Taxonomy and phylogenetic relationships of the coral genera *Australomussa* and *Parascolymia* (Scleractinia, Lobophylliidae)

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Abstract

Novel micromorphological characters in combination with molecular studies have led to an extensive revision of the taxonomy and systematics of scleractinian corals. In the present work, we investigate the macro- and micromorphology and the phylogenetic position of the genera *Australomussa* and *Parascolymia*, two monotypic genera ascribed to the family Lobophylliidae. The molecular phylogeny of both genera was addressed using three markers, the partial mitochondrial COI gene and the nuclear histone H3 and the ribosomal ITS region. Based on molecular data, *Australomussa* and *Parascolymia* belong to the Lobophylliidae and they cluster together with the genera *Lobophyllia* and *Symphyllia* within the same clade. While *A. rowleyensis* and *P. vitiensis* are closely related based on the three gene regions examined, their macro and micromorphology suggest that these species are distinct, differing in several characters, such as continuity and thickness of the costosepta, the number of septa, septal tooth height, spacing, and shape, and the distribution and shape of granules. Thus, we revise the taxonomic status of the genus *Australomussa* as a junior synonym of *Parascolymia*.

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Introduction

Over the last two decades, our understanding of the evolution and the systematics of hard corals (Cnidaria, Anthozoa, Scleractinia) has rapidly advanced due to the progressive increase of molecular studies (Romano and Pulumbi, 1996; Chen et al., 2002; Fukami et al., 2004, 2008; Kitahara et al., 2010; Stolarski et al., 2011). The new molecular phylogenies are, however, often very different from phylogenies based on macro-morphology (Fukami et al., 2004, 2008; Budd and Stolarski, 2009, 2011; Huang et al., 2011). Several recent papers integrating molecular and morphological approaches have led to formal taxonomic revisions of scleractinian corals at different taxonomic ranks (e.g. Wallace et al., 2007; Gittenberger et al., 2011; Kitahara et al., 2012, 2013; Benzoni et al., 2012a, 2014; Hoeksema, 2014; Kitano et al., 2014; Huang et al., 2014a, b). This integrated approach has proved effective at resolving long-standing issues, for example a comprehensive revision of the taxonomy and systematics of 23 nominal species of *Psammocora* Dana, 1846 (Stefani et al., 2008; Benzoni, 2006; Benzoni et al., 2010, 2012b) and 21 nominal species of *Pocillopora* Lamarck, 1816 (Schmidt-Roach et al., 2012, 2013, 2014).
The stony coral family Lobophylliidae Dai and Horng, 2009 has recently been studied by several authors using an integrated morpho-molecular approach, and is undergoing several taxonomic changes, although this process is far from complete. For example, Indo-Pacific taxa traditionally ascribed to the Mussidae Ortmann, 1890 have been moved to the Lobophylliidae as a result of the molecular work by Fukami et al. (2004, 2008), and the finding of a deep divergence between Indo-Pacific and Atlantic species based on morphological characters (Budd and Stolarski, 2009; Budd et al., 2012). The family Lobophylliidae is now comprised of the genera Lobophyllia de Blainville, 1830, Acanthastrea Milne Edwards and Haime, 1848, Cynarina Brüggemann, 1877, Echinophyllia Klunzinger, 1879, Homophyllia Brüggemann, 1877, Micromussa Veron, 2000, Moseleya Quelch, 1884, Oxypora Saville Kent, 1871, Parascolymia Wells, 1964 and Symphyllia Milne Edwards and Haime, 1848 (Budd et al., 2012). Also included in the family are two genera that have not been examined at a molecular level, namely Echinomorpha Veron, 2000 and Australomussa Veron, 1985, hence their phylogenetic placement is unresolved.

The macromorphology (budding, colony form, size and shape of corallites, numbers of septal cycles), the micromorphology (shapes and distributions of septal teeth and granules), and the microstructure (arrangement of calcification centres and thickening deposits within costosepta) of the lobophylliid genera Acanthastrea, Cynarina, Echinophyllia, Homophyllia, Lobophyllia, Micromussa, Oxypora, Parascolymia, and Symphyllia were examined by Budd and Stolarski (2009) and Budd et al. (2012). They concluded that the shape and distribution of septal teeth and granules, the area between teeth, and the development of thickening deposits are informative characters for distinguishing the Lobophylliidae from representatives of the other coral families. Arrigoni et al. (2014) presented a comprehensive molecular phylogeny that shows that the Lobophylliidae is a monophyletic family comprising nine main molecular clades (clades A-I), and that several genera are not monophyletic. The authors also showed that the monospecific genus Parascolymia belongs to clade I (sensu Arrigoni et al., 2014) together with all the species of Lobophyllia and Symphyllia for which molecular data is available, including the two type species Lobophyllia corymbosa (Forskål, 1775) and Symphyllia radians Milne Edwards and Haime, 1849. The authors did not, however, undertake any formal taxonomic revision of the status of the genus Parascolymia.

Australomussa rowleyensis Veron, 1985 was described from Western Australia and ascribed to the Mussidae. It is a colonial and zooxanthellate scleractinian coral, characterized by flattened, helmet- or dome-shaped coralla, valleys approximately 20mm wide, with very thick walls and a well-developed columella (Veron, 1985). In the original description of A. rowleyensis, Veron (1985) stated that this genus showed ‘little resemblances to any other genus’ with the exception of Parascolymia and Symphyllia, and ‘its closest affinities are probably with the former’. The author referred only to the macromorphology of the coralla for the comparison of Australomussa with Parascolymia and Symphyllia and did not consider any micromorphological characters. Budd and Stolarski (2009) and Budd et al. (2012) showed that the majority of macromorphological characters traditionally used in the taxonomy and systematics of Lobophylliidae and Mussidae exhibit homoplasy. In contrast, novel micromorphological characters separate these two families and are useful for the description and formalization of species. Nevertheless, while the micromorphology of P. vitiensis (Brüggemann, 1877) was described by Budd and Stolarski (2009) and Budd et al. (2012), A. rowleyensis was not analysed in these studies.

The known distribution of A. rowleyensis includes the Western Pacific region known as the Coral Triangle (for definition see Hoeksema, 2007; Veron et al., 2009) and partially overlaps with the distribution of P. vitiensis which is absent from Western Australia but extends to the west in the Indian Ocean and to the east in the central Pacific (Veron, 2000). Australomussa rowleyensis and P. vitiensis have very different histories of nomenclature. Perhaps due to its recent description and rarity (Veron, 1985), A. rowleyensis has always been described as A. rowleyensis despite its morphological similarity to Parascolymia and Symphyllia (Veron, 1985, 2000). Conversely, P. vitiensis has a long history of nomenclatural confusion. It was originally ascribed to Litophyllia Milne Edwards and Haime, 1857 (Gardiner, 1899; Crossland, 1952) and later described as Protolobophyllia japonica Yabe and Sugiyama, 1935. In agreement with Matthai (1928), Wells (1937) and Vaughan and Wells (1943) considered Scolymia Haime, 1852 and Protolobophyllia Yabe and Sugiyama, 1935 as junior synonyms of Lobophyllia, presuming that the solitary forms were juvenile monostomatous stages of this
of a large series of mono- to polystomatous specimens (1B-D, 2F-I). The macro-morphologic observation of 1975; Veron and Pichon, 1980: figs 416-417) (Figs can also form polystomatous coralla (Chevalier, et al., 2004, 2008; Budd and Stolarski, 2009).

Although P. vitiensis is generally monocentric, it can also form polystomatous coralla (Chevalier, 1975; Veron and Pichon, 1980: figs 416-417) (Figs 1B-D, 2F-I). The macro-morphologic observation of a large series of mono- to polystomatous specimens of P. vitiensis from Papua New Guinea and New Caledonia and the similarity of the larger specimens with A. rowleyensis prompted the detailed study of the morphological affinities and molecular relationship between these two species and the two monospecific genera they are currently ascribed to.

Here we selected three DNA regions, the barcoding region of cytochrome oxydase subunit I gene, the nuclear ribosomal ITS region. In addition, the most comprehensive phylogeny reconstruction of the Lobophylliidae to date is based on these two markers (Arrigoni et al., 2012). The latter locus was revealed to be informative for a broad-based phylogeny of the Merulinidae Verrill, 1865 (Huang et al., 2011, 2014b), a family closely related to the Lobophylliidae (Fukami et al., 2008; Gittenberger et al., 2011; Huang et al., 2011; Benzoni et al., 2011, 2014) and, moreover, the most comprehensive phylogeny reconstruction of the Lobophylliidae to date is based on these two markers (Arrigoni et al., 2014). The latter locus was revealed to be informative for a broad-based phylogeny of the Merulinidae Verrill, 1865 (Huang et al., 2011, 2014b), a family closely related to the Lobophylliidae (Fukami et al., 2008; Arrigoni et al., 2012). Several phylogenetic studies of scleractinian corals achieved a well-resolved phylogeny using a concatenated species-tree, combining mitochondrial and nuclear molecular markers (Huang et al., 2009, 2011; Souter, 2010; Gittenberger et al., 2011; Benzoni et al., 2012a; Richards et al., 2013; Arrigoni et al., 2014). This kind of approach is a powerful way to obtain a robust phylogeny, resolving all key nodes and yielding good resolution at species level.

In the present paper, the phylogenetic relationships of A. rowleyensis with the rest of the Lobophylliidae are explored for the first time on the basis of three molecular loci, the barcoding region of cytochrome oxydase subunit I gene, the nuclear histone H3, and the nuclear ribosomal ITS region. In addition, we examined the macromorphology and micromorphology of A. rowleyensis and compared it to P. vitiensis.

Material and methods

Sampling

Specimens of Parascolymia vitiensis for this study were sampled in New Caledonia, Papua New Guinea, and Eastern Australia, while samples of Australomus-sa rowleyensis were collected in the Kimberley, North-West Australia (S1). Coral specimens were photographed and collected while SCUBA diving from 2 to 35 meters depth. Digital images of living corals in the field were taken with a Canon Powershot G9 in an Ikelite underwater housing system in New Caledonia and Papua New Guinea (Figs 1 and S5), and with an Olympus XZ1 in a PT-050 underwater housing in Australia (Figs 1 and S5). Coral specimens were collected, tagged, and preserved in 95% ethanol for further molecular analysis. After the sampling of fixed tissues for DNA extraction, each corallum was immersed in sodium hypochlorite for 48 hours to remove all soft parts, rinsed in freshwater and dried for microscope observation. Specimens were identified at the species level based on skeletal morphology using a Leica M80 microscope following the descriptions and illustrations by Chevalier (1975), Veron and Pichon (1980), and Veron (1985).

Morphological analyses

Images of coral skeletons were taken with a Canon G5 digital camera and through a Leica M80 microscope equipped with a Leica IC80HD camera. For high resolution and deep field close ups of three-dimensional details of corallites and septa, a series of images of the same subject at different focus intervals were taken (approximately 10) and the images were fused using the Helicon Focus 5.3 software (Kozub et al., 2000-2012). To compare macromorphology and micromorphology of P. vitiensis and A. rowleyensis we used a subset of 21 characters from Budd et al. (2012) (Table 1). We
Fig. 1. *Parascolymia vitiensis* (A-D) and *P. rowleyensis* (previously *Australomussa*) (E-G) *in situ*: A) IRD HS2984 (monostomatous); B) UNIMIB PFB056 (bistomatous); C) IRD HS3139 (polystomatous, same as in Fig. 2H); D) UNIMIB PFB057 (polystomatous, same as in Fig. 2I); E) WAM Z65786; F) WAM Z65785; G) close up of the same colony as in E; H) close up of the same colony as in F.
adopted their character name, ID number (in brackets) and state names and, when relevant quantitative differences between the two species were observed within a character state, this information was added after it. Given the large size of the skeletal structures in *P. vitiensis* and *A. rowleyensis*, the majority of macromorphological and micromorphological characters considered in this study, with the notable exception of characters 43 (granule shape and distribution) and 44 (interarea) from Budd *et al.* (2012), were examined using light microscopy. Scanning Electron Microscopy (SEM) was used to analyze the shape and distribution of granules on septal faces and the interarea of teeth on representative specimens of *P. vitiensis* (UNIMIB PFB151) and one of *A. rowleyensis* (WAM Z65789). Specimens were mounted using silver glue, sputter-coated with conductive gold film and examined using a Vega Tescan Scanning Electron Microscopy at the SEM Laboratory, University of Milano-Bicocca.

For a glossary of skeletal terms we refer to Budd *et al.* (2012).

Abbreviations:

CC1 IRD CoralCal1 Expedition, Côte Oubliée, New Caledonia, 2007

CC4 IRD CoralCal4 Expedition, New Caledonia, IRD, 2012

CCAP IRD CoralCap Expedition, New Caledonia, 2007

Cs cycle of costosepta

IRD Institut de Recherche pour le Développement, Nouméa, New Caledonia

NIUGINI Niugini Biodiversity Expedition, Papua New Guinea, 2012

RMNH Naturalis Biodiversity Center (former Rijksmuseum van Natuurlijke Historie), Leiden, the Netherlands

S cycle of septa

UNIMIB Università di Milano-Bicocca, Milan, Italy

WAM Western Australian Museum, Perth, Australia

In the list of examined material for IRD specimens the station number (ST) is provided, when available, after the sampling locality. Station numbers can be searched in the IRD online database LagPlon (http://lagplon.ird.nc/consultv2_5/rechSimple.faces) where additional details on the reef habitat, GPS coordinates, and a map of each station can be found.

Molecular analyses

Whole genomic DNA was extracted from tissue samples using the DNeasy Blood and Tissue kit (Qiagen Inc., Valencia, CA, USA) according to the manufacturer’s protocols. DNA concentration of extracts was quantified using a Nanodrop 1000 spectrophotometer (Thermo Scientific, Wilmington, DE, USA). A total of three molecular markers were amplified and sequenced for the majority of the specimens (S1): (1) a ~750 bp fragment of the cytochrome oxidase subunit I gene (COI) from mitochondrial DNA, (2) a ~350 bp portion of the nuclear histone H3, and (3) a ~800 bp portion of the ITS region, including the 3’ end of 18S, the entire ITS1, 5.8S, and ITS2, and the 5’ end of the 28S, as nuclear loci. COI was amplified using MCOIF-MCOIR primers (Fukami *et al.*, 2004) and the protocol by Benzoni *et al.* (2011), the histone H3 using H3F-H3R primers (Colgan *et al.*, 1998), and the ITS region using ITS4 (Takabayashi *et al.*, 1998) - A18S (White *et al.*, 1990) primers and the protocol by Benzoni *et al.* (2011), or alternately using 1S and 2SS primers (Wei *et al.*, 2003) and the protocol by Kitano *et al.* (2014). Sequencing was carried out by Genomics and Bioscience and Technology Co., Ltd, Xizhi City, Taipei County, Taiwan. Sequences obtained in this study have been deposited in EMBL, and accession numbers are listed in S1.

Sequences were viewed, edited and assembled using CodonCode Aligner 4.2.5 (CodonCode Corporation, Dedham, MA, USA) and manually checked using BioEdit 7.2.5 (Hall, 1999). Alignments of the four separated datasets (three single gene trees and one concatenated) were carried out using the E-INS-i option in MAFFT 7.110 (Katoh *et al.*, 2002; Katoh and Standley, 2013) under default parameters. *Plesiasteria versipora* ( Lamarck, 1816) and several species from the family Merulinidae were selected as outgroups due their divergence from the family Lobophylliidae (Fukami *et al.*, 2008; Kitahara *et al.*, 2010; Benzoni *et al.*, 2011). Indels, invariable, and parsimony informative sites were detected with DnaSP 5.10.01 (Librado and Rozas, 2009) and Indels were treated as a fifth character in phylogenetic analyses. Genetic distances and their standard deviation were calculated as p-distance with 500 bootstrap replicates using MEGA 5.2 (Tamura *et al.*, 2011). To reconstruct the single gene trees Bayesian Inference (BI) and Maximum Likelihood (ML) analyses were used as implemented in MrBayes 3.1.2 (Ronquist and Huelsenbeck, 2003) and PhyML 3.0 (Guindon and
Fig. 2. Corallum morphology in *Parascolymia vitiensis*: A) IRD HS3255; B) IRD HS2955; C) detail of the same specimen in B showing variability in shape and size of septal teeth; D) UNIMIB PFB031; E) IRD HS2985; F) UNIMIB PFB056; G) UNIMIB PFB055; H) IRD HS3139; I) UNIMIB PFB057. All specimens are in the phylogenetic trees in Figs 9, S1-S3.
Fig. 3. Corallum morphology in *Parascolymia rowleyensis* (previously *Australomussa*): A) WAM Z65785; B) WAM Z65788; C) WAM Z65786; D) lamellar linkage between centres of adjacent corallites in same specimen as in B; E) corallite polymorphism in the same specimen as in A; F) size and shape of costosepta in specimen WAM Z65787. All specimens are in the phylogenetic trees in Figs 9, S1-S3.
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Fig. 4. Comparison of the macromorphology of *Parascolymia vitiensis* (A-C) and *P. rowleyensis* (D-E): A) septa in the monocentric specimen IRD HS2964; B) top view of the costopecta in the polycentric specimen UNIMIB PFB057; C) side view of the same portion of the specimen in B; D) peripheral calices in specimen WAM Z65786; E) top view of the costopecta in the same specimen as D; F) side view of the same portion of the specimen in E. Red arrows indicate the position of the columella in adjacent corallites, red brackets placed perpendicularly to the costopecta show the number of costopecta intercepted by a 1 cm transect.

Fig. 5. SEM images of radial elements of *Parascolymia rowleyensis* (previously *Australomussa*) (WAM Z65789: A-B) and *P. vitiensis* (UNIMIB PFB151: C-D): A) top view of septa reaching the wall; B) side view of septa of different cycles showing some variability in septal teeth size between cycles but overall homogeneous shape; C) top view of septa reaching the wall; D) side view of septa of different cycles showing high variability in septal teeth size and shape between cycles.
Gascuel, 2003), respectively. The best-fit substitution model for each locus was determined using the Akaike Information Criterion (AIC) as implemented in MrModeltest 2.3 (Nylander, 2004) in conjunction with PAUP4.0b10 (Swofford, 2003). As most suitable models AIC selected the General Time-Reversible (GTR) model with a proportion of sites being invariable (+I) and the remainder following a gamma distribution (+I) for COI and rDNA, and the Kimura (K80) model with a proportion of invariable sites (+I) for histone H3.

The Maximum Likelihood (ML) tree was calculated with PhyML and a total of 500 bootstrap replicates were performed to assess the robustness of each clade. Four independent Markov Chain Monte Carlo (MCMC) runs were conducted for 1.4 × 10^7 generations for COI dataset (1.7 × 10^7 generations for histone H3 and 4 × 10^7 generations for ITS region) with trees sampled every 100 generation for each analysis. The 25% first trees were discarded as burn-in, and posterior probabilities were estimated from the remaining trees in each run (10,500 remaining trees for COI, 12,750 for histone H3, and 30,000 for ITS region). To determine if the runs had achieved stationarity, we visualized log-likelihood scores and model parameter values across each run using Tracer 1.5 (Rambaut and Drummond, 2007). Finally, the three single gene datasets were concatenated in a single partitioned alignment and the phylogeny was reconstructed using Bayesian Inference and Maximum Likelihood analyses. Four independent Markov Chain Monte Carlo (MCMC) runs were conducted for 2.2 × 10^7 generations with trees sampled every 100 generation and the 25% first trees were discarded as burn-in. The ML tree was built in PhyML and a total of 500 bootstrap replicates were performed to assess the robustness of each clade. Branches with >70% bootstrap support values and >0.90 posterior probabilities are considered significantly supported.

Results

Macromorphology

In *P. vitiensis* coralla can be solitary (Figs 1A, 2A-E) or colonial (Figs 1B-D, 2F-I) and formed by intracalicular and extracalicular budding (e. g. Fig. 2I). In colonial coralla, as a result of circumoral budding, coralites are highly polymorphic (Fig. 2G-I) and corallite integration is uni- or multiserial. Corallum shape is generally flattened or concave (Figs 1-2). Calice or valley width is larger than 2.5cm (Fig. 2) and variable. In some specimens the central part of the calice can have a shallow depression (Fig. 2D-F, H). Continuity of costosepta is mostly confluent in di-tricentric coralla (Fig. 2F-G), but becomes mostly not confluent in polycentric coralla (Fig. 2I; Veron and Pichon, 1980: Fig. 417). There are six cycles of septa in the calices, rarely seven (Fig. 2E; Chevalier, 1975), those of the sixth are free. Septa spacing is large, with 4-5 septa per 5mm (Fig. 4A-B). Relative costosepta thickness between Cs1 and Cs2 versus Cs3 is unequal (Fig. 5C-D). In polycentric coralla linkage between centres of adjacent corallites within series is lamellar (Fig. 4B-C). Columella trabecular and spongy (indicated by arrows in Fig. 4B-C) and its size relative to calice width is less
than 1/5 (Fig. 2). The endotheca is vesicular (Fig. 6A).

In *A. rowleyensis* coralla are flattened or massive and ‘helmet- or dome-shaped’ (Fig. 1E-H). Coralla are colonial as a result of primary circumoral budding and both intra and extracalicular budding occur (Fig. 3D-E). Corallites display polymorphism in smaller colonies where the central corallite is still larger as in the paratype WAM 173-84 (Veron, 1985: Fig. 25) and corallite integration is uni- or multiserial. Calice or valley width is large according to the character state in Budd *et al.* (2012) but smaller than 2.5cm (Fig. 4D-F). Calices at the periphery of the coralla can be inclined and the part of their calice which is not adjacent to other calices can be wide (Fig. 3A-B, 6D). Continuity of costosepta is mostly confluent (Fig. 3A-E, 4E-F). There are four cycles of septa (Fig. 3D), those of the fourth are free. Septa spacing is large, with five septa per 5mm (Fig. 4D-E). Relative costosepta thickness between Cs1 and Cs2 versus Cs3 is slightly unequal (Fig. 5A-B). Linkage between centres of adjacent corallites within a series is lamellar (Fig. 4D-F). Columella are trabecular and spongy (indicated by arrows in Fig. 4E-F) and the size relative to calice width less than or equal to 1/4 of calice width (Figs 3D-E, 4D-F). The endotheca is vesicular (Fig. 6B-C).

**Micromorphology**

In *P. vitiensis* tooth base at mid-septum is elliptical in shape and parallel to the direction of the septum (Fig. 5C-D). Tooth tips are irregular and overall mainly lobate (Figs 5D, 7A-B, D). Teeth on S1 are 1mm or higher (Figs 4C, 7) and their spacing is very wide, with adjacent teeth more than 2mm apart. Tooth shape and size is very variable within and between septa (Fig. 7) as also noted by previous authors (Chevalier, 1975; Veron and Pichon, 1980) with some teeth becoming round in section towards the tip and having and overall pointed, or spiniform Chevalier (1975), shape (Fig. 7C). Granulation on the side of septa is weak and granules are enveloped by thickening deposits (Fig. 7A-B). The inter-area structure is generally smooth (Fig. 5D) or with palisade. Tooth shape between Cs3 and Cs1 is unequal (Fig. 5C-D).
In *A. rowleyensis* tooth base at mid-septum is elliptical in shape and parallel to the direction of the septum (Fig. 5A-B). Tooth tips are irregular and lobate (Figs 5B, 8). Teeth on S1 range between 0.8-0.9 mm (Figs 4F, 8) and their spacing is wide, with adjacent teeth between 1 and 2mm apart. Granulation on the side of septa is strong and granules are scattered (Figs 5B, 8A). The inter-area structure has a palisade structure (Fig. 8A). Tooth shape between Cs3 and Cs1 is equal (Fig. 5B). In general, in this species tooth shape is not very variable within and between septa (Fig. 5A-B) especially when compared to the variability described in *P. vitiensis*.

**Molecular analyses**

The final alignment of COI data consisted of 580 bp, of which 48 were parsimony informative sites, with a total of 84 mutations. The aligned histone H3 matrix was 318 bp long with 86 parsimony informative sites and 122 mutations. The total alignment of ITS region was composed by 951 bp, 160 parsimony informative sites and 294 mutations. No intra-individual polymorphisms or double peaks were observed in the chromatograms of the two nuclear loci, thereby avoiding the need to clone the amplified fragments. The phylogeny reconstruction of the combined molecular data is in Fig. 9, while the three single gene trees are in the Supplementary Information (Figs S2-S4). Phylogenetic analyses under BI and ML criteria yielded congruent results, with no contrasting signals. Bayesian topologies with significant branch support indicated by ML bootstrapping support (MLs) and Bayesian posterior probability scores (BIs) are reported in Figs 9 and S2-S4.

The phylogram based on the concatenated (COI, histone H3, and ITS) molecular dataset shows high ML and BI supports at all key nodes (Fig. 9). Clade I sensu Arrigoni et al. (2014) contains all species of *Lobophyllia* and *Symphyllia* analyzed so far and all our sequences of *A. rowleyensis* and *P. vitiensis*. The latter two species group together in a strongly supported lineage (MLs = 100 and BIs = 0.9) and their genetic boundaries remain unclear being indistinguishable from each other with these molecular markers. The average genetic distance of *A. rowleyensis* from *P. vitiensis* is 1.1 ± 0.2%,
Fig. 9. Phylogenetic position of *Parascolymia vitensis* and *P. rowleyensis* (previously *Australomussa*) and their relationships within the family Lobophylliidae based on concatenated matrix (COI, histone H3, and ITS region). Bayesian topology is shown. Numbers associated with branches indicate Maximum Likelihood bootstrap (>70%) support (left) and Bayesian posterior probabilities (>0.9) (right). Clades within Lobophylliidae are coloured and labelled A to I according to Arrigoni et al. (2014).
while genetic variability within A. rowleyensis is 1.1 ± 0.3% and within P. vitiensis is 0.7 ± 0.2%. The majority of the other species in this clade, L. costata (Dana, 1846), L. robusta Yabe, Sugiyama and Eguchi, 1936, S. agaricia Milne Edwards and Haime 1849, S. radians, S. recta (Dana, 1846), and S. valenciennesii Milne Edwards and Haime, 1849 are recovered as monophyletic lineages. The only exceptions are represented by L. corymbosa and L. hemprichii (Ehrenberg, 1834) that are not monophyletic and nested in two distinct lineages.

The Bayesian COI topology (Fig. S2) indicates that all newly obtained sequences of A. rowleyensis and P. vitiensis are nested together with the genera Lobophyllia and Symphyllia within clade I (MLs = 94% and BIs = 0.93). While the two species are not monophyletic and they occur together in two main groups within clade I, the mitochondrial phylogenetic reconstruction is similar to that of the nuclear histone H3 (Fig. S3). Again, all newly obtained sequences of A. rowleyensis and P. vitiensis form clade I sensu Arrigoni et al. (2014) (MLs = 95% and BIs = -) together with several species of Lobophyllia and Symphyllia. Clade I is composed of 10 species represented by a total of 28 sequences, of which 26 share the same haplotype and they are thus identical, while the remaining two sequences differ from the others by only one bp substitution. Moreover, all of the Merulinidae subclades defined by Budd and Stolarski (2011) and Huang et al. (2011) are recovered with the exception of D/E. Interestingly, also in the family Lobophylliidae, all of the molecular clades defined by Arrigoni et al. (2014) based on COI and rDNA molecular markers, except F, are supported in our BI and ML analyses. The Bayesian topology obtained from the ITS region alignment is similar to both COI and histone H3 ones, but has a higher resolution at species level with significant supports for the majority of key nodes (Fig. S4). Again, all our sequences of A. rowleyensis and P. vitiensis are found together in a strongly supported group (MLs = 90 and BIs = 1) within clade I (Fig. S4). A similar situation is apparent for L. hemprichii and S. agaricia which occur in a strongly supported monophyletic group. The other Lobophyllia and Symphyllia species within clade I, i.e. L. costata, L. diminuta Veron, 1985, L. flabeliformis Veron, 2000, L. robusta, S. erythraea (Klunzinger, 1879), S. radians, S. recta, and S. valenciennesii, are recovered as monophyletic lineages, while the only specimen of Acanthastrea ishigakiensis Veron, 1990 is closely related to S. recta.

Discussion

In this study we explore the gross- and fine-scale morphology and the phylogeny of the two traditionally described monotypic genera Australomussa and Parascolymia. We provide a detailed description of diagnostic micromorphological characters of A. rowleyensis and P. vitiensis and we define the phylogenetic position of both these two species within the family Lobophylliidae using three molecular markers. As a result we propose a taxonomic revision for the genus Australomussa and we formally consider Australomussa as a junior synonym of Parascolymia.

Morphology of P. rowleyensis and P. vitiensis and consequences for taxonomy

The lack of genetic resolution between P. vitiensis and P. rowleyensis in all our molecular analyses might suggest that these two species are in fact synonyms. The skeleton morphology, however, indicates that although the two species share some macro- and micromorphologic character, they are morphologically distinct and they have a different state for 10 of the 21 characters used by Budd et al. (2012) (in bold in Table 1). Veron (1985) stated that Australomussa ‘differs from Symphyllia in having an initial central corallite which buds daughter corallites extracalicularly, in lacking meandering valleys (which some Symphyllia ecomorphs also lack) and in having widely separated series of centres without a true common wall between them’. However, he provided no detailed information on the morphologic characters that differentiate Australomussa from Parascolymia (=Scolymia). Our observations of the macro- and micromorphology of P. rowleyensis and P. vitiensis confirm that these species share a number of characters, namely intracalicular and extracalicular budding, corallite polymorphism associated with circumoral budding, uni or multiserial corallite integration, free septa, wide septa spacing with less than six septa per 5mm, discontinuous linkage between corallite centres (lamellar linkage), a trabecular spongy columella, a vesicular endotheca, septum tooth elliptical at the base, and irregular lobate tooth tips (Table 1). Nevertheless P. vitiensis has wider calices or series, a larger variability of continuity of costosepta over the wall, more cycles of septa, different relative costosepta thickness, a smaller columella size relative to calice width, higher and more widely spaced septum teeth, weakly developed septa granulation, a smoother inter-area structure, and unequal tooth shape between costosepta of different cycles.
(Table 1 in bold). We propose therefore that these morphological differences are sufficient to distinguish two species despite the fact that the unresolved genetic boundaries based on multiple markers strongly argue against retaining the species as distinct. Thus we formally consider Australomussa as a junior synonym of Parascolymia and retain *P. vitiensis* and *P. rowleyensis* as separate sister species.

In *P. vitiensis* the teeth in different septal cycles differ significantly in shape as already discussed by Veron (1985) and Budd and Stolarski (2009). In *P. rowleyensis* the teeth in different septal cycles do not differ significantly in shape as described by Veron (1985) in the species original description. The type specimen of *P. rowleyensis* displays an obvious variability of thickening of costosepta between specimens as remarked by Veron (1985). However, the variability of shape and size of septal dentation is far more reduced in this species than in *P. vitiensis*. One of the specimens of *P. rowleyensis* in the series we examined, Z65786, has relatively thin septa and costosepta and is similar in this respect to the paratype WAM 173-84 (Veron, 1985: Fig. 25). The remainder have a similar thickness of costosepta to the holotype WAM 171-84 (Veron, 1985: Fig. 23). However, none of the specimens we examined in this study has radial elements as thick as paratype WAM 172-84 (Veron, 1985: Fig. 24). The thickness of radial elements of this paratype comes close to that of the radial elements of higher cycles of some *P. vitiensis*. Nevertheless, the number of septal cycles, and the relative thickness of septa from different cycles, as well as the size of the dentation of the septa fall within the range of *P. rowleyensis* rather than in that of *P. vitiensis*.

In some genera of lobophylliids (*e.g.* *Lobophyllia, Symphyllia, Parascolymia*), the teeth in different septal cycles differ significantly in shape while in other genera (*e.g.* *Acanthastrea* and *Homophyllia*) such differentiation is not observed (Budd and Stolarski, 2009). Our results confirm that the size and shape of septal teeth of *P. vitiensis* is highly variable within and between septa of the same specimen (Chevalier, 1975;...
Veron and Pichon, 1980; Budd and Stolarski, 2009) (Figs 4A-C, 5C-D, 7). The remarkable variation of these characters in *P. vitienensis*, and their within and between septa variability was also described by other authors (Veron and Pichon, 1980) and led Chevalier (1975) to describe the variety *dentorotundata* (namely, with rounded teeth) for some specimens from New Caledonia (*e.g.* Fig. 2B-C). However, in *P. rowleyensis* the variability in size and shape of septal teeth is much less developed, with septal and costoseptal teeth being of more uniform size and shape than in *P. vitienensis* (*for P. rowleyensis* see Figs 4D-F, 5A-B, 8).

**Molecular phylogeny of *P. rowleyensis* and *P. vitienensis***

Our multi-locus molecular analyses showed that *P. rowleyensis* belongs to the family Lobophylliidae (Fig. 9), as proposed by Dai and Horng (2009) and Budd et al. (2012) based on the macromorphology of the colony and on traditional taxonomy (Veron, 1985, 1992, 2000). Moreover, the species, traditionally ascribed to the monotypic genus *Australomussa*, does not occur in a distinct molecular clade, rather it is nested within the well-supported clade I sensu Arrigoni et al. (2014), which comprises the genera *Lobophyllia*, *Symphyllia*, and *Parascolymia* (Fig. 9).

*Parascolymia rowleyensis* and *P. vitienensis* could not be separated in any single gene tree or the concatenated phylogeny (Figs 9, S2-S4) and the intraspecific and interspecific divergences within and between the two species completely overlap. The lack of genetic variation suggests that these two nominal species could be just one species or that lineage sorting is incomplete because the two species have a recent common ancestor. The former explanation is unlikely because *P. rowleyensis* and *P. vitienensis* differ in several micro-morphological characters (Table 1) and, therefore, it is more likely these two species have not completely diverged although divergence time estimates are not available. An alternative hypothesis is hybridization between the two species, as reported for other genera (Diekmann et al., 2001; van Oppen et al., 2002; Vollmer and Palumbi, 2004; Richards et al., 2008). However, the lack of intra-individual polymorphism in nuclear sequences of both species and the absence of intermediate morphologies challenges this hypothesis.

**Utility of the examined molecular markers***

The three single gene trees gave congruent phylogeny reconstructions (Figs S2-S4), however higher resolution at the species level was achieved by the ITS region (Fig. S4). The best overall BI and ML support was obtained for the concatenated dataset (Fig. 9).

The scleractinian COI gene is usually characterized by low evolution rate and consequently by an overlap of intraspecific and interspecific divergences that do not allow this marker to be used as a barcoding gene in the order Scleractinia (Hellberg, 2006; Shearer and Croftroth, 2008; Huang et al., 2008). The main exception to this general scenario in scleractian corals is *Stylophora pistillata* Esper, 1797, for which Keshavmurthy et al. (2013) detected four deeply divergent lineages corresponding to four particular geographic regions. COI can also be informative when combined or compared in multi-marker analyses (Fukami et al., 2008; Forsman et al., 2009; Huang et al., 2011; Benzioni et al., 2011, 2012a; Gittenberger et al., 2011) (Figs 4, S2). This mitochondrial region does however resolve the majority of the inner nodes, *i.e.* older relationships, within the family Lobophylliidae (this study and Arrigoni et al., 2012, 2014), Fungiidae Dana, 1846 (Gittenberger et al., 2011), and Poritidae Gray, 1842 (Kitano et al., 2014). In our phylogenetic reconstruction based on this mtDNA region, *P. vitienensis* and *P. rowleyensis* are nested within clade I (*sensu* Arrigoni et al., 2014) but they appear to be polyphyletic (Fig. S2). The intra-specific variability of *P. vitienensis* (0.9 ± 0.2%) and *P. rowleyensis* (0.9 ± 0.2%) overlaps the inter-specific distance between the two species (0.9 ± 0.2%) and the last value is comparable to the mean closest congeneric inter-specific distances among Anthozoa (0.71 ± 0.15%) found by Huang et al. (2008).

The nuclear histone H3 gene has been extensively used in phylogenetic studies of arthropods (Colgan et al., 1998; Maxmen et al., 2003), annelids (Novo et al., 2011), and mollusks (Colgan et al., 2000; Pola and Gosliner, 2010) because it is easily amplifiable, highly conserved at the amino acid level, (transiently) highly expressed, and the presence of multiple histone repeats is an uncommon feature (Colgan et al., 1998, 2000; Maxson et al., 1983). It has recently been used in a coral phylogenetic analysis by Huang et al. (2011, 2014b) where it supported all higher-level lineages within the Merulinidae except clade D/E. Our phylogeny reconstruction based on histone H3 resolved all molecular clades within the Lobophylliidae with high node-support values (Fig. S3). These results suggest that histone H3 could be used to evaluate the broad-base phylogeny of other families, in both the Robust and Complex groups, and the phylogenetic relationships among their genera.
The ITS region has been extensively used to resolve species boundaries in scleractinian corals (Diekmann et al., 2001; Forsmann et al., 2009; Benzoni et al., 2010, 2012b, 2014; Flot et al., 2011; Gittenberger et al., 2011; Stefani et al., 2011; Schmidt-Roach et al., 2012; Arrigoni et al., 2012, 2014; Keshavmurthy et al., 2013; Kitano et al., 2013, 2014). Despite the phylogenetic utility of this marker being questioned because of its unique pattern of secondary structure in the genus Acropora Oken, 1815 (van Oppen et al., 2002; Vollmer and Palumbi, 2004; Chen et al., 2004; Wei et al., 2006), it is currently accepted and considered as the most suitable molecular locus to resolve phylogenetic relationships among closely related species. Here, the ITS region resolved the majority of lobophylliid species (Fig. S4), except for species in clade E (Arrigoni et al., 2014). Within clade I (sensu Arrigoni et al., 2014) the majority of species included were monophyletic, with the notable exception of P. rowleyensis and P. vitienensis. Therefore, these results confirmed the usefulness of this marker in phylogenetic studies and we strongly encourage its application for the delimitation of species boundaries in scleractinian corals until new highly variable markers are discovered.

In conclusion, this study demonstrated that comprehensive studies conducted both at molecular and micromorphological levels are and will be essential to evaluate the evolutionary relationships of scleractinian corals and their taxonomy. We strongly believe that different disciplines, such as morphology, molecular systematics, ecology, and reproduction, should be used for taxonomical studies to reach a more complete and comprehensive approach towards the understanding of coral species diversity and biogeography.

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**References**


Dai CF, Hong S. 2009. Scleractinia fauna of Taiwan II. The robust group. Taipei: National Taiwan University. pp. 1-162.


Gardiner JS. 1899. On the solitary corals, collected by Dr. A. Willey Zoological Results based on Material from New Britain, New Guinea, Loyalty Islands and Elsewhere 2: 161-180.


Hoeksema BW. 2014. The “*Fungia patella* group” (Scleractinia, Fungiidae) revisited with a description of the mini mushroom coral *Cycloseris boschmai* sp. n. *Zookeys* 371: 57-84.


Wei NWV, Wallace CC, Dai CF, Pillay KRM, Chen CA. 2006. Analyses of the ribosomal Internal Transcribed Spacers (ITS) and the 5.8S gene indicate that extremely high rDNA heterogeneity is a unique feature in the scleractinian coral genus *Acropora* (Scleractinia; Acroporidae). *Zoological Studies* 45: 404-418.


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**On-line Supplementary Information**

S1. List of the material examined in this study. For each specimen we list code, identification, molecular clade within the Lobophylliidae, sampling locality, collector, and COI, histone H3, and ITS region sequences used for the phylogenetic reconstructions.

S2. Phylogenetic position of *Parascolymia vitiensis* and *P. rowleyensis* (previously *Australomussa*) within the family Lobophylliidae based on partial mitochondrial COI gene. Bayesian topology is shown. Numbers associated with branches indicate Maximum Likelihood bootstrap (>70%) support (left) and Bayesian posterior probabilities (>0.9) (right). Clades within Lobophylliidae are coloured and labelled A to I according to Arrigoni *et al.* (2014).

S3. Phylogenetic position of *Parascolymia vitiensis* and *P. rowleyensis* (previously *Australomussa*) within the family Lobophylliidae based on nuclear histone H3. Bayesian topology is shown. Numbers associated with branches indicate Maximum Likelihood bootstrap (>70%) support (left) and Bayesian posterior probabilities (>0.9) (right). Clades within Lobophylliidae are coloured and labelled A to I according to Arrigoni *et al.* (2014).

S4. Phylogenetic relationships between *Parascolymia vitiensis* and *P. rowleyensis* (previously *Australomussa*) within the family Lobophylliidae based on nuclear ITS region. Bayesian topology is shown. Numbers associated with branches indicate Maximum Likelihood bootstrap (>70%) support (left) and Bayesian posterior probabilities (>0.9) (right). Clades within Lobophylliidae are coloured and labelled A to I according to Arrigoni *et al.* (2014).

S5. *In situ* photos of the specimens of *Parascolymia vitiensis* analyzed in this study: A) 6816, B) 6830; C) IRD HS2955; D) IRD HS2964; E) IRD HS2985, F) IRD HS3255; G) UNIMIB PFB031; H) UNIMIB PFB032; I) UNIMIB PFB033; J) UNIMIB PFB052; K) UNIMIB PFB053; L) UNIMIB PFB055; M) UNIMIB PFB151; N) UNIMIB PFB152.
Appendix

Based on the aforementioned molecular data and morphologic observations discussed above, Australomussa is considered a junior synonym of Parascolymia and A. rowleyensis is hereafter formally moved to this genus.

Family Lobophylliidae Dai and Horng, 2009
Genus Parascolymia Wells, 1964
Type species. Scolymia vitiensis Brüggemann, 1877: 304

Revised diagnosis. Corallum attached, monocentric or polycyclic by intracalicular and extracalicular budding. Corallite can display polymorphism. Corallite integration uni or multiserial. Calice or valley width is large (see Budd et al., 2012). Continuity of costosepta mostly confluent in policentric coralla. Septa of the last cycle free. Septa spacing large. Relative costosepta thickness between Cs1 and Cs2 versus Cs3 unequal or slightly unequal. In polycentric coralla linkage between centres of adjacent corallites within series is lamellar. Columella trabecular and spongy. Endotheca vesicular. Septal tooth base at mid-septum is elliptical in shape and parallel to the direction of the septum. Tooth tips irregular and lobate. Teeth on S1 high and their spacing is wide. The inter-area structure is generally smooth or with palisade. Tooth shape between Cs3 and Cs1 equal or unequal.

Parascolymia vitiensis (Brüggemann, 1877)

Parascolymia rowleyensis (Veron, 1985)
Australomussa rowleyensis Veron, 1985, p.171, figs 23-25; Veron, 2000, p. 80, figs 1-5; Dai and Horng, 2009, p. 69, figs 1-2; Turak and DeVantier, 2011, p. 175.

Parascolymia rowleyensis (Veron, 1985)
Australomussa rowleyensis Veron, 1985 from Phuket Peninsula, western Thailand, dry specimen.

Examined material

Parascolymia vitiensis (Brüggemann, 1877)

Parascolymia rowleyensis (Veron, 1985)
Australia – (Woodside Collection (Kimberley) Expeditions 2009-2012, Coll. Z. Richards): WAM Z65785, Stn. 114/K12, Patricia Is. (14.25.298°S; 125.30.443°E) 12m, 22/10/2012 (K10); WAM Z65789, Stn. 6, Adele Is. (15.26.676°S; 123.10.249°E) 12m, 15/10/2009 (K60);
WAM Z65788, Stn. 9/K12, Adele Is. (15.30.248°S; 123.05.766°E) 12m, 16/10/2009 (K93); WAM Z65787, Stn. 75/K12, Beagle Reef (15.35.217S; 123.53.654°E) 12m, 20/10/2011 (K94); WAM Z65786, Stn. 78/K12, Mavis Reef (15.50.519°S; 123.60.824°E) 12m, 21/10/2012 (K141); Solomon Islands – (Coll. E. Turak) MTQ G57901 Santa Isabel Island, Palunuhukura (07°50.8’S; 158°43.3’E), 2-26m, 16/05/2004; Indonesia – (Coll. B.W. Hoeksema) RMNH Coel. 23309, W Sumatra, off Padang, Gusung Sipakal reef, 02/05/1995; RMNH Coel. 24941, SW Sulawesi, Spermonde Archipelago, Kudingareng Keke reef, 22/05/1996; RMNH Coel. 24178, SW Sulawesi, Spermonde Archipelago, Bone Lola reef, 11m, 05/03/1996.