be used to

separately color the branches based on user-specified conditions, and a name map can be

- 143 used to rename nodes in a single command or web request. Iroki follows a specific order of
- 144 precedence when applying multiple customizations. The color gradient inferred from the
- 145 biom file is applied first. Next, the color map is applied to specified labels or branches,
- 146 overriding the gradient applied in the previous step, if necessary. Finally, the name map is
- 147 used to map current names to the new names (Fig. 1).

148

149 Output

- 150 Iroki outputs the modified tree in the Nexus format. When building the phylogenetic tree,
- 151 FigTree uses the Nexus format file and interprets the color metadata output of Iroki.

152

## 153 **Results & Discussion**

- 154
- 155 Global diversity of bacteriophage

156 Viruses are the most abundant biological entity on Earth, providing an enormous reservoir 157 of genetic diversity, driving evolution of their hosts, influencing composition of microbial 158 communities, and affecting global biogeochemical cycles [19,20]. The viral taxonomic 159 system developed by the International Committee on Taxonomy of Viruses (ICTV) is based 160 on a suite of physical characteristics of the virion rather than on genome sequences. Noting 161 this limitation, the phage proteomic tree was created to provide a genome-based taxonomic 162 system for bacteriophage classification [21]. The phage proteomic tree was recently 163 updated to include hundreds of new phage genomes from the Phage SEED reference 164 database [22], as well as long assembled contigs from viral shotgun metagenomes (viromes) 165 collected from the Chesapeake Bay (SERC sample) [23] and the Mediterranean Sea [24]. 166

167 Taxonomy and host information metadata was collected for the viral genome sequences, a 168 color map was created to assign colors based on viral family and host phyla, and Iroki was 169 used to add color metadata to branches and labels of the phage proteomic tree. Since a 170 large number of colors were required on the tree, Iroki's Kelly color palette was used to 171 provide clear color contrasts. The tree was rendered with FigTree (Fig. 2).

172

173	Adding color to the phage proteomic tree with Iroki shows trends in the data that would be
174	difficult to discem without color. Uncultured phage contigs from the Chesapeake Bay and
175	Mediterranean viromes make up a large portion of all phage sequences shown on the tree,
176	and are widely distributed among known phage. In general, viruses in the same family
177	claded together, e.g., branch coloring highlights large groups of closely related Siphoviridae
178	and Myoviridae. This label-coloring scheme also shows that viruses infecting hosts within
179	same phylum are, in general, phylogenetically similar. For example, viruses within one of
180	the multiple large groups of Siphoviridae across the tree infect almost exclusively host
181	species within the same phylum, e.g., Siphoviridae infecting Actinobacteria clade away from
182	Siphoviridae infecting Firmicutes or Proteobacteria.
183	
184	Bacterial community diversity and prevalence of E. coli in beef cattle
185	Shiga toxin-producing Escherichia coli (STEC) are dangerous human pathogens that
186	colonize the lower gastrointestinal tracts of cattle and other ruminants. STEC-contaminated

187 beef and STEC shed in the feces of these animals are major sources of foodborne illness.

188 To identify possible interactions between STEC populations and the commensal cattle

189 microbiome, a recent study examined the diversity of the bacterial community associated

190 with beef cattle hide [25]. Fecal and hide samples were collected over twelve weeks and

191 SSU rRNA amplicon libraries were constructed and analyzed by Illumina sequencing [26].

192 The study indicated that the community structure of hide bacterial communities was altered

193 when the hides were positive for STEC contamination.

194

195	Iroki was used to visualize changes in the relative abundance of each cattle hide bacterial
196	OTU according to the presence or absence of STEC. A Mann-Whitney U test comparing
197	OTU abundance between STEC positive and STEC negative samples was performed, and
198	those bacterial OTUs showing a significant change in relative abundance (p < $0.5$ ) were
199	placed on a phylogenetic tree according to the 16S rRNA sequence. Branches of the tree
200	were colored based on whether there was a significant change in relative abundance with
201	STEC contamination (red: $p < 0.1$ , blue: $p > = 0.1$ ). Node labels were colored along a
202	blue-green color gradient representing the abundance ratio of OTUs between samples with
203	STEC (blue) and without (green). Additionally, label luminosity was determined based on
204	overall abundance of each OTU (lighter: less abundant, darker: more abundant) (Fig. 3).
205	Iroki makes it clear that most OTUs on the tree showed a significant difference in
206	abundance (branch coloring) between STEC positive and STEC negative samples (node
207	coloring). Furthermore, we can see that most OTUs are at low abundance with only a few
208	highly abundant OTUs (label luminosity). The color gradient added by Iroki allows us to
209	see that the abundant OTUs were evolutionarily distant from one another and thus spread
210	out across many phylogenetic groups.

211

Iroki can be used to quickly test hypotheses without investing a large amount of time
annotating trees manually. A UPGMA tree was created based on unweighted UniFrac
distance [27] between 356 bacterial community profiles based on SSU rRNA amplicon
sequences from cattle hide and fecal samples (Fig. 4). Iroki was used to evaluate similarities

216 in sample bacterial communities according to the sampling location. Iroki colored branches 217 based on whether the sample originated from feces (blue) or from hide (red). The coloring 218 added by Iroki shows a clear partitioning of bacterial communities on the tree based on their 219 sampling location (hide or feces). However, four fecal samples grouped with hide samples. 220 and two hide samples grouped with fecal samples, highlighting the ability of Iroki to easily 221 identify good candidates for more in-depth examination. Additionally, Iroki was used to 222 illustrate a correlation between one of the most abundant bacterial families, 223 Ruminococcaceae, and sampling location. Iroki colored node labels with a color gradient 224 based on Ruminococcaceae family abundance, utilizing both a single color gradient (Fig. 225 4A) and a two color gradient (Fig. 4B). Custom trees were visualized using FigTree. Iroki's 226 automatic color gradient and ability to label branches and nodes based on different criteria 227 clearly show that Ruminococcaceae is more abundant in fecal samples than in hide 228 samples.

229

230 Short-term dynamics of virioplankton

The gene encoding Ribonuclotide reductase (RNR) is common within viral genomes and thus can be used as a marker gene for studying viral diversity [23]. Moreover, RNR polymorphism is predictive of some of the biological and ecological features of viral populations [28]. A mesocosm experiment examined the short-term dynamics of phage populations using RNR amplicon sequences, specifically, sequences of class II RNRs of bacteriophages infecting cyanobacterial hosts. A phylogenetic tree was created from the Cyano II RNR amplicon sequences and Iroki was used to color nodes and branches based

238	on the time point (0 h, 6 h, $12$ h) at which each amplicon sequence was observed. The
239	customized tree was then visualized using FigTree (Fig. 5). Iroki's coloring showed that no
240	phylogenetic clade was dominated by OTUs observed in any particular time point; rather,
241	time points were spread relatively evenly across clades. This analysis demonstrates Iroki's
242	utility for exploring sequence datasets, allowing the researcher to quickly and easily test
243	hypotheses.

244

## 245 Phylogeny of Zetaproteobacteria within a biogeographic context

246 Biogeographical studies assess the distribution of an organism's biodiversity across space 247 and time. The extent to which microorganisms exhibit geographic distribution patterns is an 248 open question in microbial ecology. The isolated nature of the microbial communities 249 associated with deep-ocean hydrothermal vents provides an ideal system for studying the 250 biogeography of microbes. In particular, iron-oxidizing bacteria have been shown to thrive 251 in vent fluids, sediments, and iron-rich microbial mats associated with the vents. Globally, 252 iron-oxidizing bacteria make significant contributions to the iron and carbon cycles. A 253 recent study analyzed multiple SSU rRNA clone libraries to investigate the biogeography of 254 Zetaproteobacteria, a phyla containing many iron-oxidizing bacterial species, between three 255 sampling regions of the Pacific Ocean (central Pacific—Loihi seamount, western Pacific— 256 Southern Mariana Trough, and southern Pacific (Vailulu'u Seamount/Tonga Arc/East Lau 257 Spreading Center/Kermadec Arc) [29]. Sequences were aligned and a phylogeny was 258 inferred as described in [29]. Iroki was used to examine the relationship between sampling 259 location and phylotype by adding branch and label color based on geographic location and

260	renaming original node labels with OTU and location metadata. The custom tree was
261	visualized using FigTree (Fig. 6). In some cases, OTUs contained sequences from only one
262	sampling location (e.g., OTUs 12, 15, and 16), whereas other OTUs are distributed among
263	more than one sampling location (e.g., OTUs 1, 2, and 4). Often, sequences sampled from
264	the same geographic location are in the same phylotype despite being members of different
265	OTUs (e.g., OTUs 10 and 19).
266	
267	Availability and requirements
268	A web browser version of Iroki can be accessed online at <u>www.iroki.net</u> or through the
269	VIROME portal (http://virome.dbi.udel.edu/). For users who wish to run Iroki locally, a
270	command line version of the program is installable via RubyGems, from GitHub
271	( <u>https://github.com/mooreryan/iroki</u> ). A Docker image is available for users who desire the
272	flexibility of the command line tool, but do not want to install Iroki or manage its
273	dependencies ( <u>https://hub.docker.com/r/mooreryan/iroki</u> ). Docker is a popular software
274	container platform that allows bundling of an application with its dependencies in a
275	portable, self-contained system [30,31]. The README file, accompanying the source code,
276	provides detailed instructions for setting up and running Iroki. Further documentation and
277	tutorials can be found at the Iroki Wiki (https://github.com/mooreryan/iroki/wiki).
278	
279	License
280	Iroki and its associated programs are released under the GNU General Public License

281 version 3 [32].

282

# 283 Conclusions

284	Iroki is a command line program and web browser application for fast, automatic
285	customization of large phylogenetic trees based on user specified configuration files
286	describing categorical or continuous metadata information. The output files include Nexus
287	tree files with color metadata tailored specifically for use with FigTree. Various example
288	datasets from microbial ecology studies were analyzed to demonstrate Iroki's utility. In each
289	case, Iroki simplified the processes of data exploration, data presentation, and hypothesis
290	testing. Though these examples focused specifically on applications in microbial ecology,
291	Iroki is applicable to any problem space with hierarchical data that can be represented in
292	the Newick tree format. Iroki provides a simple and convenient way to rapidly customize
293	trees, especially in cases where the tree in question is too large to annotate manually or in
294	studies with many trees to annotate.
295	

- 296 List of Abbreviations
- 297 OTU: operational taxonomic
- 298 RNR: Ribonucleotide reductase
- 299 STEC: Shiga-toxengenic Escherichia coli

- 301 Ethics approval and consent to participate
- 302 Not applicable
- 303

## 304 **Consent for publication**

- 305 Not applicable
- 306

## 307 Availability of data and materials

- 308 Data and code used to generate figures are available on GitHub at
- 309 https://github.com/mooreryan/iroki\_manuscript\_data
- 310

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- 315

### **Competing Interests**

- 317 The authors declare that they have no competing interests.
- 318

## 319 Authors' contributions

320 RMM and SMM conceived the project. RMM wrote the manuscript and implemented Iroki.

- 321 AOH and RLM processed and analyzed Cyano II amplicons. All authors read, edited, and
- 322 approved the final manuscript.

323

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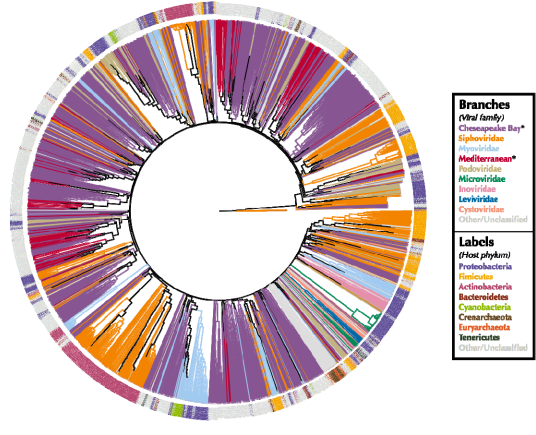
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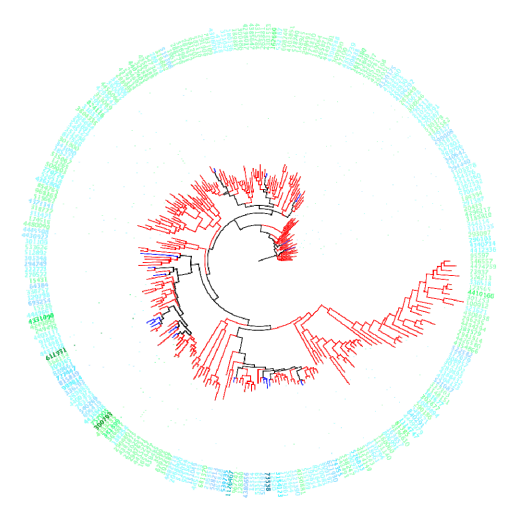


402

0.9

# 403 **Fig. 2: Comparing phage and their host phyla**

All phage genomes from Phage SEED with assembled virome contigs from the Chesapeake
Bay and Mediterranean Sea. Iroki highlights phylogenetic trends after coloring branches
according to viral family or sampling location in the case of virome contigs (marked with an
asterisk in the legend), and coloring node labels according to host phylum of the phage.



409

0.1

### 410 Fig. 3: Changes in OTU abundance in two sample groups

411 Approximate-maximum likelihood tree of OTUs that showed significant differences in

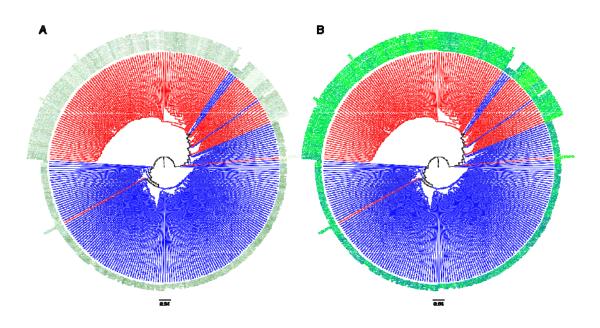
412 relative abundance between STEC positive and STEC negative cattle hide samples.

413 Branches show significance based on coloring by the p-value of a Mann-Whitney U test

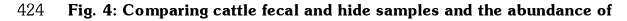
414 examining changes in abundance between samples positive for STEC (p < 0.1 - red) and

- 415 samples negative for STEC, (p > = 0.1 blue). Label color on a blue-green color gradient
- 416 highlights OTU occurrence based on the abundance ratio between STEC positive samples
- 417 (green) and STEC negative samples (blue). Labels that are darker green had a higher

- 418 abundance in STEC positive samples, and a lower abundance in STEC negative samples.
- 419 For example, OTU 300793 (bottom left corner) is darker than most (indicating high overall
- 420 abundance) and more green than blue (indicating higher abundance in STEC positive
- 421 samples than in STEC negative samples). Node luminosity represents overall abundance
- 422 with lighter nodes being less abundant than darker nodes.

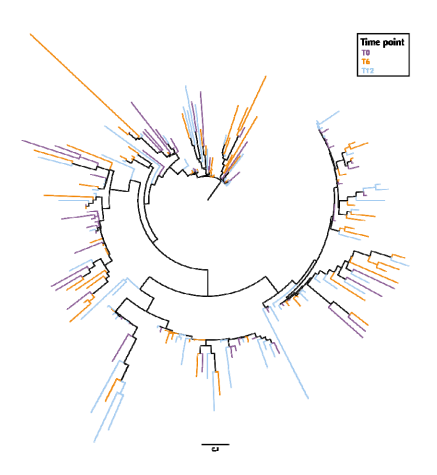






### 425 **Ruminococcaceae**

426 Phylogeny based on UPGMA tree of pairwise unweighted UniFrac distance between 356 427 bacterial community profiles based on SSU rRNA amplicon sequences from cattle hide and 428 feces. Branches are colored by feces (blue) and hide (red). Rapid testing of the hypothesis 429 that the abundance of one of the most abundant families, Ruminococcaceae, and sample 430 origin are correlated is enabled through node label coloring by (A) a green single-color 431 gradient (color saturation increases with increasing abundance of Ruminococcaceae OTUs) 432 and (B) a light green (low abundance of Ruminococcaceae OTUs) to dark blue (high 433 abundance of Ruminococcaceae OTUs) color gradient.

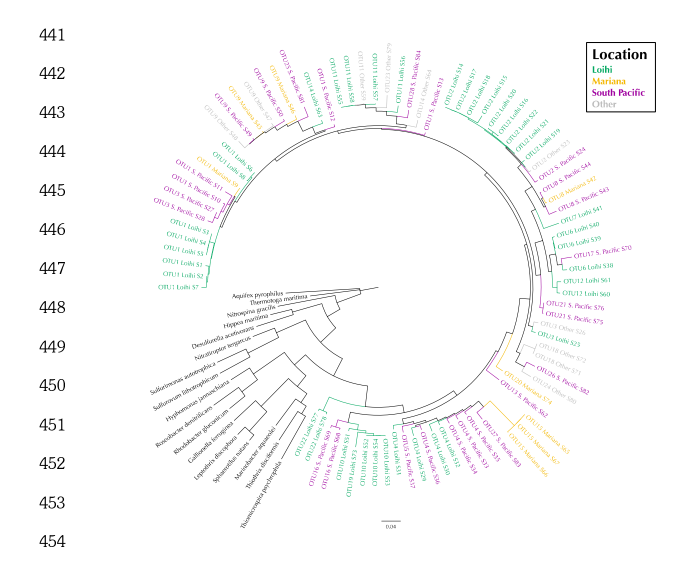


# 435

# 436 Fig. 5: Temporal dynamics of virioplankton populations according to Cyano II

## 437 **RNR amplicon phylogeny**

- 438 An approximately-maximum-likelihood phylogenetic tree of 200 randomly selected class II
- 439 Cyano RNR representative sequences from 98% percent clusters. Iroki was used to color
- 440 branches by time point: zero hours purple, six hours orange, and twelve hours blue.



#### 455 Fig. 6: Zetaproteobacteria show biogeographic partitioning

Phylogenetic tree showing placement of 84 full-length Zetaproteobacteria SSU rRNA
sequences collected from three Pacific Ocean locations and 17 reference sequences. Iroki
was used to color labels and branches by geographic location of the sampling site (Loihi –
green, Mariana – gold, South Pacific – purple, and Other – gray), as well as to rename the
nodes with OTU and sampling site metadata. Known reference Zetaproteobacterial species
are shown in black.