LETTER 1 2 EVIDENCE THAT INCONSISTENT GENE PREDICTION CAN MISLEAD ANALYSIS OF ALGAL GENOMES 3 4 Yibi Chen 5 Institute for Molecular Bioscience, The University of Queensland, Brisbane, QLD 4072, 6 Australia; School of Chemistry and Molecular Biosciences, The University of Queensland, 7 Brisbane, QLD 4072, Australia 8 9 Timothy G. Stephens 10 Institute for Molecular Bioscience, The University of Queensland, Brisbane, QLD 4072, 11 Australia 12 13 Debashish Bhattacharya 14 15 Department of Biochemistry and Microbiology, Rutgers University, New Brunswick, NJ 08901, USA 16 17 Raúl A. González-Pech 18 Institute for Molecular Bioscience, The University of Queensland, Brisbane, QLD 4072, 19 Australia 20 21 22 Cheong Xin Chan<sup>1</sup> Institute for Molecular Bioscience, The University of Queensland, Brisbane, QLD 4072, 23 Australia; School of Chemistry and Molecular Biosciences, The University of Queensland, 24 Brisbane, Queensland 4072, Australia 25 26 27

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Comparative algal genomics often relies on predicted gene models from de novo assembled genomes. However, the artifacts introduced by different gene-prediction approaches, and their impact on comparative genomic analysis, remains poorly understood. Here, using available genome data from six dinoflagellate species in Symbiodiniaceae, we identified potential methodological biases in the published gene models that were predicted using different approaches. We developed and applied a comprehensive customized workflow to predict genes from these genomes. The observed variation among predicted gene models resulting from our workflow agreed with current understanding of phylogenetic relationships among these taxa, whereas those published earlier were largely biased by the distinct approaches used in each instance. Importantly, these biases mislead the inference of homologous gene families and synteny among genomes, thus impacting biological interpretation of these data. Our results demonstrate that a consistent gene-prediction approach is critical for comparative genomics, particularly for non-model algal genomes. We implemented a customized, comprehensive workflow to predict protein-coding genes in six published draft Symbiodiniaceae genomes: Breviolum minutum (Shoguchi et al. 2013), Symbiodinium tridacnidorum, Cladocopium sp. C92 (Shoguchi et al. 2018), Symbiodinium microadriaticum (Aranda et al. 2016), Cladocopium goreaui and Fugacium kawagutii (Liu et al. 2018). These draft genomes, generated largely using short-read sequence data, remain fragmented (e.g. N50 lengths range from 98.0 Kb for C. goreaui to 573.5 Kb for S. microadriaticum); we treat these genome assemblies independently as is standard practice. The published gene models from these four studies were predicted using three different approaches: (a) ab initio using AUGUSTUS (Stanke et al. 2006) guided by transcriptome data (Shoguchi et al. 2013, Shoguchi et al. 2018), (b) ab initio using AUGUSTUS guided by

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and average length of intergenic regions.

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length, proportion of splice-donor site motifs (GT, GC or GA), number of intergenic regions,

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This result suggests that variation among  $\alpha$  genes is predominantly due to methodological

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biases, and that these biases are larger compared to those of  $\beta$  genes. Variation in the latter appears to be more biologically relevant and consistent with Symbiodiniaceae evolution. Genomes that are phylogenetically closely related are expected to share greater synteny than those that are more distantly related. Here, we defined a collinear syntenic gene block as a region common to two genomes in which five or more genes are coded in the same order and orientation. These gene blocks were identified using SynChro (Drillon et al. 2014) at *Delta* = 4. Overall, 421 collinear syntenic blocks (implicating 2454 genes) between any genome-pairs were identified among  $\alpha$  genes, compared to 426 blocks (implicating 2553 genes) among  $\beta$ genes (Figs. 2a and 2b). Based on  $\alpha$  genes comparison (Fig. 2a), S. microadriaticum and S. tridacnidorum shared the largest number of syntenic blocks (130; 760 genes), whereas S. microadriaticum and F. kawagutii shared the fewest (1; 6 genes). Surprisingly, S. tridacnidorum and Cladocopium sp. C92 shared 38 blocks (222 genes). This close relationship is not evident between any other pair of genomes from these two genera (e.g. only 3 blocks implicating 15 genes between S. microadriaticum and C. goreaui), and is even closer than the relationship between the two *Cladocopium* species (i.e. *C. goreaui* and C92: 33 blocks, 187 genes). The unexpectedly high conserved synteny between S. tridacnidorum and Cladocopium sp. C92 may be explained by the fact that these genes were predicted with the same method (Shoguchi et al. 2018). In contrast, based on the  $\beta$  genes comparison (Fig. 2b), the number of syntenic blocks shared between any Symbiodinium and Cladocopium species did not vary to the same extent; e.g. 7 blocks (38 genes) between S. tridacnidorum and Cladocopium sp. C92, and 10 blocks (55 genes) between S. microadriaticum and C. goreaui. The number of  $\beta$  genes implicated in blocks shared by these two genera is also smaller than those between the two *Cladocopium* species (263 genes in 48 blocks), consistent with their closer phylogenetic relationship.

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To assess the impact of methodological biases on the delineation of homologous gene families, Orthofinder v2.3.1 (Emms and Kelly 2018) was used to infer "orthogroups" from protein sequences (i.e. homologous protein sets) encoded by the  $\alpha$  and  $\beta$  genes (Figs 2c and 2d). More homologous sets were inferred among the  $\alpha$  genes (33,580) than among the  $\beta$ genes (26,380), likely due to the higher number of  $\alpha$  genes in all genomes. Genomes from closely related taxa are expected to share more homologous sequences (and therefore more sets) than those that are phylogenetically distant. Most of the identified homologous sets (6431 from  $\alpha$  genes, 4941 from  $\beta$  genes) contained sequences from all analyzed taxa; these represent core gene families of Symbiodiniaceae. Similar to the results of the synteny analysis described above, the pattern of homologous sets shared between members from Symbiodinium and Cladocopium varies among the  $\alpha$  genes (Fig. 2c). For instance, 638 homologous sets are shared only between S. tridacnidorum and Cladocopium sp. C92, compared to 89 between C. goreaui and S. tridacnidorum. In contrast, the corresponding number of homologous sets inferred based on  $\beta$  genes are closer to each other (Fig. 2d); i.e. 100 between S. tridacnidorum and Cladocopium sp. C92, and 132 between C. goreaui and S. tridacnidorum. Our results indicate that comparative genomics using the  $\alpha$  genes (i.e. simply based on published gene models) could lead to the inference that S. tridacnidorum and Cladocopium sp. C92 are more closely related with each other than is each of them with other isolates in their corresponding genus. The bias introduced by different gene-prediction approaches can significantly impact downstream comparative genomic analyses and lead to incorrect biological interpretations. We therefore urge the research community to consider a consistent gene-prediction workflow when pursuing comparative genomics, particularly among highly divergent, non-model algal genomes. Although we only considered dinoflagellate genomes

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LaJeunesse, T. C., Parkinson, J. E., Gabrielson, P. W., Jeong, H. J., Reimer, J. D., Voolstra, 170 C. R. & Santos, S. R. 2018. Systematic revision of Symbiodiniaceae highlights the 171 antiquity and diversity of coral endosymbionts. Curr. Biol. 28:2570-80. 172 Liu, H., Stephens, T. G., González-Pech, R. A., Beltran, V. H., Lapeyre, B., Bongaerts, P., 173 Cooke, I., Aranda, M., Bourne, D. G., Forêt, S., Miller, D. J., van Oppen, M. J. H., 174 Voolstra, C. R., Ragan, M. A. & Chan, C. X. 2018. Symbiodinium genomes reveal 175 176 adaptive evolution of functions related to coral-dinoflagellate symbiosis. Commun. Biol. 1:95. 177 178 Shoguchi, E., Beedessee, G., Tada, I., Hisata, K., Kawashima, T., Takeuchi, T., Arakaki, N., Fujie, M., Koyanagi, R., Roy, M. C., Kawachi, M., Hidaka, M., Satoh, N. & Shinzato, 179 C. 2018. Two divergent *Symbiodinium* genomes reveal conservation of a gene cluster 180 for sunscreen biosynthesis and recently lost genes. BMC Genomics 19:458. 181 Shoguchi, E., Shinzato, C., Kawashima, T., Gyoja, F., Mungpakdee, S., Koyanagi, R., 182 Takeuchi, T., Hisata, K., Tanaka, M. & Fujiwara, M. 2013. Draft assembly of the 183 Symbiodinium minutum nuclear genome reveals dinoflagellate gene structure. Curr. 184 Biol. 23:1399-408. 185 Stanke, M., Keller, O., Gunduz, I., Hayes, A., Waack, S. & Morgenstern, B. 2006. 186 AUGUSTUS: ab initio prediction of alternative transcripts. Nucleic Acids Res. 187 34:W435-W39. 188 189 **Data accessibility** All genome data (after removal of microbial contaminants), and the predicted gene models 190 from this study are available at: <a href="https://cloudstor.aarnet.edu.au/plus/s/JXALPndBKLNYgF9">https://cloudstor.aarnet.edu.au/plus/s/JXALPndBKLNYgF9</a> 191

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Author contribution

YC, RAGP and CXC conceived the study and designed the experiments. YC conducted all computational analyses. YC, TGS, RAGP and CXC analyzed and interpreted the results. YC and RAGP prepared all figures, tables, and the first draft of this manuscript. YC, TGS and RAGP provided analytical tools and scripts. All authors wrote, reviewed, commented on and approved the final manuscript.

Competing interests

The authors declare no competing interests.

## Figures

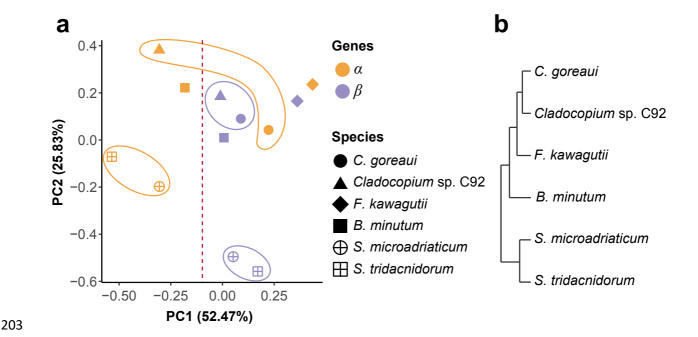


Fig. 1. Gene metrics of  $\alpha$  and  $\beta$  genes from six Symbiodiniaceae genomes. (a) Principal Component Analysis plot based on ten metrics of the predicted gene models, shown for the  $\alpha$  genes in orange, and the  $\beta$  genes in purple, for each of the six genomes (noted in different symbols) as indicated in the legend. The two *Cladocopium* species and the two *Symbiodinium* species were highlighted for clarity. (b) A tree topology depicting the phylogenetic relationship among the six taxa, based on (LaJeunesse et al. 2018).

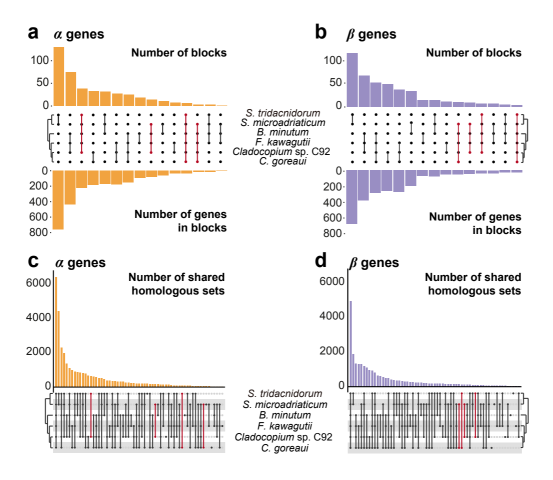


Fig. 2. Conserved synteny and homologous sets among six Symbiodiniaceae genomes.

The number of collinear syntenic gene blocks between each genome-pair is shown for those inferred based on (a)  $\alpha$  and (b)  $\beta$  genes; the upper bar chart shows the number of blocks, the lower bar chart shows the number of implicated genes in these blocks, and the middle panel shows the genome-pairs corresponding to each bar with a line joining the dots that represent the implicated taxa. The number of homologous sets inferred from (c)  $\alpha$  and (d)  $\beta$  genes is shown, in which the taxa represented in the set corresponding to each bar are indicated in the bottom panel. The most remarkable differences between (a) and (b), and (c) and (d), focusing on *Symbiodinium* and *Cladocopium* species are highlighted in red.