A broadly resolved molecular phylogeny of New Zealand cheilostome bryozoans as a framework for hypotheses of morphological evolution

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\textbf{A B S T R A C T}

Larger molecular phylogenies based on ever more genes are becoming commonplace with the advent of cheaper and more streamlined sequencing and bioinformatics pipelines. However, many groups of inconspicuous but no less evolutionarily or ecologically important marine invertebrates are still neglected in the quest for understanding species- and higher-level phylogenetic relationships. Here, we alleviate this issue by presenting the molecular sequences of 165 cheilostome bryozoan species from New Zealand waters. New Zealand is our geographic region of choice as its cheilostome fauna is taxonomically, functionally and ecologically diverse, and better characterized than many other such faunas in the world. Using this most taxonomically broadly-sampled and statistically-supported cheilostome phylogeny comprising 214 species, when including previously published sequences, and 17 genes (2 nuclear and 15 mitochondrial) we tested several existing systematic hypotheses based solely on morphological observations. We find that lower taxonomic level hypotheses (species and genera) are robust while our inferred trees did not reflect current higher-level systematics (family and above), illustrating a general need for the rethinking of current hypotheses. To illustrate the utility of our new phylogeny, we reconstruct the evolutionary history of frontal shields (i.e., a calcified body-wall layer in ascus-bearing cheilostomes) and ask if its presence has any bearing on the diversification rates of cheilostomes.

\textbf{1. Introduction}

Large and broadly-sampled phylogenies are vital to robustly answering many different classes of evolutionary questions, including those involving trait evolution, origins and evolution of biogeographic distributions and rates of taxonomic diversification. While mega-phylogenies with hundreds to thousands of species (Smith et al., 2009) are available for many groups of vertebrates (Meredith et al., 2011; Prum et al., 2015) and plants (Zanne et al., 2014), and also for some non-vertebrate terrestrial groups (Varga et al., 2019), the molecular phylogenetics of many marine invertebrate groups remains relatively neglected (Arrigoni et al., 2017; Kocot et al., 2018; O’Hara et al., 2017).

In this contribution, we begin to rectify the paucity of large and/or taxonomically broadly sampled molecular phylogenies for marine invertebrates, targeting a phylum whose rich fossil record can be subsequently integrated for evolutionary analyses. Our focal group is Cheilostomatida, the dominant living order of the colonial metazoan phylum Bryozoa, with c. 5200 described extant species, corresponding to >80% of the living species diversity of the phylum (Bock and Gordon, 2013). Cheilostomes first appeared in the fossil record in the Late Jurassic (c. 160 million years ago) and then displayed a spectacular diversification c. 55 million years later in the mid-Cretaceous (Taylor, 2020). Cheilostomes, common in benthic marine habitats globally, are lightly- to heavily-calcified and largely sessile as adults. Most species are encrusting, while fewer are erect, with some forming robust structures whereas many are small and inconspicuous (Fig. 1). Although a number of cheilostome bryozoans have been sequenced and placed in a molecular phylogenetic context (Fuchs et al., 2009; Knight et al., 2011; Orr et al., 2019a; Waeschenbach et al., 2012) the systematics of cheilostome bryozoans aimed at reflecting their evolutionary relationships still remain largely based on morphological characters (Bock and Gordon, 2013; Martha et al., 2020; Taylor and Waeschenbach, 2015). This is in

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part because assumed cheilostome phylogenetic relationships have only recently benefited from high-throughput sequencing (HTS) techniques and the increased phylogenetic support it provides (Orr et al., 2019a, b, 2020). HTS yields more sequence data with lesser effort compared with traditional PCR and Sanger sequencing techniques (Fuchs et al., 2009; Knight et al., 2011; Waeschenbach et al., 2012). By applying genome-skimming approaches to greatly expand on the taxonomic sampling of cheilostomes for molecular phylogenetics, we independently test phylogenetic hypotheses implicit in their current systematics (Bock, 2020), and also facilitate future studies.

We focus our sequencing effort in this contribution primarily on New Zealand cheilostomes for a number of reasons. Cheilostomes play a conspicuous role as habitat-building organisms in New Zealand as well as other temperate areas (Cook et al., 2018; Wood et al., 2012). In fact, some cheilostome thicket communities (Fig. 1A) are protected in New Zealand because of their function as nurseries for commercial fish stocks (Bradstock and Gordon, 1983). As important components of marine communities, cheilostomes are crucial members of the marine food chain globally. This is because, like all bryozoans, they are efficient suspension-feeders (Gordon et al., 1987) while also providing food for other organismal groups (Lidgard, 2008). Cheilostomes are highly diverse in New Zealand, thanks to a combination of factors, including New Zealand’s geological and hydrographic setting, constituting the major part of the geological continent of Zealandia, which is 94% submerged (Campbell and Mortimer, 2014). Additionally, the New Zealand Exclusive Economic Zone, plus its extended continental shelf, is one of the largest in the world (5.7 million km²) with a wide latitudinal spread from subtropical to subantarctic (c. 23°–57.5° S). It also has varied seafloor topography, including extensive deep shelves, plateaus, ridges and seamounts (Gordon et al., 2010). Within this area, New Zealand has 359 genera and 1053 species of marine Bryozoa, including 867 cheilostomes (of which 285 species remain to be formally described). About 61% of New Zealand’s marine Bryozoa are endemic (Gordon et al., 2019), making New Zealand a doubtless diversity hotspot for cheilostome bryozoans. Complementing Recent diversity, the published Cenozoic record of cheilostome bryozoans is also rich, though relatively less studied (Brown, 1952; Gordon and Taylor, 2015; Rust and Gordon, 2011), comprising 531 species (of which 240 are in open nomenclature). This complementarity of living and fossil species renders a molecular phylogeny of New Zealand taxa amenable to modern statistical methods that integrate molecular and fossil data for inferring evolutionary processes (Heath et al., 2014). Last, but not least, New Zealand is one of the better-studied marine regions taxonomically and ecologically for Bryozoa (e.g. Gordon, 1984; 1986; 1989; Gordon et al., 2009; Schack et al., 2020), a phylum that is somewhat neglected in many other parts of the world. Bryozoan research has been continuously conducted in New Zealand since 1841 (Gordon et al., 2009) and a governmental agency, the National Institute of Water and Atmospheric Research (NIWA), is both the data manager and custodian for fisheries and invertebrate research data, hence assuring knowledge curation. All of this means that a cheilostome phylogeny with New Zealand species broadly represented allows us to begin to ask evolutionary and ecological questions while controlling for phylogenetic non-independence.

Here we apply a genome-skimming approach to New Zealand cheilostome bryozoans and present a robustly supported molecular phylogeny based on 15 mitochondrial and 2 rRNA genes. The molecular

Fig. 1. New Zealand Bryozoans. (A) Foliose branching colonies of the cheilostome bryozoan Euthyroides episcopalis from Fiordland, New Zealand (photo by Dr Mike Page, NIWA). (B–M) Scanning electron micrographs of various New Zealand cheilostome bryozoans (B–E: anascans;grade; F–M: ascophoran-grade). (B) Steginoporella perplexa (Steginoporellidae; BLEED 1651). (C) Monoporella n. sp. (Monoporellidae; BLEED 1360). (D) Ellisina sericea (Ellisinidae; BLEED 697). (E) Beania stonycha (Beaniidae; BLEED 84). (F) Arachnopusia unicorns (Arachnopusidae; BLEED 221). (G) Orthoscuticella fusiformis (Catenicellidae; BLEED 1623). (H) Chiastosella watersi (Escharinidae; BLEED 56). (I) Callopotina angustipora (Microporellidae; BLEED 793). (J) Smittina rosacea (Smittinidae; BLEED 1700). (K) Bitectipora cincta (Bitectiporidae; BLEED 801). (L) Galeopisis n. sp. 2 (Celleporidae; BLEED 1618). (M) iodoxyum yaldwynii (Philodborporidae; BLEED 1387). All scale bars are 0.5 mm.
sequences of 199 cheilostome colonies sampled in New Zealand are presented here for the first time. Using 180 species and 96 genera from New Zealand and previously sequenced, non-New Zealand species, we construct the largest and most taxonomically broadly sampled cheilostome phylogeny to date, with 263 in-group colonies, representing 214 species and 120 genera. The inclusion of non-New Zealand taxa allows us to explore the robustness of the inferred relationships among New Zealand species but also reduces phylogenetic inference errors by nature of a broader taxonomic sampling (Pollock et al., 2002). To illustrate the utility of our inferred tree for understanding cheilostome evolution, we reconstruct the evolutionary history of a morphological trait (the frontal shield), where it is thought that there have been repeated gains, and perhaps losses, of a calcified (ascophoran) shield (Gordon, 2000). The state of the frontal shield is crucial for the mechanics of extrusion of the feeding tentacles (lophophore) and the protection of the retracted polypide (the soft tissue of the bryozoan) (Taylor, 2020). To demonstrate our inferred tree’s utility, we also ask if the diversification (i.e. speciation and extinction) rates of cheilostomes that have such a shield (ascophoran-grade; Fig. 1F–M) are different from those that do not (anascan-grade; Fig. 1B–E). We also discuss several other key taxonomic traits, including the presence of a frontal shield opening (ascophore; Fig. 1I), widely thought to be evolutionarily stable and the consequences our highly resolved cheilostome phylogeny has for these. Our contribution is a first step towards a global cheilostome megaphylogeny, needed for answering biological questions that go beyond those probing genealogical relationships.

2. Methods

2.1. Sampling & SEM

Sequences are provided here for 207 New Zealand cheilostome colonies that were collected during several field expeditions by NIWA and University of Otago, New Zealand. While we have newly sequenced 199 colonies, we also supply unpublished sequences for 8 extra colonies we previously presented (see Supplementary Table S2). Samples were sorted, preserved in 70–96% ethanol, then shipped to the University of Oslo, Norway, for processing. Each bryozoan colony, preliminarily identified to the lowest possible taxonomic level (usually genus but sometimes species) using a stereoscope, was subsampled for DNA isolation, and also for scanning electron microscopy. The scanning electron micrographs (SEMs), taken with a Hitachi TM4000 Plus after bleaching to remove tissue (where appropriate), are required for species-level confirmation. All SEM digital vouchers are supplied as a supplementary data file. Taxonomic identifications are made independently of the phylogenetic inference and metadata to avoid identification bias.

2.2. DNA isolation, sequencing and assembly

The 199 subsamples of colonies (henceforth “samples”) were dried before genomic DNA isolation using the DNeasy Blood and Tissue kit (QIAGEN, Germantown, MD, USA). Samples were homogenized in lysis buffer, using a pestle, in the presence of protease-K. Genomic DNA were sequenced at the Norwegian Sequencing Centre (Oslo, Norway) using Illumina HiSeq4000 150 bp paired-end (PE) sequencing with a 350 bp insert size. Approximately 20 samples (library preps) were genome-skimmed (multiplexed) on a single lane. Illumina HiSeq reads were quality checked using FastQC v0.11.8 (Andrews, 2010), then quality- and adapter-trimmed using TrimmGalore v0.4.4 with a Phred score cutoff of 30 (Krueger, 2015). Trimmed reads were de novo assembled with SPAdes 3.13 (Bankevich et al., 2012) using k-mers of 21, 33, 55, 77, 99 and 127. The mitogenome and rRNA operon of each sample were identified separately with blastn (Altschul et al., 1990) using blast + against a database constructed from broadly sampled cheilostome sequences already deposited in NCBI (Orr et al., 2020). An E-value of 1.00e–185 and maximum target sequence of 1 were used to filter any blast hits of non-cheilostome origin.

2.3. Annotation

Mitogenomes for each of the samples were annotated with Mitos2 using a metazoan reference (RefSeq 89) and the invertebrate genetic code (Bernt et al., 2013) to identify two rRNA genes (rrnL and rrnS) and 13 protein coding genes (ap6, ap8, cox1, cox2, cox3, cob, nad1, nad2, nad3, nad4, nad5, nad6). In addition, two nuclear rRNA operon genes (ssu/18 s and lsa/28 s) were identified and annotated using RnaMMer (Lagesen et al., 2007). The internal transcribed spacer regions (ITS1 and 2) and the 5.8 s rRNA were not utilized in this study. Thirty published (Orr et al., 2019a, b, 2020) New Zealand samples were included in the subsequent workflow to bring the total number to 229 (Supplementary Table S2). Further, the mitogenomes and rRNA operons of 38 non-New Zealand bryozoans (Orr et al., 2020), were aligned with our samples to compile a broader cheilostome ingroup and ctenostome outgroup taxon sample.

2.4. Aligning

MAFFT (Katoh and Standley, 2013) was used for alignment with default parameters: for the four rRNA genes (nucleotide) the Q-INS-i model, considering secondary RNA structure, was utilized; for the 13 protein-coding genes, in amino acid format, the G-INS-i model was used. The 17 separate alignments were edited manually using Mesquite v3.61 to remove any uncertain characters (Maddison and Maddison, 2017). Ambiguously aligned characters were removed from each alignment using Gblocks (Talavera and Castresana, 2007) with least stringent parameters. The single-gene alignments were concatenated to a supermatrix using the catfasta2phyml perl script (Nylander, 2010). The alignments (both masked and unmasked) are available through Dryad (https://doi.org/10.5061/dryad.7pmcvdr3).

2.5. Phylogenetic reconstruction

Maximum likelihood (ML) phylogenetic analyses were carried out for each single gene alignment using the “AUTO” parameter in RAxML v8.0.26 (Stamatakis, 2006) to establish the evolutionary model with the best fit. The general time reversible (GTR + G) was the preferred model for the four rRNA genes (18 s, 28 s, rrnL and rrnS), and MtZoa + G for all 13 protein coding genes. The two concatenated datasets (“New Zealand” and “global” = New Zealand + non-New Zealand, see section above), divided into 17 separate RNA and protein gene partitions each with its own distinct gamma distribution to accommodate for different substitution patterns among sites, were analyzed using RAxML. The topology with the highest likelihood score of 100 heuristic searches was chosen. Bootstrap values were calculated from 500 pseudo-replicates.

Bayesian inference (BI) was performed using a modified version of MrBayes incorporating the MtZoa evolutionary model (Huelsenbeck and Ronquist, 2001; Tanabe, 2016). The datasets were executed, as before, with 17 separate rRNA and protein gene partitions under their distinct gamma distributions. Two independent runs, each with three heated and one cold Markov Chain Monte Carlo (MCMC) chain, were initiated from a random starting tree. The MCMC chains were run for 20,000,000 generations with trees sampled every 1,000th generation. The posterior probabilities and mean marginal likelihood values of the trees were calculated after the burnin phase (5,000,000 generations). The average standard deviation of split frequencies between the two runs was < 0.01, indicating convergence of the MCMC chains.

Congruence between the topological signal of the bryozoan nuclear rRNA operon (Supplementary Fig. S7) and mitogenome (Supplementary Fig. S8) was tested, to support their concatenation, using the Icong index (de Vienne et al., 2007). As the Icong index is dependent on identical leaves between topologies the analysis was performed on a subset (218...
of 267) where only taxa present in both the nuclear rRNA and mitogenome datasets, with < 70% missing characters for each alignment, were represented (see Table S4 and available through Dryad).

2.6. Ancestral state reconstruction and BiSSE analyses

The tips states of whether the sampled species is anascan (0, having a non-calcified frontal membrane; e.g. Fig. 1D) or ascophoran (1, having a calcified frontal shield; e.g. Fig. 1K), both states decipherable from SEMs, is given in Fig. 3. We use a standard Markov model of binary character evolution (Pagel, 1994) implemented in ape (Paradis and Schliep, 2018) to estimate the ancestral states of the nodes on our inferred phylogeny. We use a standard binary state speciation and extinction model, also termed BiSSE (Maddison et al., 2007) implemented in diversitree (FitzJohn, 2012) to investigate any differences in diversification rates due to the anascan or ascophoran frontal shield state of the species involved. As input for this latter analysis, we estimate that of the 1876 anascans and 3358 ascophoran species in Bock (2020), we have sampled 4.4% and 3.9% respectively to account for biases due to the sampling of species given the trait. We perform ancestral state reconstruction and BiSSE analyses.
reconstruction and BiSSE analyses for both ML and Bayesian “global” trees (Supplementary Figs. S3 and S4 respectively) to account for minor differences in the topological signal (see Results) and present posterior distributions of estimated speciation ($\lambda$) and extinction ($\mu$) rates given an anascan (0) or ascophoran (1) state, as well transition ($q$) rates between the two states. Note that as we do not (fossil) calibrate the branch lengths (average number of substitutions per site over the alignment) to absolute time, the estimated rates from the BiSSE will be presented in units of substitutions. In cases where there are multiple representatives within a species, we choose the colony with the highest number of nucleotides/amino-acids/genes to represent the species for these analyses.

3. Results

3.1. Sequencing and concatenation

We successfully sequenced and assembled 199 New Zealand...
chelostome colonies, representing 165 species (SEM vouchers in Supplementary file) that have never been presented previously (Supplementary Table S1). We supply additional sequence data for a further eight species previously presented (Supplementary Table S2 and Orr et al., 2019b). The final 17 gene and 267 taxa “global” supermatrix constitutes 77% total character completeness for the dataset used to infer Fig. 2. For the convenience of future workers interested in only the New Zealand taxa, we supply also trees based on these data (Supplementary Figs. S1 (ML) and S2 (Bayesian), where character completeness is 78%). The assembled rRNA and mitogenomes are deposited at NCBI with accession numbers (Supplementary Table S2).

3.2. A global chelostome phylogeny

3.2.1. Broad taxon-sampling

Our inferred “global” chelostome phylogeny, encompassing 214 species and 120 genera, from 56 families (Fig. 2) of which 229 colonies, 186 species and 96 genera, currently distributed in 48 families, are from New Zealand (Supplementary Figs. S1 and S2). The New Zealand and global trees represent c. 21% described species of chelostomes from New Zealand and c. 15% of the described chelostome genera globally, respectively. Both phylogenies (Fig. 2 and Supplementary Fig. S1) are robustly resolved with most branches (146 of 195 branches, or approximately 75% based on Fig. 2) receiving either high (>90 bootstrap (BS))/>0.99 Posterior Probability (PP)) or full support (100 BS / 1 PP). Our ingroup chelostome taxa form a fully supported monophyletic clade, when we infer the global tree including a ctenostome outgroup (Fig. 2).

We summarize only general ingroup observations while referring the reader to topological details in Fig. 2 and Supplementary Fig. S3 that are not discussed here or in the Discussion. We also refrain from summarizing results above the family-level for reasons stated in the Discussion.

3.2.2. Family relationships

Several families for which we have three or more genera represented form supported monophyletic (Fig. 2), e.g. the fully supported Cateneciliidae (Orthoscuticella, Costaticella, Paracrucibrillina, Talivittaticella, Pterocella, Catenicella, Cornuticella and Terminocella; Fig. 1G), Adeoideae (Adeoidopsis, Adeona, Reptadeonella, Laminopora, Cucellipora, Adeonella), Flistridae (Flistra, Hinchskia, Securiflistra), Hippothoidae (Cellepora, Hippothoa, Antarctothoa) and Phidoloporidae (Iodictyum, Hippolossoo, Phidolora, Stephanonella, Rhyhchoossoo; Fig. 1M). The monophyly of the Candidae (Menipea, Amastigia, Caberea, Canda, Emma) conversely, receives poor support (-/0.97).

Of the 29 nominal families represented by two or more genera in our phylogeny, only 12 (c. 41%) are monophyletic in our inference (Fig. 2 green boxes). Families such as the Microporidiae (Micropora, Oparoophora, Calponia), Calloporidae (Valdemunitella, Crassimarginatella, Callopora, Amphibletrum), Bugulidae (Dimetopita, Bugula), Romancheinidae (Hippomenella, Escharoides and Eoxochella), and Microporidiae (Micro- porella, Calloporina), all currently accepted in Bock (2020), are recovered as polyphyletic with high support (Fig. 2 brown boxes), while others such as Euthyroididae are parapatric (Fig. 2, orange boxes). Monogenic families (e.g. Cripdacidathideae, Macronorideae and Powellidheicidae) recovered as fully supported monophyly comprising multiple species are not considered here.

3.2.3. Genus relationships

In contrast to family-level systematics, the 50 currently morphologically defined nominal genera for which we have two or more representatives in general (approximately 70%) form monophyletic groupings (e.g. Parasmitina, Bectipora, Rhynchosoo, Micro- porella, Amphibletrum, Micropora, Steginoporella and 27 others) with either high or full support. A few genera are non-monophyletic (approximately 30% of those for which we have at least two representatives): several are recovered as parapatric in our tree (Chiastosella, Fenestrulina, Smittioidea, Schizosmitina, Chaperiopis and Valdemunitella), while only a handful are polyphyletic (Celleporina, Galeopsis and Osthimia).

Because there are indications that some species are phenotypically highly variable and others have morphologies that are not yet well-understood, we also sequenced multiple colonies of the same species in several cases even though our goal was to sequence one colony of each species. Morphologically identified species match genetic species inferred by phylogenetic inferences in these cases, including, Parkerinella punciterta (98.86% id over 15078 bp), Chiastosella longaevitas (99.56% id over 15142 bp), C. enigma (99.85% id over 14403 bp), Microporella agonistes (99.85% id over 14303 bp) and M. intermedia (99.62% id over 13938 bp). We note that while our three Parasmittina aotea samples form a monophyletic clade, there is somewhat greater genetic variability than the species mentioned above (86.91% id over 14236 bp). For more details, please see individual SEM cards in the Supplementary data file.

3.2.4. Congruent trees and a single incongruent branch

We show the inferred global nuclear rRNA (Supplementary Figs. S7 and S9) and mitogenome trees (Supplementary Figs. S8 and S10) to be topologically more congruent than expected by chance (Ips index = 3.77; probability that they are topologically unrelated = 6.54e–36). The result supports the concatenation of rRNA and mitogenome data in chelostome bryozoans, as previously demonstrated (Orr et al., 2019a), albeit on a smaller dataset.

For the concatenated nuclear rRNA and mitogenome datasets, we highlight the incongruent placement of the Euthyroidia, Figularia and Valdemunitella clade between the ML and Bayesian trees. Note that this clade is highly supported as a monophyly in both sets of trees, but its placement within the trees is contested; the ML trees, whether based only on the New Zealand taxa or all taxa (Fig. 2, Supplementary Figs. S1 and S3), place this clade in a basal position with an affinity to the Macropora/Monoporella groupings. The Bayes trees, however, infer a more derived position. In all instances (ML and Bayes), support for the inferred placement is lacking.

3.3. Ancestral state reconstruction and BiSSE

A different rates model for the transition of the anascan to ascophoran state has a less negative log-likelihood (-29.24) than that for an equal rates model (-35.16), suggesting that it describes our ML tree better. Parameter estimates indicate that the ascophoran state never goes to anascan, and anascan state goes to ascophoran at rate of 0.207 (std err 0.0273); in our ML tree. The estimated node states are shown in Fig. 3. Plots of posterior distributions of speciation and extinction rates (in terms of average number of substitutions) given the frontal shield trait show a substantial overlap (Fig. 4) where the means of each group (anascan or ascophoran) are encompassed in the 95% CI (Credibility Intervals in parenthesis) of the other group in each comparison: μanascan = 17.74 (13.15, 26.05); μascophoran = 15.32 (9.43, 29.62), μanascan = 12.57 (7.51, 21.56); μascophoran = 8.48 (1.39, 24.23). However, the transition rates of the states are non-overlapping in their 95% CI; q01 (anascan to ascophoran) = 0.19 (0.08, 0.42), q10 (ascophoran to anascan) = 0.03 (0.00, 0.11). If we assume that chelostomes originated in the Late Jurassic approximately 160 million years ago (Taylor and Waeber, 2015), then μanascan = 0.12, μascophoran = 0.10, μanascan = 0.08, μascophoran = 0.06 (in units of million years). Note that BiSSE is prone to type I errors (Rabosky and Goldberg, 2015) but that we actually cannot soundly reject the null hypothesis, given the posterior speciation and extinction rate distributions and are hence on safe ground. Ancestral state reconstruction for the frontal shield states and BiSSE analyses for the alternative Bayesian tree (Supplementary Figs. S5 and S6) are highly comparable with those estimates from the ML tree (Figs. 3 and 4).
4. Discussion

It has long been known that molecular and morphological approaches (the latter including fossil taxa) must be simultaneously embraced for robust phylogenetic inferences (Pyron, 2015). In this contribution, we have taken a substantial step in contributing new molecular data and a greatly expanded and robustly supported phylogeny for an understudied but ecologically and evolutionarily important phylum (Pages-Escoff and Costello, 2020). Although we are interested primarily in New Zealand cheilostome bryozoans for reasons stated in our introduction, we have also now filled out numerous previously unsampled parts of the global cheilostome tree (compare Orr et al., 2020 with Fig. 2).

4.1. Higher-level cheilostome systematics needs revision

Cheilostome systematics is in a state of flux as molecular studies, coupled with the introduction of genome-skimming, are starting to take off for this diverse clade (Orr et al., 2019a, b, 2020). In providing a broadly sampled and robustly supported framework to evaluate evolutionary hypotheses we find that less than half of the 29 currently recognized families for which we have multiple genera represented are phylogenetically coherent. Our result emphasizes that much of the current family and higher-level bryozoan systematics, based largely on morphology, is unreliable, and further corroborates previous studies with statistically well-supported, but less broadly sampled, phylogenies (Orr et al., 2019a, b, 2020). One implication of this observation is that higher-level systematics (involving families, superfamilies and sub-orders) likely require substantial revision. We have hence refrained from detailing the mismatches of higher-level systematics (Bock, 2020) prematurely, but highlight new evolutionary hypotheses that have emerged, that are potentially supportable by morphological traits, given our molecular inferences (Fig. 2, Supplementary Figs. S3 and S4).

Notwithstanding some discrepancies between morphology-based hypotheses (Bock, 2020) and molecular data (this study), there is frequently mutual support. Take, for example, the basal grouping of Scrupariidae (Scruparia) as sister taxa to Electridae (Electra), Membraniporidae (Biflustra and Membranipora) and Aeteidae (Aetea) plus Steginoporellidae (Steginoporella) and Calpensiidae (Calpensia) (Fig. 2): these families are understood to have acquired different reproduction patterns (non-brooding in Membraniporidae and Electridae; different modes of embryonic incubation in the remaining families).
independently from the rest of incubating cheilostomes (Ostrovsky, 2013, 2020). Our tree now corroborates this hypothesis with full bootstrap and posterior probability support. In addition, our analysis resurrects Calpensiidae (Canu and Bassler, 1923), as sister clade of Steginoporellidae, suggesting that Calpensia also broods its embryos in an internal sac as does Steginoporella (Ostrovsky, 2013), and supports the hypothesis of multiple independent evolution of internal brooding (e.g. in Chaperiidae, Inversiulidae, Watersiporidae, Cryptosulidae, Urcellioridae) (e.g. Ostrovsky et al., 2006; Ostrovsky et al., 2009b) and placentation (e.g. in Bugulidae + Beaniidae, Catenicellidae, Watersiporidae, Urcellioridae, and Celleporella hyalina) (e.g. Ostrovsky et al., 2009a; Ostrovsky et al. 2016).

A closely positioned clade formed by Monoporella (Monoporellidae) and Macropora (Macroporidae) shares the presence of large ooecia (Fig. 1C), exceptionally able to incubate several embryos at the same time, that evolved from basally articulated spines or costae (e.g. Ostrovsky, 2013) (but see next paragraph). The fully supported Arachnopusiidae (Arachnopusia + Foveolariidae (Foveolaria and Odontioneilla) relationship is not indicated in present classification schemes (Bock, 2020), as species of Arachnopusia have an ascophoran state (Fig. 1F), while the Foveolariidae has an anascan state. However, we note that not only is the arachnopusid frontal shield a straightforward structure to form (unlike other ascophoran structures), but some species in Arachnopusiidae (e.g. A. gigantea) are anascan-like, where the frontal shield is practically non-existent (Hayward, 1995).

4.2. A need for even broader taxon sampling to fill gaps

Our ML (Fig. 2, Supplementary Fig. S3) and Bayesian (Supplementary Fig. S4) trees are largely in agreement with only one clade demonstrating incongruence. This is the fully supported clade comprising Valdemunitella (currently Calloporidae), Figularia (currently Cribrilinidae) and Euthyroides (currently Euthyroidiidae). Based only on morphology, we might have hypothesized that the Valdemunitella clade (based on 4 species represented by 6 colonies) is closely associated with other representatives of the family Calloporidae (e.g. Crassimarginatella or Callopora), but neither of our trees inferred this position. Rather, our ML tree places this clade (including Figularia and Euthyroides, both currently belonging to other families) in a position close to Monoporellidae and Macroporidae (see paragraph above) and our Bayesian tree places it in a more derived position. However, note that nodes sub-tending this clade in both trees are poorly supported. Rather than speculating on evolutionary and/or morphological arguments for either or both of these placements, we argue this indicates that there are many crucial unsampled taxa that would potentially allow a more robust placement of this clade, such as other cribrilinids in addition to Figularia and other calliporids such as Cauloramphus which, similarly to Valdemunitella, has spines encircling the frontal uncalcified membrane, forming a costate shield in some species (Dick et al., 2011). In the event, Valdemunitella, Figularia and Euthyroides are morphologically united, not by a costate shield, but by identical bilobate ooecia with a median suture (e.g. Ostrovsky, 2013), and the presence of vicarious avicularia in most of their species.
4.3. The evolution of the cheilostome frontal-shield

Historical studies of cheilostome body-wall development and morphology led to the conclusion that ascophoran frontal shields were phylogenetically informative (Banta, 1970; Gordon and Voigt, 1996; Sandberg, 1977). Our results substantiate the observation that characters considered to have deep phylogenetic information such as frontal shields are more evolutionarily labile than previously thought, and sometimes may even be convergent rather than homologous traits (Knight et al., 2011; Orr et al., 2019a). It has already been suggested, for instance, that anascan and ascophoran states, respectively regarded as stemward and crownward, have evolved more than once (Dick et al., 2009; Gordon, 2000; Waeschenbach et al., 2012). We show here that the anascan state is basal in the cheilostome tree and that the change from an anascan to ascophoran state has occurred multiple times independently (seven times in Fig. 3), hence likely more times in the history of cheilostome evolution, given our taxon sampling, which is far from complete. It is also striking that an ascophoran-state never reverts back to the anascan state, suggesting that it is evolutionarily more problematic to evolve a more complex calcified skeleton, but that once this structure is in place, it has not been lost again (Fig. 3). This is the “reverse” of what is the modern view of Dool’s law (Simpson 1953) where complex structures cannot be re-evolved (e.g. Collin and Miglietta 2008), where the morphological apparently “simpler” state (anascan) is not “gained” by the loss of the “complex” state (ascophoran). This could be due to genetic or developmental constraints (Smith et al., 1985), and/or because the advantages conferred by a calcified frontal shield vastly outweigh its disadvantages. Testing a classic idea that morphological complexity may predict diversification rates (Schoff et al., 1975), we found that the (potentially) more morphologically complex ascophoran-grade cheilostomes do not have distinguishable speciation and extinction rates compared with anascan-grade ones.

The frontal shield clearly contains phylogenetic information, but more research is needed to understand when it is informative, and why. As a further example, frontal shields produced by different developmental processes (e.g., lepralioid or umbonuloid (Hayward and Ryland, 1999; Martha et al., 2020; Taylor, 2020)) leave such distinct morphological tell-tale signs that it was commonly assumed that members within families constituted only a single type of frontal shield development. Our tree, however, places ascophoran taxa with lepralioid frontal shields (e.g., Powellitheca/Cyclicopora; Celleporina, Galeopsis, Osthomisoa) and umbonuloid ones (Exochella; Celleporaria) in the same clades (Fig. 2), as already shown to a lesser extent in earlier extensive studies (Dick et al., 2009; Orr et al., 2019a; Waeschenbach et al., 2012). Yet, at the more derived part of our inferred tree, the structure of the frontal shield seems to be more phylogenetically informative than seemingly simpler characters, such as the liruula (Beming et al., 2014). This is an anvil-shaped tooth-like structure projecting from the orifice that functions in water compensation. Specifically, the clade containing Parminitina to Hemismitinidea (containing four and three genera of the families Celleporidae and Bitectiporidae respectively) has a non-pseudoporous umbonuloid frontal shield (Gordon, 2000), while the next one containing Schizastenitina to Bivicopora (containing two smittiniid and two bitectiporid genera) has a pseudoporous lepralioid shield. The presence of a liruula seems haphazard among these genera, where those in the Smittiniidae have liruula and those in the Bitectiporidae have a sinus (Fig. 1J, K). Our tree suggests new ways of partitioning some of the families and genera of Smittiniidea, which unexpectedly also includes the genera Porella (Bryocryptellidae) and Ostrurkova (Umbonulidae). To summarize, it is clear that a much more thorough and systematic investigation of the development and evolution of frontal shields, and greater taxonomic sampling, is necessary for a deeper understanding of ascophoran cheilostomes.

4.4. Molecules suggest morphological hypotheses and pinpoint research needs

Another example of traits thought to be phylogenetically related and hence informative is the sinus versus the ascopore, pertaining to the ascophoran plumbing system. Because Microporella, Fenestrulina and Calloporina all have ascopores, they were historically united in the Microporellidae. A previous molecular study has clearly shown that Fenestrulina does not belong in the same clade as Microporella (Orr et al., 2019b). Here, we give molecular support to the hypothesis that Calloporina is a more microporellid and further suggest that Chiastosella (having a sinus, currently belonging to the Escharinidae; Fig. 1H) and Calloporina (having a slit-like ascopore; Fig. 1I) belong in the same clade, a relationship supported also by their shared distinctive ooeia (Brown, 1954; Cook et al., 2018). Supporting the long-held hypothesis that an ascopore should evolve by the cutting-off of a sinus, Chiastosella should be basalwards of Calloporina (Cook et al., 2018, p. 218). This is supported by our tree, which also suggests that Chiastosella may be paraphyletic with respect to Calloporina. In multiple cases, taxa that are considered unique or unusual have placed in phylogenetic positions that suggest hypotheses of their evolutionary relationships based on morphology. For instance, Rhabdozoum, currently placed in its own family because of its highly distinctive morphology, is basal to Candidae, suggesting that they are closely related and that Candidae sensu stricto may have been derived from a Rhabdozoum-like ancestor. In fact, the initial zooid of the colony (ancestrula) of Rhabdozoum resembles those in some Scrupocellaria species and several of its mature zooidal features such as its ooeia, frontal avicularia and spines are reminiscent of species of Amastigia and Meriopia (all Candidae s.s.). Margareta, another rather distinct genus, is in a family with only one other monospecific genus (Tubucella). Here, Margareta is inferred to be basal to Catenicellidae, suggesting that Catenicellidae s.s. may have been derived from a Margareta-like ancestor, although it has always been thought that catenicellids are derived from cibrimorphs (Gordon, 2000; Gordon and Braga, 1994). Much research is required to unravel the mystery of this grouping, given that they are both so distinctive, sharing apparently only rhizoids, rootlets fixing the colony to the substrate which have independently evolved multiple times in all major orders of marine bryozoans (Schack et al., 2019). Note that we infer two distinct clades of Catenicellidae, one represented by Catenicella and Cymbicellaria, which are vittate (frontal pore chambers are long and narrow) and the second including Orthoscuticella and Pterocella, which are foraminate (frontal shield has numerous windows in the gymnocyct; Fig. 1G). Yet another example is the erect and branching calweliid Malakosaria whose zooidal features resemble Fenestrulina (Fenestrulinidae), the genus in which Malakosaria nests in our tree.

One taxonomically challenging family deserves special mention. The speciose Celleporidae, with at least 252 described living taxa globally, is mostly characterized by nodular/massive colonies as a result of rapid frontal budding (the building of zooids on top of existing ones). As a consequence, autozooids are somewhat irregularly disposed and difficult to characterize morphologically. These genera are currently distinguished by the morphology of their ooeia (development of endoecium/tabula) and orifices (always sinuate but the sinus varies from a narrow slit to a broad and shallow concavity). Genus-level hypotheses based on these characters are problematic as indicated by our tree, in which Celleporina, Galeopsis (Fig. 11) and Osthomisoa are non-monophyletic. Buffonellaria is excluded from the family and allied with Buffonellodidae, whereas Celleporaria, historically included in Celleporidae but subsequently split off because of its umbonuloid frontal shield (Harmer, 1957; Cook et al., 2018, p. 182), is reinstated.

4.5. Lower-level cheilostome systematics are very robust

Although higher-level systematics are in need of revision, we report
that lower-level morphological hypotheses (i.e., species and genera) are very robust, supporting inferences based on common-garden experiments, to put forward the idea that “morphological species” are as good as “genetic species” in cheilostome bryozoans (Jackson and Cheetham, 1990). While Jackson and Cheetham experimented only with a handful of species, we now confirm their hard-earned insight implies that many more species and genera can be treated as distinct evolutionary lineages. This is an important result as many evolutionary and paleontological studies use morphospecies or even morpho-genera as the unit of analyses (Alroy, 2010; Heim et al., 2015). We also note that there are many New Zealand species in our tree that are yet undescribed (c. 20% of those newly sequenced here), indicating that continued exploration in the EEZ of New Zealand is crucial even for such a geographically well-characterized marine clade.

5. Conclusions

Our work shows that lower-level taxonomic sampling in phylogenetics is vital for understanding higher-level systematics, especially in an understudied group like cheilostome bryozoans. While we have contributed a substantial number of sequences from diverse species, many more must be included for the phylogenetic inferences and reliable systematic groupings for cheilostomes. By contributing molecular data and robustly supported phylogenetic inferences, we have supplied the basis for evolutionary (including phylogenetic) hypotheses that can be further examined. Once we are confident in the topology of at least parts of the cheilostome tree, we can start asking further questions on evolutionary processes.

CRediT authorship contribution statement

R.J.S. Orr: Data curation, Formal analysis, Investigation, Methodology, Project administration, Resources, Software, Validation, Visualization, Writing - original draft, Writing - review & editing. D.P. Gordon: Validation, Writing - original draft, Writing - review & editing. M.H. Ramsfjell: Methodology, Writing - review & editing. H.L. Mello: Resources, Writing - review & editing. A.M. Smith: Resources, Writing - review & editing. L.H. Liow: Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Methodology, Project administration, Resources, Validation, Visualization, Writing - original draft, Writing - review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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