# 1 Intraspecies genomic divergence of coral algal symbionts shaped

# 2 by gene duplication

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# 9 Abstract

10 Dinoflagellates of Order Suessiales include the diverse Family Symbiodiniaceae known for 11 their role as essential coral reef symbionts, and the cold-adapted Polarella glacialis. These 12 taxa inhabit a broad range of ecological niches and exhibit extensive genomic divergence, 13 although their genomes are in the smaller size ranges (haploid size < 3 Gbp) compared to 14 most other dinoflagellates. Different isolates of a species are known to form symbiosis with 15 distinct hosts and exhibit different regimes of gene expression, but intraspecies whole-16 genome divergence remains little known. Focusing on three Symbiodiniaceae species (the 17 free-living Effrenium voratum, and the symbiotic Symbiodinium microadriaticum and 18 Durusdinium trenchii) and the free-living outgroup P. glacialis, all for which whole-genome 19 data from multiple isolates are available, we assessed intraspecies genomic divergence at 20 sequence and structural levels. Our analysis based on alignment and alignment-free methods 21 revealed greater extent of intraspecies sequence divergence in symbiodiniacean species than 22 in P. glacialis. Our results also reveal the implications of gene duplication in generating 23 functional innovation and diversification of Symbiodiniaceae, particularly in D. trenchii for 24 which whole-genome duplication was involved. Interestingly, tandem duplication of single-25 exon genes was found to be more prevalent in genomes of free-living species than in those of 26 symbiotic species. These results in combination demonstrate the remarkable intraspecies 27 genomic divergence in dinoflagellates under the constraint of reduced genome sizes, shaped 28 by genetic duplications and symbiogenesis events during diversification of Symbiodiniaceae.

# 29 Introduction

30 Dinoflagellates of the Order Suessiales include the Family Symbiodiniaceae, which

- 31 predominantly consists of symbiotic lineages essential to coral reef organisms.
- 32 Symbiodiniaceae taxa collectively exhibit a broad spectrum of symbiotic associations (i.e.,

33 facultativeness) and variable degrees of host specificity (i.e., host-specialist vs host-

34 generalist), although some are described as solely free-living (Thornhill et al. 2014;

35 LaJeunesse et al. 2018). A comparative analysis of whole-genome sequences from 15 taxa

36 revealed extensive sequence and structural divergence among Symbiodiniaceae taxa, which

37 was more prevalent in isolates of the symbiotic species, *Symbiodinium microadriaticum* 

38 (González-Pech et al. 2021). This was supported by a metagenomics survey of single-

39 nucleotide polymorphisms in the genomes of symbiotic *Symbiodinium fitti* from different

40 coral taxa and biogeographical origins, revealing intraspecies sequence divergence correlated

41 to coral host taxa (Reich et al. 2021).

42 A recent comparative genomic analysis incorporating genomes from three isolates of 43 the free-living species E. voratum revealed genome features representative of the 44 Symbiodiniaceae progenitor, due to the absence of symbiogenesis in the Effrenium lineage 45 (Shah et al. 2023). These features include longer introns, more extensive RNA editing, less 46 pseudogenisation, and, perhaps most surprisingly, similar genome sizes when compared to 47 symbiotic counterparts. The genome size of E. voratum suggests that genome reduction (to 48 haploid genome size < 3Gbp) occurred in symbiodiniacean dinoflagellates before 49 diversification of Order Suessiales (Shah et al. 2023). These results further hint at a role of 50 symbiotic lifestyle in shaping intraspecies genomic divergence and the evolution of these 51 taxa. Intragenomic variation of the ITS2 phylogenetic marker sequences is known among 52 Symbiodiniaceae taxa (Wilkinson et al. 2015; Hume et al. 2019). However, intraspecies 53 whole-genome divergence in these taxa relative to symbiotic versus free-living lifestyle 54 remains little known. Whole-genome data from multiple isolates of a species provide an 55 excellent analysis platform to address this knowledge gap.

56 Here, we investigate intraspecies genomic divergence in four Suessiales species (of 57 which three are Symbiodiniaceae); these taxa represent two free-living species and two 58 symbiotic species, for which whole-genome data from multiple isolates are available. We 59 focus specifically on sequence and structural conservation, gene family dynamics, and gene 60 duplication, and how these features may reflect adaptation to the distinct lifestyles.

# 61 **Results and Discussion**

62 We used four Suessiales species for which multi-isolate genome data are publicly available, 63 to investigate patterns of intraspecies genomic divergence related to facultative lifestyle. The two symbiotic symbiodiniacean species, S. microadriaticum (González-Pech et al. 2021; 64 65 Nand et al. 2021) and Durusdinium trenchii (Dougan et al. 2022a), represent taxa that arose from independent origins of symbiogenesis (Figure 1 and Supplementary Table S1). The 66 67 remaining two are free-living species, the symbiodiniacean E. voratum (Shah et al. 2023) and Polarella glacialis that is sister to the Symbiodiniaceae in the Order Suessiales (Stephens et 68 69 al. 2020). The available genome data were generated from isolates collected over vast 70 geographic areas: the thermotolerant symbiont D. trenchii from the Caribbean Sea and 71 Pacific Ocean, the free-living E. voratum from the Mediterranean Sea and both sides of the 72 Pacific Ocean, the symbiotic S. microadriaticum from the Red Sea, Pacific Ocean, and the 73 Caribbean Sea, and the psychrophilic P. glacialis from the Antarctic and Arctic oceans 74 (Figure 1). Collectively, these data provide a robust analytic framework for interrogating 75 intraspecies genomic divergence.

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#### 77 Genomes of facultative symbionts exhibit higher sequence divergence

We investigated divergence of genome sequence following the approach of González-Pech et 78 79 al. (2021). For each pairwise comparison of genome sequences, we calculated the percentage 80 of aligned bases, *Q*, and overall sequence identity of aligned regions, *ID*. Genome sequences 81 from isolates of the same species are highly similar (Q > 70.2%, ID > 98.6% with minimum 82 alignment length 100 bp; Figure 2A, see Supplementary Figure S1 for detail), compared to those between species (Q < 10.0%, ID < 98.6%). High intraspecies sequence similarity was 83 84 observed despite the diverse geographic origins for isolates from each species (Figure 1). Genome sequences of the free-living P. glacialis were the most similar (Q = 95.5%, ID =85 86 98.7%; CCMP1383 against CCMP2088), followed by the symbiotic D. trenchii (Q = 93.3%, 87 ID = 99.8; CCMP2556 against SCF082), the free-living E. voratum (Q = 92.0%, ID = 99.4%; 88 RCC1521 against rt-383), and the symbiotic S. microadriaticum (Q = 78.5%, ID = 99.7%; 89 CCMP2467 against CassKB8). Among the three E. voratum isolates, CCMP421 showed

- smaller percentage of aligned genome bases against rt-383 (Q = 70.2%) and against
- 91 RCC1521 (Q = 79.2%), compared to Q = 92.0% observed between RCC1521 and rt383; this
- 92 is likely due to the more-fragmented CCMP421 genome assembly, also reflected in the low

percentage of mapped sequence reads (Supplementary Table S2). Between the two symbiotic
species, the greater divergence observed in *S. microadriaticum* might represent its much
earlier emergence and diversification (LaJeunesse et al. 2018). Alternatively, the lower
divergence in *D. trenchii* may be due to the recent whole-genome duplication (WGD) in this
lineage (Dougan et al. 2022a). Genome data of multiple isolates from a broader taxon
representation of Symbiodiniaceae lineages will help clarify the possible link between
intraspecies divergence and facultative lifestyle of these symbionts.

To extend genome comparisons beyond alignable sequence regions, we further assessed sequence divergence using an alignment-free *k*-mer-based approach. This approach was found to be robust against the contiguity of genome assemblies (Dougan et al. 2022c), and has been applied successfully to discover distinct phylogenetic signals in different genomic regions of Symbiodiniaceae (Lo et al. 2022; Shah et al. 2023). We followed Lo et al. (2022) to derive pairwise  $D_2^S$  distances, *d*, based on shared *k*-mer profiles at k = 23 observed in whole-genome sequences (see Methods). As shown in Figure 2B, the lowest sequence

107 divergence was seen in *P. glacialis* (d = 0.30), followed by *E. voratum* (d = 0.53 between

108 RCC1521 and rt-383; d = 0.9 when implicating the more-fragmented CCMP421 assembly),

109 *D. trenchii* (0.54), and the three *S. microadriaticum* isolates (0.72-0.76). This pattern of

110 divergence is consistent with our observations based on Q and ID in Figure 2A.

111 We further assessed the conserved core 23-mers in each species (i.e., k-mers common 112 in genomes of all isolates within a species). For each species, we assessed the extent of 113 genome content shared among the isolates based on x, the percentage of core 23-mers relative 114 to all distinct 23-mers; in the perfect scenario where genomes of all isolates are identical, x =115 100%. Using this approach, E. voratum and S. microadriaticum show similar extent of shared genome content among their corresponding isolates (x ranges between 19.5% and 25.2%; 116 117 Supplementary Table S3). Approximately two-fold greater x was observed for P. glacialis 118 (52.3-54.9%) and D. trenchii (55.6-55.7%); this observation likely reflects the impact of a 119 diploid genome assembly in the former (Stephens et al. 2020) and WGD in the latter (Dougan 120 et al. 2022a). Duplicated genomic regions arising from WGD are resolved over long 121 evolutionary time scales of hundreds of millions of years (Carretero-Paulet and Van de Peer 122 2020). Given the recent (~1 MYA) WGD in D. trenchii, this species likely has not had 123 sufficient time to resolve genetic redundancy. Regardless, our results here lend support to the 124 general utility of k-mer-derived distances in clarifying genome-sequence divergence beyond 125 gene boundaries, which may serve as evidence to guide or complement taxonomic

126 classification of Symbiodiniaceae, and potentially of other dinoflagellates (Dougan et al.

127 2022c).

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#### 129 Intraspecies structural divergence in the genomes of Symbiodiniaceae

130 To assess intraspecies structural genomic divergence, we identified collinear gene blocks in 131 all possible pairwise genome comparisons for each species (see Methods); the greater 132 recovery of these blocks and their implicated genes indicates a greater conserved syntemy 133 among the isolates in a species. As expected, due to recent WGD, the two symbiotic D. 134 trenchii isolates CCMP2556 and SCF082 displayed the greatest conserved synteny (1,613 135 blocks implicating ~22% of total genes spanning 181-199 Mbp; Supplementary Table S4). 136 On the other hand, genomes of the symbiotic S. microadriaticum (100-196 blocks, 2.7-3.6% 137 of genes, 8.1-16 Mbp) showed less conserved synteny than the free-living E. voratum 138 RCC1521 and rt383 (344 blocks, 6.6-8.1% of genes, 51-60 Mbp; Supplementary Table S4); 139 at first glance this result appears to support observations in an earlier study (González-Pech et 140 al. 2021) that the extent of structural rearrangements is greater in genomes of facultative symbionts than those of free-living taxa. However, the greater contiguity of the E. voratum 141 142 assemblies (scaffold N50 length = 720 Kbp for RCC1521, 252 Kbp for rt-383) than that of S. 143 *microadriaticum* assemblies (e.g., scaffold N50 length = 43 Kbp for CassKB8 and 50 Kbp for 144 04-503SCI.03) represents a systematic bias that would affect recovery of collinear gene 145 blocks. S. microadriaticum CCMP2467 (N50 length 9.96 Mbp) (Supplementary Table S1), 146 the sole representation of a chromosome-level assembly, lacks comparative power in this 147 instance. As a case in point, the inclusion of the fragmented assembly of E. voratum 148 CCMP421 (N50 length 304 Kbp; 38,022 scaffolds) lowers the extent of conserved synteny 149 identified in E. voratum (195-331 blocks, 4.4-7.9% of genes, 30-65 Mbp; Supplementary 150 Table S4), and we identified no collinear gene blocks between the outgroup P. glacialis 151 isolates due in part to sparsity of genes on the assembled genome scaffolds (Stephens et al. 152 2020). These results in combination suggest that while structural rearrangements contribute to 153 structural divergence of Symbiodiniaceae genomes as postulated in those of facultative 154 symbionts (González-Pech et al. 2019) even at intraspecies level, such an analysis based on 155 collinear gene blocks is sensitive to contiguity of assembled genome sequences. An in-depth 156 assessment of structural divergence would require genome assemblies of comparably high 157 quality.

#### 159 Genetic duplication enables functional innovation

160 We assessed the evolution of protein families for evidence of functional innovation and 161 divergence within species, and its relation to lifestyle. For each species, we inferred 162 homologous protein sets with OrthoFinder using sequences predicted from all corresponding 163 isolates (see Methods); the homologous sets that are specific to an isolate may reflect 164 instances of contrasting divergence in and/or specialisation of protein functions (e.g., putative 165 remote homologs), occurring at distinct evolutionary rates. First, we assessed number of 166 isolate-specific sets for each species based on OrthoFinder results ran at default parameters 167 (i.e., inflation parameter I = 1.5). The highest percentage of isolate-specific sets was observed 168 in D. trenchii (17.2% of total sets), followed by P. glacialis (16.0%); these numbers are 169 nearly four-fold greater than that observed in S. microadriaticum (4.0%) and E. voratum 170 (4.1%; Figure 3). To investigate the robustness of this result, we increased the inflation 171 parameter (1) for clustering within OrthoFinder that controls the granularity (i.e., higher 172 inflation parameter produces smaller clusters). As expected in all cases, the increase of I 173 resulted in an increase of isolate-specific protein sets; at I = 10, the percentage of these sets is 174 37.8% (D. trenchii), 32.4% (P. glacialis), 15.6% (S. microadriaticum), and 10.8% (E. 175 voratum). Despite the high synteny and sequence conservation in D. trenchii, the substantial 176 number of protein families retained in duplicate after WGD show evidence of isolate-specific 177 divergence and/or specialization in D. trenchii where facultative lifestyle has been hypothesized to be the main driver of post-WGD adaptation (Dougan et al. 2022a). On the 178 179 other hand, the comparable extent of isolate-specific protein sets in P. glacialis may represent 180 heterozygosity inherent to a diploid representation of the genome assembly (Stephens et al. 181 2020), distinct from the haploid genome assemblies among the Symbiodiniaceae taxa. None 182 of the *E. voratum* and *S. microadriaticum* isolates showed evidence of WGD (Supplementary 183 Table S5), and thus the similar level of isolate-specific divergence in these species supports 184 the notion of massive genome reduction in the Suessiales ancestor, with WGD a mechanism 185 for escaping this process to generate functional innovation, as observed in D. trenchii 186 (Dougan et al. 2022a). 187

# 188 Genomes of free-living species exhibit greater extent of tandemly duplicated single-exon 189 genes

190 Tandemly duplicated (TD) genes, i.e., duplicated genes found next to each other on the

191 genome, are part of unidirectional gene clusters commonly found in dinoflagellates, thought

192 to facilitate their expression (Nand et al. 2021; Chen et al. 2022). In an earlier study 193 (Stephens et al. 2020), ~40% of the gene repertoire in P. glacialis genomes were located in 194 unidirectional gene clusters, many of which encoded functions associated with cold and low-195 light adaptation. Here we defined a TD block as a block comprising two or more consecutive 196 genes with high sequence identity on a genome scaffold (see Methods). In our independent 197 survey of TD genes in all 19 available Suessiales genomes, we found the largest number and 198 proportion of TD genes in the free-living lineages of P. glacialis (7.8% in CCMP1383, 9.2% 199 in CCMP2088) and S. natans (7.1%), followed by the symbiotic S. tridacnidorum 200 CCMP2592 (6.5%) and C. goreaui SCF055 (6.0%), with smaller proportions observed in the 201 free-living E. voratum (3.9% in rt-383, 4.4% in RCC1521), and the smallest in S. 202 microadriaticum (1.0-2.2%) (Table 1). Some of the largest TD blocks consisted of 13-16 203 genes, found in genomes of free-living lineages (S. natans, and the P. glacialis CCMP1383 204 and CCMP2088). Among the free-living *E. voratum* isolates, the TD block sizes were slightly 205 smaller, implicating genes encoding ribulose bisphosphate carboxylase (the largest block of 9 206 genes in RCC1521), HECT and RLD domain-containing E3 ubiquitin protein ligase 4 (rt-207 383, 7 genes), calmodulin (rt-383, 7 genes), and solute carrier family 4 (rt-383, 7 genes) 208 (Supplementary Table S6); these implicated functions are essential for photosynthesis, ion 209 binding, and transmembrane transport. However, we cannot dismiss the possibility of 210 genome-assembly contiguity in affecting recovery of TD blocks. For instance, the recovery of 211 TD genes in the chromosome-level assembly of S. microadriaticum CCMP2467 is 2.2% 212 versus ~1.0% in the other two assemblies, and the recovery of 1.5% in E. voratum CCMP421 213 contrasts to 3.9-4.4% in the other two *E. voratum* genomes. Despite this, a greater extent of 214 TD genes in free-living lineages (P. glacialis: 55.2-59.4%; E. voratum RCC1521: 23.1% and 215 rt-383: 22.5%; S. natans: 21.8%) were single-exon genes, in contrast to the symbiotic D. 216 trenchii and S. microadriaticum (4.2-9.2%) (Table 1). Our results lend support to the notion 217 that tandem duplication may facilitate transcription of genes encoding essential functions 218 implicating single-exon genes, and is potentially more prominent in genomes of free-living 219 taxa than those of symbiotic lineages (Stephens et al. 2020). 220 Introner elements (IE) are non-autonomous mobile elements characterised by inverted 221 repeat motifs within introns that are hypothesised to propagate introns into genes (Worden et

al. 2009; van der Burgt et al. 2012; Huff et al. 2016), which have been found to be more

prevalent in genomes of free-living dinoflagellate species (Farhat et al. 2021; Dougan et al.

224 2022b; Shah et al. 2023). We examined the presence of these elements in the assembled

225 genomes and TD genes for the multi-isolate Suessiales species (Supplementary Table 1). We

- found the proportion of IE-containing genes overall to be less in Symbiodiniaceae (3.2-6.3%)
- than *P. glacialis* (10.7-11.5%), a trend also observed in the genome of bloom-forming
- dinoflagellate species, *Prorocentrum cordatum* (10.4%) (Dougan et al. 2022b). Nonetheless,
- IEs were only found in a small proportion of TD genes (2.5-5.7%) per Suessiales isolate,
- 230 suggesting they are neither connected to lifestyle nor play a major role in propagating TD
- 231 genes in Suessiales (Supplementary Table S1).
- 232

#### 233 Most tandemly duplicated genes undergo purifying selection

- To assess selection acting on TD genes, we focused on the two best-quality genome
- assemblies (based on number of scaffolds and N50 length) from each species (i.e., total of
- eight isolates), excluding the fragmented assemblies of *E. voratum* CCMP421 and *S.*
- 237 *microadriaticum* CassKB8. We calculated the ratio  $\omega$  as the nonsynonymous substitution rate
- 238  $(K_a)$  to synonymous substitution rate  $(K_s)$  between all possible gene pairs within each TD
- block (Supplementary Table S6; see Methods); in general,  $\omega > 1.0$  indicates positive
- 240 selection,  $\omega = 1.0$  indicates neutral selection, whereas  $\omega < 1.0$  indicates purifying selection
- 241 (Yang and Bielawski 2000) among TD genes within a block. Based on this analysis,
- 242 compared to genomes of symbiotic species, those of free-living species yielded larger
- proportions of TD blocks with mean  $\omega < 1.0$ , indicating purifying selection, i.e., 71.7% in *P*.
- 244 glacialis and 67.7% in E. voratum, compared to 64.2% in D. trenchii and 49.1% in S.
- 245 *microadriaticum* (Figure 4A; Supplementary Table S7). In all cases, the mean K<sub>s</sub> value per
- TD block is less than 0.5 (Figure 4B). The observed mean  $\omega$  values are similar between two
- isolates of a species, e.g., mean variance of  $\omega = 0.26$  for both *P. glacialis* isolates
- 248 (Supplementary Figure S2), suggesting a common pattern of selective pressures acting on TD
- 249 genes for the species. An exception is the symbiotic S. microadriaticum (mean variance of  $\omega$
- 250 = 0.16 for 04-503SCI.03 and 0.95 for CCMP2467; Supplementary Figure S2), but more
- 251 genome data from other multi-isolate symbiotic species will enable the systematic
- investigation of the possible links between selection acting on TD genes and lifestyles.
- To assess functions encoded by TD genes, we focused on TD gene blocks that were recovered in genomes of both isolates in one or more species. Functional annotation of these gene blocks is shown in Figure 4C, and the mean  $\omega$  value for the corresponding block is shown in Figure 4D. Genes encoding calmodulin, sulfotransfer domain-containing proteins, and disulfide-isomerase proteins were recovered in TD blocks in all eight isolates. Fructosebisphosphate aldolase, dinoflagellate viral nucleoproteins, and caltractin were recovered in at

259 least 7 of the 8 isolates. Genes in TD blocks recovered only in free-living P. glacialis and E. 260 voratum encode functions related to photosynthesis (i.e., photosystem I reaction centre 261 subunit III, chloroplast TIC 20-II protein, PS II complex 12 kDA extrinsic protein, and peridinin-chlorophyll *a*-binding protein). In comparison, those in TD blocks found only in the 262 263 two symbiotic species encode for Nek1 protein that is involved in maintaining centrosomes, 264 and NaCP60E, a sodium channel protein. Most of these functions were encoded by no more 265 than 50 TD genes per isolate (Figure 4C) in which the mean  $\omega$  per gene block was < 1 266 (Figure 4D). These results do not speak directly to the specificity of gene functions to tandem 267 duplication in the genomes we analysed, given that some gene copies may also occur 268 elsewhere in the genomes. However, our results suggest a tendency for TD genes within a 269 block to undergo purifying selection, regardless of lifestyle.

270

#### 271 Concluding remarks

272 Our results, based on multi-isolate whole-genome data from representative species,

- 273 demonstrate how facultative lifestyle or the lack thereof has shaped the genome evolution of
- 274 Symbiodiniaceae dinoflagellates. Generation of genetic and functional diversity at the
- 275 intraspecies level implicates genetic duplication, including tandem duplication of genes. All
- these evolutionary regimes are under the constraint of genome reduction that is hypothesised
- to pre-date the diversification of Order Suessiales (Shah et al. 2023). Although our results
- 278 hint at the potential linkages of facultative lifestyles to some of the varying features observed
- between free-living versus symbiotic species, whole-genome data from a broader taxonomic
- representation (and from multiple isolates) will enable a more-systematic investigation toestablish these linkages.
- 282

#### 283 Methods

#### 284 Data

For this study, we used publicly available genome assemblies and gene models of *D. trenchii* 

- 286 CCMP2556 and SCF082 (Dougan et al. 2022a), E. voratum isolates RCC1521, rt-383, and
- 287 CCMP421 (Shah et al. 2023), S. microadriaticum CCMP2467 (Nand et al. 2021), 04-
- 288 503SCI.03 and CassKB8 (González-Pech et al. 2021), and P. glacialis CCMP1383 and
- 289 CCMP2088 (Stephens et al. 2020) (Supplementary Table S1). To contrast the contiguity of

- 290 these genome assemblies, we obtained chromosome numbers from cytological observations
- 291 (Blank and Trench 1985; Jeong et al. 2014; Wham et al. 2017). For tandem gene duplication
- analysis, we used genomic datasets from 9 more Symbiodiniaceae isolates (Supplementary
- Table S1) generated in Chen et al. (2020; 2022), González-Pech et al. (2021), and Shoguchi
- et al. (2013; 2018). To determine the intraspecific identity of the three *E. voratum* genome
- datasets, we mapped the short-read gDNA of each isolate obtained from (Shah et al. 2023) to
- 296 each other using Bowtie2 v2.4.4 (Langmead and Salzberg 2012) with the --very-fast
- algorithm.

#### 298 Assessment of genome-sequence similarity based on alignment

- 299 To assess genome-sequence similarity of the four target species based on sequence
- 300 alignment, we used nucmer (--*mum*) implemented in MUMmer 4.0.0beta2 (Marçais et al.
- 301 2018) at minimum alignment lengths of 100 bp, 1 Kb, and 10 Kb to align assembled genome
- 302 sequences for every possible pair of isolates in each species. For each pairwise comparison,
- 303 we calculated the percentage of aligned bases, *Q*, and overall sequence identity of aligned
- 304 regions, *ID*. Maximum values of for both *Q* and *ID* at 100% indicate that two genome
- 305 sequences are identical. We then used mummerplot (*-f -- layout*) and dnadiff to generate
- 306 figures and reports for these alignments.

#### 307 Assessment of genome-sequence similarity using an alignment-free approach

- 308 Adopting the same approach described in Lo et al. (2022), we calculated  $D_2^S$  statistic based on
- 309 shared *k*-mers for each pair of genomes, from which a distance (*d*) was derived. Briefly,
- Jellyfish v2.3.0 (Marçais and Kingsford 2011) was used to derive k-mers (at k = 23) from
- 311 each genome assembly, from which distances were calculated using *d2ssect*
- 312 (https://github.com/bakeronit/d2ssect) from all possible pairs of genomes. Following the
- 313 earlier studies (Lo et al. 2022; Shah et al. 2023), core 23-mers among isolates of each species
- 314 were identified from the extracted 23-mers, using the bash command *comm* (-12). BEDtools
- 315 (Quinlan and Hall 2010) *intersect* was used to find regions of overlap between the core k-
- 316 mers and different genomic features.

# 317 Gene family evolution and introner element search

- 318 To infer homologous protein sets among isolates for a species, all protein sequences predicted
- 319 from all isolates were used as input for OrthoFinder v2.5.4 (Emms and Kelly 2019). The
- analysis was conducted at different inflation parameters (I = 1.5, 2.0, 4.0, 6.0, 8.0, or 10.0).

- 321 From the generated homologous protein sets, the proportion of isolate-specific sets was
- 322 identified. To identify introner elements, we used the introner element sequences identified in
- 323 Shah et al. (2023) from eight Suessiales isolates as a reference for Pattern Locator (Mrázek
- and Xie 2006) to search for inverted and direct repeat motifs within introns.

#### 325 Identification of collinear gene blocks and types of gene duplication

- 326 To identify collinear gene blocks shared by isolates of a species, we first identified
- homologous protein sequences using BLASTp (e-value  $< 10^{-5}$ , query or subject cover > 50%,
- 328 filtered for top five hits for each query). This output was used as input for MCScanX (Wang
- et al. 2012) (-*b* 2) to search for collinear gene blocks between all possible pairs of isolates.
- 330 For *D. trenchii*, we filtered out duplicated genes (Dougan et al. 2022a) from the MCScanX
- 331 output by selecting gene pairs that were more similar to each other (i.e., low nonsynonymous
- 332  $(K_a)$  + synonymous  $(K_s)$  substitution score), then chose gene blocks that still contained  $\geq 5$
- 333 genes. Gene Ontology (GO) terms were assigned to all gene sets via UniProt (version
- 334 2022\_01) to GO (version December 2022) ID mapping on the UniProt website
- 335 (uniprot.org/id-mapping). The *duplicate\_gene\_classifer* implemented in MCScanX was used
- to assess five distinct type of gene duplications: 1) singleton = not duplicated, 2) dispersed =
- 337 duplicated with > 10 genes in between, 3) proximal = duplicated with < 10 genes in between,
- 4) WGD = whole or segmental genome duplication inferred by anchor genes in collinear gene
- blocks comprising at least 5 genes, 5) tandem = duplicated one after the other, i.e., two or
- 340 more consecutive genes on the same scaffold.

#### 341 Analysis of tandemly duplicated genes

- 342 Tandemly duplicated (TD) genes were identified based on the results of MCScanX above.
- 343 For this analysis, we focused on two best-quality genome assemblies from each species, i.e.,
- 344 for a total of eight genomes. For each TD block, we calculated the nonsynonymous
- 345 substitution rate  $(K_a)$  and synonymous rate  $(K_s)$  between all possible pairs of genes within the
- block, using the *add\_ka\_and\_ks\_to\_collinearity.pl* script implemented in MCScanX (Wang
- et al. 2012). The ratio  $\omega$  was defined as  $K_a/K_s$ . When assessing mean  $\omega$  for each TD block,
- instances of infinity values, e.g., due to  $K_s = 0$ , were ignored.

# 349 **Competing interests**

350 Authors declare that they have no competing interests.

# 351 Author contributions

- 352 Conceptualization, SS, KED, DB and CXC; methodology, SS, KED, YC, and CXC; formal
- analysis, SS, KED, and YC; investigation, SS, KED; writing—original draft preparation, SS;
- writing—review and editing, SS, KED, DB, and CXC; visualisation, SS; supervision, KED,
- 355 DB, CXC; funding acquisition, DB and CXC. All authors have read and agreed to the
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- 368
- 369

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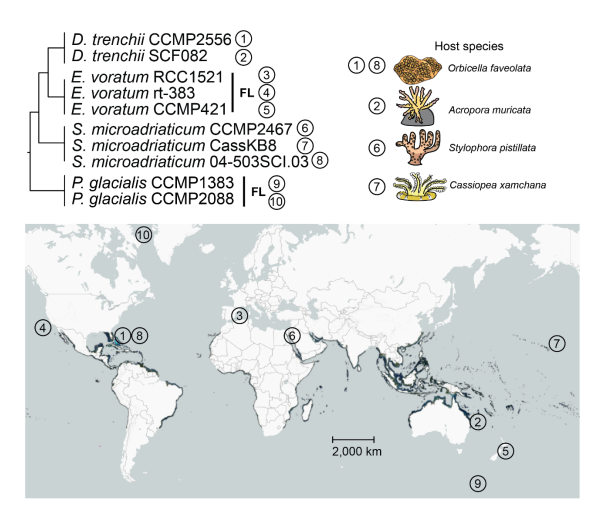
# 468 Table

# 469 Table 1. Tandemly duplicated (TD) genes within 19 Suessiales isolates.

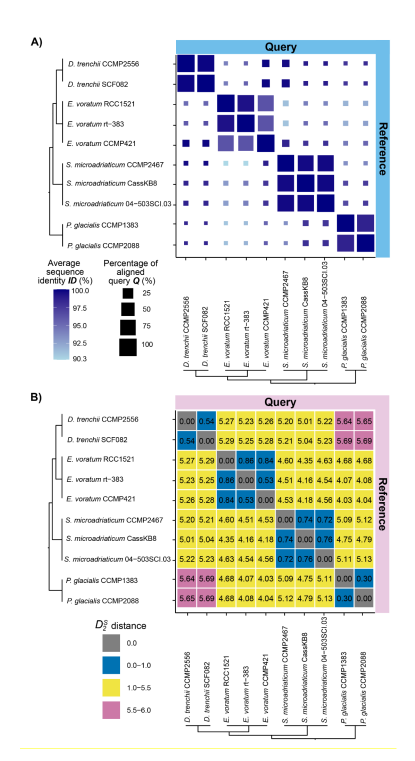
- 470 TD genes were defined as  $\geq 2$  consecutive genes on the same scaffold making up a "block",
- 471 with its size represented by the total number of consecutive TD genes.

Species and isolate	Number of TD genes	Number of TD blocks	Median of TD block size	Maximum TD block size	Number of single-exon genes in the genome	% of single- exon genes among TD genes
B. minutum Mf1.05b.01	1,225 (3.7%)	569	2	7	2,054 (6.3%)	9.9
Cladocopium sp. C92	1,148 (2.5%)	536	2	8	789 (1.7%)	2.2
C. goreaui SCF055	2,017 (6.0%)	937	2	7	1,870 (5.6%)	9.6
D. trenchii CCMP2556	1,031 (1.8%)	745	2	6	3,828 (6.9%)	9.2
D. trenchii SCF082	1,045 (2.0%)	645	2	6	5,677 (10.6%)	7.5
E. voratum CCMP421	495 (1.5%)	233	2	4	1,420 (4.4%)	5.1
E. voratum RCC1521	1,405 (4.4%)	559	3	9	3,983 (12.0%)	23.1
E. voratum rt-383	1,567 (3.9%)	635	3	7	3,574 (9.0%)	22.5
S. linucheae CCMP2456	737 (2.3%)	348	2	6	255 (0.8%)	8.4
S. microadriaticum 04- 503SCI.03	437 (1.1%)	206	2	4	2,734 (7.1%)	5.9
S. microadriaticum CassKB8	418 (1.0%)	200	2	4	3,074 (7.2%)	5.7
S. microadriaticum CCMP2467	1,060 (2.2%)	475	2	7	2,770 (5.7%)	4.2
S. natans CCMP2548	2,499 (7.1%)	1,021	2	13	5,099 (14.5%)	21.8
S. necroappetens CCMP2469	577 (1.6%)	274	2	6	3,187 (8.9%)	14.9
S. pilosum CCMP2461	496 (2.1%)	236	2	4	1,431 (6.1%)	8.3
S. tridacnidorum CCMP2592	2,491 (6.5%)	1,254	2	10	5,192 (11.4%)	19.2
S. tridacnidorum Sh18	581 (2.3%)	272	2	5	3,033 (11.8%)	9
P. glacialis CCMP1383	5,376 (9.2%)	2,095	2	16	15,263 (26.2%)	59.4
P. glacialis CCMP2088	4,028 (7.8%)	1,634	2	14	12,619 (24.4%)	55.2

#### **Figures** 473



- 475 476 Figure 1. Suessiales species, following LSU rDNA phylogeny (LaJeunesse et al. 2018),
- 477 for which genome data of multiple isolates are available.
- 478 Coral reef (in dark blue and cyan) world map by Allen Coral Atlas (2022). Those not marked
- 479 FL (free-living) are symbiotic and their host species are represented on the top right.
- 480
- 481



483 Figure 2. Intra/interspecies genome sequence identity among the four Suessiales species.

- 484 (A) Alignment-based identity (minimum alignment length = 100 bp) with query genome
- 485 sequences (y-axis) aligned to the references (x-axis). The colour of the squares corresponds to
- 486 percent sequence identity *ID* (darker blue = higher identity) and the sizes represent the
- 487 percentage of the query genome sequence Q aligned to the reference. (B) Alignment-free  $D_2^S$
- 488 distances (d) showing delineation between species (d < 1 in blue), Family (d between 1.0 and
- 489 5.5 in yellow), and the longest evolutionary distance across the Order (d > 5.5 in pink).

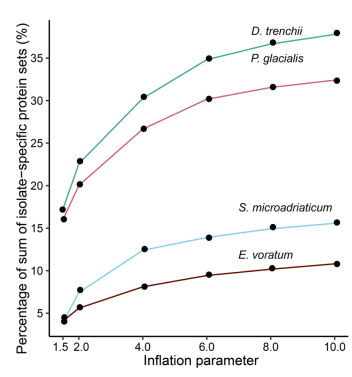
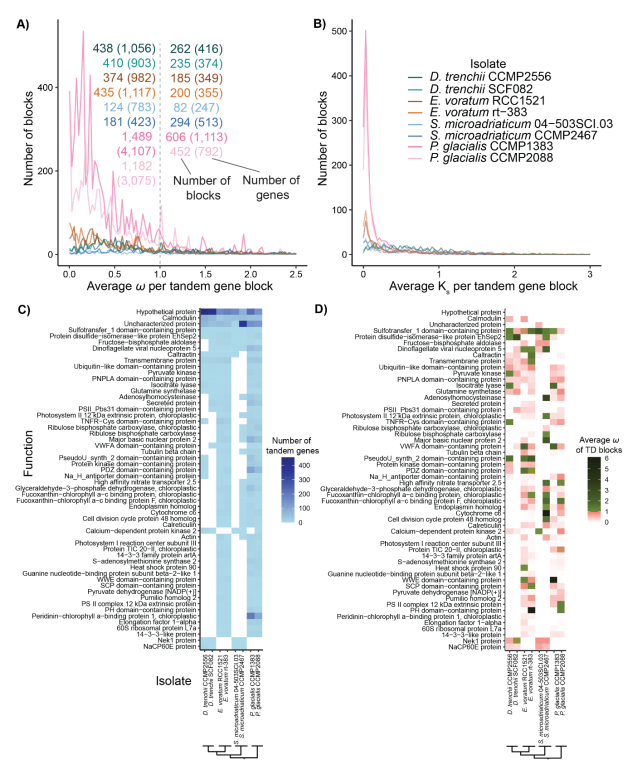




Figure 3. The percentage of isolate-specific protein sets in each Suessiales species.

- 493 Protein sequences were clustered at inflation parameter I between 1.5 and 10 using
- 494 OrthoFinder.
- 495
- 496





498 Figure 4. TD genes and their functions in eight Suessiales isolates.

499 The number of TD blocks showing distribution respectively for (A) mean  $\omega$  and (B) mean  $K_s$ 

of each TD block and its associated TD genes with  $\omega < 1$  or > 1. Functions encoded by TD

- 501 blocks that were recovered in genomes of both isolates in one or more species, showing the
- 502 (C) sum of TD genes, (D) mean  $\omega$ .