Comparative Genomics Highlight the Importance of Lineage-Specific Gene Families in Evolutionary Divergence of the Coral Genus, Montipora

Yuki Yoshioka
Graduate School of Frontier Science, The University of Tokyo

Go Suzuki
Fisheries Technology Institute, Japan Fisheries Research and Education Agency

Yuna Zayasu
Marine Genomics Unit, Okinawa Institute of Science and Technology Graduate University

Hiroshi Yamashita
Fisheries Technology Institute, Japan Fisheries Research and Education Agency

Chuya Shinzato (✉ c.shinzato@aori.u-tokyo.ac.jp)
Atmosphere and Ocean Research Institute, The University of Tokyo

Research Article

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Abstract

**Background:** Scleractinian corals of the genus Montipora (Anthozoa, Cnidaria) possess some unusual biological traits, such as vertical transmission of algal symbionts; however, the genetic bases for those traits remain unknown. We performed extensive comparative genomic analyses among members of the family Acroporidae (Montipora, Acropora and Astreopora) to explore genomic novelties to explain unique biological traits of Montipora using improved genome assemblies and gene predictions for M. cactus, M. efflorescens and Astreopora myriophthalma.

**Results:** We obtained genomic data for the three species, of comparable high quality to other published coral genomes. Comparative genomic analyses revealed that the number of gene families restricted to Montipora are significantly higher than those of Acropora and Astreopora, but their functions are largely unknown. The number of gene families specifically expanded in Montipora was much lower than the number specifically expanded in Acropora. In addition, we found that evolutionary rates of the Montipora-specific gene families were significantly higher than other gene families shared with Acropora and/or Astreopora. Of 40 gene families under positive selection (Ka/Ks ratio > 1) in Montipora, 30 were specifically detected in Montipora-specific gene families. Comparative transcriptome analysis of early life stages of Montipora, which possesses maternally inherited symbionts, and Acropora, which lacks them, revealed that most gene families continuously expressed in Montipora, but not expressed in Acropora do not have orthologs in Acropora. Among the 30 Montipora-specific gene families under positive selection, 27 are expressed in early life stages.

**Conclusions:** Lineage-specific gene families were important to establish the genus Montipora, particularly genes expressed throughout early life stages, which under positive selection, gave rise to biological traits unique to Montipora. Our findings highlight evolutionarily acquired genomic bases that may support symbiosis in these stony corals and provide novel insights into mechanisms of coral-algal symbiosis, the physiological foundation of coral reefs.

**Background**

Coral reefs are the most biologically diverse shallow water marine ecosystems [1]. Reef-building corals and endosymbiotic algae of the family Symbiodiniaceae, photosynthetic products of which provide host corals with energy and nutrients, establish mutualistic relationships that are fundamental to coral reefs [2-4]. However, reef-building corals have declined in recent decades due to a variety of anthropogenic stresses, including ocean warming associated with climate change [5-7]. These stresses result in coral bleaching (the breakdown of the symbiosis between corals and their algal endosymbionts [8]), which ultimately leads to loss of habitat for numerous marine species and can precipitate the collapse of entire coral reef ecosystems [9].

The genus *Montipora* (family Acroporidae; Figure 1) is one of the most widespread reef-building corals in the Indo-Pacific [10]. Colony morphology in the genus varies from submassive to laminar, encrusting, and
branching colonies [10, 11]. *Montipora* has some unusual and interesting biological traits among acroporid corals, such as its mode of transmitting algal symbionts and higher stress tolerance. Symbiont transmission maintains symbioses across generations and strongly influences host evolution and adaptation to environments [12-14]. Two fundamental transmission modes predominate in nature (reviewed in [14]): horizontal transmission (symbionts acquired from the environment) and vertical transmission (symbionts acquired maternally). While most coral species (~71%), including *Acropora*, acquire symbionts from the ocean in each generation [15], all known *Montipora* species acquire algal symbionts vertically [16, 17] (Figure 1). Offspring of horizontal transmitters generally associate with a broad range of symbiont types and acquire optimal symbionts from new environments [18, 19]; however, there is no guarantee that optimal symbionts will be available. By contrast, offspring of vertical transmitters inherit symbionts that are suitable for their environments [20], but if they encounter an environment that differs significantly from that of their parents, or if the environment changes too drastically, the inherited symbionts may be disadvantageous. *Montipora* also exhibits low sensitivity to ocean acidification and thermal stressors compared to other coral species [21, 22]. These distinct differences between *Montipora* and its close congener, *Acropora*, may have occurred after their divergence (approx. 125 Mya [23]).

In the family Acroporidae, whole-genome data are becoming more readily available, now including 16 species of *Acropora* [23-26], 3 species of *Montipora* [23, 27, 28] and 1 *Astreopora* species [23], the latter being the basal genus of the Acroporidae [29] (Figure 1). Recently, Shinzato et al. [23] performed a large-scale genomic comparison of acroporids (using genomes of *Acropora*, *Montipora*, and *Astreopora*) and proposed that the evolutionary success of *Acropora* may have occurred by virtue of gene duplications. Although there have been some studies performing genome-wide analysis using *Montipora* genomes [27, 28], the genomic basis for their unique biological traits remains unknown. Exploiting abundant acroporid genomic resources, we performed comparative genomic analyses using improved genomic data of *Montipora* and *Astreopora*. We further identified genes with high evolutionary rates in *Montipora* that may be associated with adaptive evolution, and we specifically attempted to identify genes related to maintenance of maternally inherited symbionts by comparing gene expression during early life stages of *Montipora* and *Acropora*.

**Results**

*Improvement of genome assemblies and gene predictions for Montipora and Astreopora*

Assembly error, including retention of allelic contigs in haploid assemblies, is problematic for downstream analyses, mainly due to redundant genome sequences (alleles from the same genetic locus). We curated scaffold sequences of *M. cactus* and *M. efflorescens* by removing scaffold sequences with high or low coverage and those that may have originated from one of two allelic copies in heterozygous regions. Numbers of scaffold sequences were significantly reduced from the previous version, from 4,925 to 3,521 in *M. cactus* and from 5,162 to 3,589 in *M. efflorescens* (Table 1). For *Astreopora*, possible allelic scaffold sequences were removed from the genome assembly during the previous study [23]. The
previous version of gene models for *M. cactus*, *M. efflorescens*, and *Astreopora* were predicted using AUGUSTUS, based solely on a training set built for *Acropora* or for protein homology with gene models of other corals [23]. Thus, it was highly possible that lineage-specific genes were missed in the previous version. In this study, we performed gene prediction for *M. cactus*, *M. efflorescens*, and *Astreopora myriophthalma* using a combination of *ab initio* and RNA-seq evidence-based prediction. We predicted 29,158 protein-coding genes for *M. cactus*, 29,424 for *M. efflorescens* and 25,406 for *Astreopora myriophthalma* (Table 1). Benchmarking universal single-copy orthologs (BUSCO) completeness scores were 93.3% (of which 0.8% were duplicated) for *M. cactus*, 91.2% (of which 1.4% were duplicated) for *M. efflorescens* and 94.5% (of which 1.3% were duplicated) for *Astreopora myriophthalma*, which were considerably better scores than the previous version (Table 1). In comparison to other *Montipora* gene models, gene models reported by Shumaker et al. [28] may have contained a higher fraction of diploid copies (93.4% complete BUSCO, with 18.3% duplicated; Table1). Completeness of gene models reported by Helmkampf et al. [27] was lower than that reported by Shumaker et al. [28] (64.2%, with 0.5% duplicated; Table1). Thus, the gene models reported by Shumaker et al. [28] contained many duplicates, but those reported by Helmkampf et al. [27] lacked many genes. In contrast, BUSCO completeness scores of *M. cactus*, *M. efflorescens* and *Astreoporamyriophthalma* reported in this study were comparable to published gene models of other coral species, including *A. millepora*, predicted using the NCBI annotation pipeline (Table 1). These improvements to the *Montipora* and *Astreopora* genomes enabled more accurate comparative genomics among acroporids.

**Comparison of gene families within the Acroporidae**

Identifying orthologous relationships between sequences is fundamental for comparative genomic analyses. To obtain orthologous relationships among acroporid genomes, we used three *Acropora* species (*A. digitifera*, *A. millepora*, and *A. tenuis*), for which BUSCO completeness scores are high (Table 1), two *Montipora* species (*M. cactus* and *M. efflorescens*), and *Astreoporamyriophthalma*, representing the basal clade of the Acroporidae [29]. We obtained 12,769 gene families for *Montipora*, 11,007 for *Acropora* and 11,309 for *Astreopora* (Figure 2). We then categorized each gene family into seven groups, (1) common to all three genera (9,690 gene families), (2) common to *Montipora* and *Acropora* (743 gene families), (3) common to *Montipora* and *Astreopora* (665 gene families), (4) common to *Acropora* and *Astreopora* (257 gene families), (5) restricted to *Montipora* (1,670 gene families), (6) restricted to *Astreopora* (696 gene families) and (7) restricted to *Acropora* (316 gene families) (Figure 2). 75.8% (9,690/12,769) of the gene families in *Montipora*, 88% (9,690/11,007) in *Acropora*, and 85.7% (9,690/11,309) in *Astreopora* were shared among all three genera (Figure 2), indicating that a large number (~ 80 - 90%) of gene families are shared throughout the Acroporidae, and these are likely to be the core-gene families the Acroporidae.

The two major clades of reef-building corals possess different metabolic pathways [30]. From the six species, we compared 303 functional modules comprising ten categories in the Kyoto Encyclopedia of Genes and Genomes (KEGG) metabolic pathways and found that metabolic pathways were basically conserved in the three genera (Supplementary Table S1). An enzyme involved in cysteine biosynthesis
(KEGG module ID: M00338) and methionine degradation (KEGG module ID: M00035) was not detected among the six species (Supplementary Table S1), as reported in Shinzato et al. [23, 24]. Although one gene (KEGG entry ID: K04486) involved in the histidine biosynthetic pathway (KEGG module ID: M00026) was detected in acroporid corals used in this study, the remaining genes required to complete the pathway were not detected (Supplementary Table S1), as reported in Ying et al. [30]. Taken together, gene families involved in common features, such as amino acid synthesis, are widely conserved in the three genera.

While we identified 696 lineage-specific gene families in Astreopora and 316 in Acropora, we identified 1,670 gene families restricted to Montipora (2,307 genes in M. cactus and 2,303 in M. efflorescens) (Figure 2). The proportion of lineage-specific gene families in Montipora (13.07%) was significantly larger than those in Acropora (2.87%) and Astreopora (6.15%) (Pairwise proportion test: \( p < 0.05 \)). In addition, although we performed gene annotation with BLAST searches against the SwissProt database (BLASTP, e-value cutoff: 1e-5), the proportion of Montipora-specific gene families with SwissProt annotation was significantly lower than in Acropora and Astreopora (Pairwise proportion test: \( p < 0.05 \) for Montipora versus Acropora, \( p < 0.05 \) for Montipora versus Astreopora, and \( p = 0.59 \) for Acropora versus Astreopora; Figure 2). This indicates that functions of gene families restricted to Montipora are largely unknown.

**Gene expansions in Montipora and comparisons among acroporids**

Gene duplication has contributed to acquisition of new gene functions during evolution [31, 32]. To explore gene families that underwent expansions, we first compared gene numbers of 9,690 gene families common to the three genera and 743 gene families common to Montipora and Acropora (Figure 2). In these two groups, genes in families that underwent gene expansions in either Montipora or Acropora might have been duplicated after Montipora and Acropora diverged from their common ancestor. Three gene families, similar to dimethylsulfoniopropionate (DMSP) lyase (Alma; HOG0000829), Endonuclease-reverse transcriptase (GP1; HOG0000531), and Spondin (Spon1; HOG0001590), and three non-annotated gene families (NA; HOG0000965, HOG0001135, and HOG0001312), were significantly expanded in Acropora (Fisher’s exact test: \( p < 0.05 \); Figure 3a and 3b). Recently, it was reported that DMSP lyase is the most expanded gene family in Acropora [28], and our result is consistent with a previous report, supporting the accuracy of this analysis. We found that three gene families, transient receptor potential protein (TRPC; HOG0002487), collagen alpha-1 (VII) chain (COL7A1; HOG0003259) and non-annotated gene family (NA; HOG0001797) are significantly expanded in Montipora compared with Acropora (Fisher’s exact test: \( p < 0.05 \); Figure 3a and 3b).

Next, we compared gene numbers of 665 gene families common to Montipora and Astreopora (Figure 2), in which gene duplication may have occurred after divergence of Montipora or Astreopora. These genes may have been lost in Acropora. Two gene families (HOG0003949 and HOG0004557) lacking SwissProt annotation were significantly expanded in Astreopora (Fisher’s exact test: \( p < 0.05 \); Figure 3c), whereas one other gene family, tetramericopeptide repeat protein 28 (TTC28; HOG0000387), which is involved in the
cell cycle in humans [33], was significantly expanded in Montipora compared with Astreopora (Fisher’s exact test: $p < 0.05$; Figure 3c).

**Estimation of evolutionary rate in each Montipora gene family group**

The ratio of nonsynonymous (Ka) to synonymous substitutions (Ks) reflects the strength of selective pressure on protein sequences [34]. For example, when Ka is less than Ks ($Ka/Ks < 1$), selection has occurred to eliminate mutations of protein sequences (negative or purifying selection). In contrast, when Ka is larger than Ks ($Ka/Ks > 1$), selection has occurred to mutate the protein sequences (positive selection). In order to evaluate the strength of selective pressure acting on protein sequences in each Montipora gene family, we calculated pairwise Ka/Ks ratios between Montipora single-copy orthologous gene pairs ($M. cactus$ versus $M. efflorescens$) for each of the four groups: 1) gene families common to the three Acroporidae genera, 2) gene families common to Montipora and Acropora, 3) gene families common to Montipora and Astreopora, and 4) gene families restricted to Montipora (Figure 4). When we compared Ka/Ks ratio between groups, gene families restricted to Montipora showed a highest Ka/Ks ratio (Wilcoxon rank sum test: $p < 0.05$; Figure 4), indicating that this gene family group has undergone a relaxation of negative selection, and that functional constraints on this gene family group are relaxed. This could explain why the deduced gene functions of gene families restricted to Montipora are largely unknown.

**Positive selection specific to Montipora**

To identify genes with fast evolutionary rates that may be associated with adaptive evolution in Montipora, we focused on gene families exhibiting $Ka/Ks > 1$. We found evidence of positive selection in 40 gene families (rapidly evolving gene families) (Table 2). Of those, 10 families are common to the three genera or common to Montipora and Acropora, while the remaining 30 families are restricted to Montipora (Table 2), suggesting that these 30 gene families arose specifically in that lineage and likely contribute to biological traits unique to Montipora. Although 28 of the 30 gene families restricted to Montipora were without annotation, their possible subcellular localization ranging from membrane to organelle was predicted by DeepLoc, a deep learning neural networks model (Table 2).

**Gene expression unique to early life stages of Montipora**

Presence of maternally inherited algal symbionts at an early life stage is the most obvious difference between vertical and horizontal transmitters (Figure 1). In order to identify gene families specifically involved in symbiosis in vertical transmitters, we compared the repertoire of expressed genes in early life stages of Montipora with those expressed in Acropora. In this analysis, a gene family was considered expressed if even only one gene in that family was expressed (Transcript per million (TPM) > 1). We confirmed that 11,930 and 10,838 gene families were expressed at early life stages of Montipora and Acropora, respectively (Figure 5a). Of these, 10,051 gene families (84% in Montipora and 93% in Acropora) were common to both at early life stages (Figure 5a), suggesting that these are essential for early development of acroporid corals; thus, we did not focus on these in the present study. We identified
1,879 gene families that were exclusively expressed in Montipora (Figure 5b). Among those, 60% (1,132 gene families) were expressed in planula larvae, metamorphosed larvae and recruit stages (Figure 5b), suggesting that these genes may be related to maintenance of algal symbionts in Montipora. Interestingly, 97% of these gene families ((753 + 344) / 1,132, Figure 5b) that were expressed throughout the three life stages were specific to Montipora or shared by Astreopora (Supplementary Table S2). In contrast, the remaining 3% of gene families ((22 + 13) / 1,132, Figure 5b) have orthologs in Acropora, but were not expressed in Montipora. Nonetheless, they were expressed throughout early life stages of Montipora (Supplementary Table S3). Within gene families containing gene duplications in the Montipora genomes above, two gene families (HOG0001797 and HOG0000387) were exclusively expressed in at least one early life stage in Montipora, and one of them (HOG0000387) was expressed throughout all three early life stages (Supplementary Table S2). Among the identified 30 rapidly evolving gene families restricted to Montipora, we detected gene expression of 90% of these families. Expression of nine families was detected in at least one early life stage of Montipora, and the remaining 18 gene families were continuously expressed throughout all three early life stages (Table 2).

**Discussion**

*Improved genome information for genus Montipora and Astreopora*

BUSCO completeness scores for improved gene models of *M. cactus*, *M. efflorescens*, and Astreoporamyriophthalma were 93.3% (0.8% duplicates), 91.2% (1.4% duplicates), and 94.5% (1.3% duplicates), respectively (Table 1). They are considerably better than those of *M. capitata* (93.4% (18.3% duplicates) from Shumaker et al. [28] and 64.2% (0.5% duplicates) from Helmkampf et al. [27]; Table 1), and were comparable to those of other coral species (Table 1). These numbers indicate that we successfully obtained high-quality gene models from Montipora and Astreopora species. Numbers of genes in *M. cactus* and *M. efflorescens* genomes were not quite as large as that of *M. capitata* reported by Shumaker et al. [28]. Previously, it was reported that *M. capitata* has fewer exons and shorter coding regions per gene than other corals [27, 28]; however, this was not the case with *M. cactus* and *M. efflorescens* (Table 1). Fewer exons and shorter coding regions per gene could be an unusual feature of the *M. capitata* genome or could reflect the quality of the genome assembly. Indeed, the N50 size, one of the indices to evaluate the quality of genome assembly, was larger for both *M. cactus* and *M. efflorescens* genome assemblies than for *M. capitata* (Table 1).

*Possible genomic evolutionary strategy unique to Montipora*

Recent large-scale genome comparisons of acroporid genomes showed that 28 gene families were specifically expanded in Acropora, but none in Montipora [23], but we identified four expanded gene families in Montipora (Figure 3). Although the number of gene families in Montipora is not much different from those of Acropora and Astreopora, the proportion of lineage-specific gene families in Montipora was significantly larger than in Acropora and Astreopora (Figure 2). Montipora does not appear to have duplicated existing gene families, as has Acropora. Lineage-specific gene families contribute larger gene
numbers in Montipora genomes, and emergence of lineage-specific genes may have helped to establish unique biological traits of Montipora corals. In particular, Montipora-specific gene families under positive selection may be major contributors.

Three gene families, similar to Trpc6, TTC28, and COL7A1, and one gene family without annotation were significantly expanded in Montipora compared with Acropora or Astreopora (Figure 3). Known functions of transient receptor potential (TRP) proteins encoded by Trpc are diverse (reviewed in [35]). For example, TRP proteins respond to hypertonicity in yeasts [36, 37], detect and avoid noxious chemicals in nematodes [38], and discriminate warmth, heat, and cold in humans [35]. In each case, TRP proteins mediate sensory transduction in cells [35]. In corals, expression levels of Trp-like genes change when the concentration of CO₂ in seawater changes [39]. They also change diurnally [40, 41] or when exposed to symbiotic algae [42, 43]. The Trpc6-like gene family, specifically expanded in Montipora, may also be involved in sensory transduction during environmental transitions. The TTC28-like gene family has tetratricopeptide repeats (PF12176 and/or PF13424) and caspase HetF associated with Tprs (CHAT) domains (PF12770) (Supplementary Table S4). Canonical TTC28 is composed of tetratricopeptide repeats and CHAT domains (Q96AY4: TTC28_HUMAN; [33]) and genes in the gene family (HOG0000387) are also composed of tetratricopeptide repeats and CHAT domains, indicating that this gene family may have been duplicated from canonical TTC28, which is conserved in all acroporids examined in this study (HOG0016559 in Supplementary Data S1). TTC28 is required for the cell cycle in humans [33]. The expanded TTC28-like gene family may also be involved in cell cycle in Montipora. Collagen is expressed in gastrodermis at a specific developmental stage of cnidarian larvae [44-46] and the expanded collagen-like gene family may function in early development of Montipora.

In this study, we identified 40 genes under positive selection in Montipora (Table 2). Positive selection has often been detected in genes involved in immunity in vertebrates [47]. In corals, genes related to immunity, such as lectin and antimicrobial peptide, are also under positive selection [23, 48, 49]. In the 40 rapidly evolving gene families found in this study, with one exception, no genes appeared homologous to immune-related genes (Table 2). In addition, 28 of 30 rapidly evolving gene families restricted to Montipora have no annotated function (Table 2). Generally, genes with no homology to genes of other lineages are called orphan genes [50]. They may arise principally by two processes: gene duplication or de novo evolution from non-coding regions [50]. If a gene originates by duplication, the protein domains tend to be conserved in the new genes, since a functional protein domain cannot easily be changed by mutations [51], suggesting that the 28 rapidly evolving gene families originated by de novo evolution from non-coding regions. Orphan genes are expected to interact specifically with the environment as a consequence of lineage-specific adaptation [50]. Therefore, orphan genes may contribute to adaptive evolution in Montipora. In particular, 18 rapidly evolving gene families with expression throughout the three early life stages, planula larvae, metamorphosed larvae and recruits, may have important functions in symbiosis during early life stages of Montipora.

Conclusions
In this study, we highlighted possible genomic underpinnings of unusual biological traits of Montipora using high-quality genomic information of Montipora and Astreopora. We found that the driving force behind evolution of Montipora was lineage-specific gene families, rather than gene duplication, as among Acropora corals. The importance of rapidly evolving gene families in Montipora for their unique biological traits was particularly highlighted. Our dataset and findings offer novel insights into mechanisms of coral-algal symbiosis. Although genetic tools for manipulating corals have been established [52, 53], development of more efficient methods to deliver gene-knockdown or -knockout reagents into large numbers of zygotes will facilitate rapid screening for relevant phenotypes of candidate genes. In addition, coral cell lines which have the capacity to incorporate algal symbionts has been developed [54], allowing us to observe ongoing symbiosis at the single cell level. Together, these advances will facilitate a deeper understanding of cellular and molecular mechanisms of coral-algal symbiosis.

Methods

Sample preparation, RNA extraction, and RNA-Seq

Colonies of Montipora cactus, M. efflorescens, and Astreopora myriophthalma were collected in Sekisei Lagoon, Okinawa, Japan in May 2015, and were maintained in aquaria at the Research Center for Subtropical Fisheries, Seikai National Fisheries Research Institute, until spawning. Permits for coral collection were kindly provided by the Okinawa Prefectural Government for research use (Permits #29-74). We isolated total RNA from adult colonies of M. cactus, M. efflorescens, and Astreopora myriophthalma using an RNeasy Plant Mini Kit (QIAGEN). A TruSeq Stranded mRNA Library Kit (Illumina) was used for mRNA sequencing library preparation, and each library was sequenced from 100-bp paired-end libraries using a NovaSeq 6000 (Illumina). For Montipora, we also isolated total RNA from eggs, sperm, planula larvae (1- and 4-days post-fertilization) using TRIzol reagent (Thermo Fisher Scientific) according to the manufacturer’s protocol. KAPA RNA HyperPrep Kit (Kapa Biosystems) and MGIEasy RNA Directional Library Prep Set (MGI) were used for total RNA and mRNA sequencing library preparation, and each library was sequenced on a NobaSeq 6000 in 150-bp paired-end and a DNBSEQ-G400RS (MGI) in 100-bp paired-end mode. This information is summarized in Supplementary Table S5.

Curating scaffold sequences of M. cactus and M. efflorescens and gene prediction

We downloaded scaffold sequences of M. cactus and M. efflorescens from the genome browser of the OIST Marine Genomics Unit (https://marinegenomics.oist.jp). We identified scaffold sequences with high or low coverage or those that may have originated from one of the two allelic copies of heterozygous regions, using Purge Haplotigs v1.1.1 [55] with default option and excluded these from subsequent analyses.

In addition to the above RNA samples, we used publicly available RNA-seq data from NCBI SRA for gene prediction (Supplementary Table S6). Low-quality reads (quality score < 20 and length < 20 bp) and sequence adaptors were trimmed using CUTADAPT v1.18 [56]. A total of 31 and 2 RNA-seq libraries were used for Montipora and Astreopora gene prediction, respectively. Repetitive elements in the scaffolds
were identified de novo with RepeatScout v1.0.6 [57] and RepeatMasker v4.1.0 (http://www.repeatmasker.org). Repetitive elements were filtered out by length (> 50 bp) and occurrence (more than 10 times for Montipora, more than 60 times for Astreopora). Gene prediction was first executed with the BRAKER pipeline v2.1.2 [58], with AUGUSTUS v3.3.3. RNA-seq reads were aligned to each genome sequence with HISAT v2.1.0 [59]. Then, the alignment information was used for BRAKER gene prediction with options “UTR=on”, “softmasking”, and “AUGUSTUS_ab_initio.” To improve gene prediction, we further executed genome-guided transcriptome assembly using StringTie [60] with option “-m 500.” Genome-based transcript structure was predicted with TransDecoder (https://github.com/TransDecoder/TransDecoder/wiki). During read alignment, we used soft-masked repeats for genome-guided transcriptome assembly and hard-masked repeats for BRAKER gene prediction. Finally, genes that were present in genome-guided assembly or ab initio prediction, but absent in predictions from the hint file were added to the prediction from the hint file using GffCompare [61], as summarized in Supplementary Figure S1. To evaluate the completeness of predicted genes, we used BUSCO v5.0 [62] with Metazoa OrthoDB10 dataset (2021-02-24, n=954).

Gene annotation, orthology inference within the Acroporidae

We used publicly available gene models for A. digitifera [23, 24], A. tenuis [23], and A. millepora [25] in addition to Montipora and Astreopora gene models. For A. millepora, we downloaded gene models from NCBI RefSeq (RefSeq assembly accession: GCF_004143615.1). We downloaded gene models of v2.0 for A. digitifera and v1.0 for A. tenuis from the genome browser of the OIST Marine Genomics Unit, respectively. We selected the longest transcript variants from each gene and translated them into protein sequences. All proteomes were annotated with BLASTP [63] (E-value cut off: 1e-5) against the SwissProt database (8 January 2021). Domains in proteomes were annotated using InterProScan v5.31-70.0 [64] with default settings. In addition, putative transposable elements in gene models were identified with TransposonPSI (http://transposonpsi.sourceforge.net/), Dfam scan (release 3.3; [65]), and Pfam keyword (“Reverse transcriptase” and “Integrase”). All proteins were also annotated with KEGG [66] in all eukaryote genes using GenoMaple v2.3.2 [67] with the GHOSTX search engine and the bi-directional best hit method. Module completion ratio (MCR) was calculated in each functional module defined by KEGG, also using GenoMaple v2.3.2. For clustering of orthologous genes (herein gene families) of the Acroporidae, we used OrthoFinder v2.4.0 [68] and Porites australiensis gene models (Shinzato et al., unpublished data) were also included as an outgroup for the Acroporidae. In this study, we used phylogenetic hierarchical orthogroups (HOG) as gene families. Gene families common to the three Acropora species were defined as Acropora gene families. Gene families shared by the two Montipora species were defined as Montipora gene families. Gene families containing transposon-like genes were excluded from subsequent analyses.

Transcriptomic comparisons between Montipora and Acropora

We used RNA-seq data of M. efflorescens (planula larvae), A. tenuis (blastula, gastrula, planula larvae and polyps) and A. digitifera (blastula, gastrula, planula larvae and polyps) (Supplementary Table S7). In
addition, publicly available RNA-seq data of *M. capitata* (planula larvae, metamorphosed larvae, and recruits) were also used in this study (Supplementary Table S7). Low-quality reads (quality score < 20 and length < 20 bp) and sequence adaptors were trimmed using CUTADAPT v1.18. Cleaned RNA-seq reads were mapped to each organism's gene models (For *M. capitata* RNA-seq data, we used *M. eorescens* gene models as a reference) using SALMON v1.0.0 [69]. Expression levels were quantified using SALMON v1.0.0. Genes with TPM > 1 were considered expressed. Then expressed genes were classified into corresponding gene families based on the above gene family inference.

*Estimation of the ratio of nonsynonymous to synonymous substitutions*

Protein sequences of putative single-copy orthologs between *M. cactus* and *M. efflorescens* were aligned pairwise with MAFFT [70]. Aligned nucleotide codon sequences without alignment gaps were retrieved using the PAL2NAL script [71]. Genes with nucleotide alignment lengths longer than 120 bp were used for further analysis. We calculated pairwise nonsynonymous (Ka) and synonymous (Ks) substitution ratios of single-copy genes between *M. cactus* and *M. efflorescens* using KaKs_Calculator 2.0 [72] with option “-MA”. Following Villanueva-Canas et al. [73], we discarded gene families showing Ks < 0.01, as such low Ks values may result in inaccurate Ka/Ks estimates, and gene families showing Ks or Ka > 2 indicating saturation of substitutions. Genes exhibiting Ka/Ks ratios with *p* < 0.05 (Fisher's exact test) were used for further analysis. Subcellular localization of gene families showing Ka/Ks > 1 was predicted using the DeepLoc-1.0 online server [74].

*Statistical analysis*

Pairwise proportion tests were conducted to compare lineage-specific gene families (“number of lineage-specific gene families” / “number of gene families in lineage”) and gene annotation proportions of lineage-specific gene families (“number of genes with annotation” / “number of genes without annotation”). Fisher's exact test was conducted to identify expanded gene families in each group (“number of genes in one gene family in species A” / “number of genes in the rest of the gene family in species A” versus “number of genes in one gene family in species B” / “number of genes in the rest of the gene family in species B”). We considered a *p* < 0.05 as significantly expanded. The Wilcoxon rank sum test was conducted to compare median Ka/Ks values between gene family groups. All statistical tests were performed in R v4.0.3 [75].

**Abbreviations**

BUSCO: Benchmarking universal single-copy orthologs.

KEGG: Kyoto encyclopedia of genes and genomes.

DMSP: Dimethylsulfoniopropionate.

Ka: nonsynonymous substitutions.
Ks: synonymous substitutions.
TPM: Transcript per million.
TRP: Transient receptor potential protein.
TTC: Tetratricopeptide repeat protein.
CHAT: Caspase HetF associated with Tprs.
HOG: Phylogenetic hierarchical orthogroups.
MCR: Module completion ratio.

Declarations
- Ethics approval and consent to participate
  Not applicable.
- Consent for publication
  Not applicable.
- Availability of data and materials
  Raw RNA-sequencing data have been deposited in the DDBJ/EMBL/GenBank databases under accession number DRA011820 (BioProject ID: PRJDB11460). A genome browser for *M. cactus*, *M. efflorescens* and *Astreopora myriophthalma* is available from the Marine Genomics Unit web site (https://marinegenomics.oist.jp/gallery). Sequence IDs of retained scaffolds were prepared as Supplementary Data S2. Gene models in GTF format for *M. cactus*, *M. efflorescens* and *Astreopora myriophthalma* are provided as supplementary Data S3-S5. For *M. capitata*, we downloaded genome assembly and gene models reported from Shumaker et al. [28] (URL: http://cyanophora.rutgers.edu/montipora/) and Helmkampf et al. [27] (Data set DOI: 10.15482/USDA.ADC/1503958).

- Competing interests
  Chuya Shinzato is an editorial board member for *BMC Ecology and Evolution*.

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- Author’s contributions
C.S. and G.S. conceptualized, and C.S. supervised the project. G.S., C.S. and Y.Z. performed coral sampling. H.Y. helped with sampling of *Montipora* larvae. Y.Y. performed molecular biological experiments and bioinformatic analyses. Y.Y. and C.S. wrote main manuscript.

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Tables

Due to technical limitations, Table 1 and Table 2 are only available as a download in the Supplemental Files section.

Figures

Figure 1

Phylogenetic relationships within the Acroporidae and their morphology. (a) Schematic phylogenetic tree representing evolutionary relationships within the Acroporidae. (b and e) Colonies of Montipora efflorescens (b) and M. cactus (e). (c and d) Eggs of M. efflorescens with algal symbionts photographed under visible light (c) and blue light (d). (f and g) A planula larva of M. efflorescens with algal symbionts, photographed under visible light (f) and blue light (g). (h) A colony of Acropora tenuis. (i and j) A planula larva of A. tenuis without algal symbionts photographed under visible light (i) and blue light (j). (k) A
A colony of *Astreopora myriophthalma*. Algal symbionts (brown dots) in eggs and planula larvae of *Montipora* (c and f). Green fluorescence was from fluorescent proteins of *Montipora* and red fluorescence was from chlorophyll in algal symbionts (d and g). Orange and cyan-green fluorescence were from fluorescent proteins of *Acropora* (j).

**Figure 2**

Gene family composition in acroporid genomes and the higher proportion of function-unknown genes in *Montipora*. Left horizontal bars indicate numbers of gene families in each genus. Vertical bars indicate numbers of gene families conserved among genera. Pie charts indicate the generic composition in a given number of gene families (vertical bars). Gene annotation was performed using BLAST searches against the SwissProt database (e-value cutoff: 1e-5), and numbers in pie charts indicate the number of assigned or unassigned gene annotations. Proportions of gene annotation were compared among gene families specific to each lineage and asterisks indicate statistical significance (Pairwise proportion test: p < 0.05). Upset plot was produced using the "UpSetR" package [76].
Figure 3

Gene family expansions in Montipora. (a) Comparison of numbers of genes in Montipora and Acropora in each gene family common to the three genera. (b) Comparison of numbers of genes in Montipora and Acropora in each gene family common to Montipora and Acropora. (c) Comparison of numbers of genes in Montipora and Astreopora in each gene family common to Montipora and Astreopora. The diagonal solid line indicates 1:1 numbers of genes in orthologous families. Possible gene names and gene family
IDs are shown for significantly expanded gene families (Fisher’s exact test: p < 0.05). Scatter plot was produced using the “ggplot2” package [77].

![Figure 4](image)

**Figure 4**

Relaxed negative selection in Montipora-specific gene families. The Y-axis represents the distribution of the ratio of nonsynonymous (Ka) to synonymous amino acid substitutions (Ks). Orthologous gene pairs in two Montipora species (M. cactus and M. efflorescens) are used for calculation of pairwise Ka/Ks rate. Ka/Ks ratios were compared among gene families and significant differences were observed in all pairwise combinations (Wilcoxon rank sum test: p < 0.05). A raincloud plot was produced using the “raincloudplots” package [78].
Figure 5

Expression patterns of gene families during early life stages of Montipora and Acropora. (a) Numbers of gene families that are commonly or exclusively expressed in early life stages of Montipora and Acropora. Numbers of gene families that are exclusively expressed in each genus are shown in upper parentheses after genera names. The number of gene families that are commonly expressed in both Montipora and Acropora early life stages are shown in an orange box. Gene families that are exclusively expressed in
early life stages of Montipora were further classified according to whether they are expressed in one of the stages (green), or throughout all stages (red). (b) Gene families expressed in early Montipora life stages. For Montipora, RNA-seq data from planula larvae, metamorphosed larvae and recruits were used. For Acropora, RNA-seq data from blastulae, gastrulae, planula larvae and polyps were used. SRA accession numbers for the RNA-seq data are provided in Supplementary Table S7.

Supplementary Files

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