Not out of the Mediterranean: Atlantic populations of the gorgonian Paramuricea clavata are a separate sister species under further lineage diversification

Márcio Coelho\textsuperscript{1}, Gareth Pearson\textsuperscript{2}, Joana Boavida\textsuperscript{3}, Diogo Paulo\textsuperscript{4}, Didier Aurelle\textsuperscript{5}, Sophie Arnaud-Haond\textsuperscript{6}, Daniel Gómez-Gras\textsuperscript{7}, Nathaniel Bensoussan\textsuperscript{5}, Paula López-Sendino\textsuperscript{7}, Carlo Cerrano\textsuperscript{8}, Silvija Kipson\textsuperscript{9}, Tatjana Bakran-Petricioli\textsuperscript{9}, Eliana Ferretti\textsuperscript{10}, Cristina Linares\textsuperscript{11}, Joaquim Garrabou\textsuperscript{12}, Ester Serrão\textsuperscript{4}, and Jean-Baptiste Ledoux\textsuperscript{13}

\textsuperscript{1}Algarve Centre of Marine Sciences
\textsuperscript{2}CCMAR-CIMAR
\textsuperscript{3}Aix Marseille Univ, Université de Toulon, CNRS, IRD, MIO, Centre for Marine Sciences (CCMAR), University of Algarve
\textsuperscript{4}Centre for Marine Sciences (CCMAR), University of Algarve
\textsuperscript{5}Aix Marseille Univ, Université de Toulon, CNRS, IRD, MIO
\textsuperscript{6}Ifremer
\textsuperscript{7}Institut de Ciències del Mar (CSIC)
\textsuperscript{8}Consorzio Nazionale Interuniversitario per le Scienze del Mare (CoNISMa)
\textsuperscript{9}University of Zagreb Department of Biology
\textsuperscript{10}Studio Associato GAIA s.n.c.
\textsuperscript{11}Facultat de Biologia, Universitat de Barcelona
\textsuperscript{12}Institut de Ciències del Mar CSIC
\textsuperscript{13}ICM

June 17, 2022

Abstract

The accurate delimitation of species boundaries in non-bilaterian marine taxa is notoriously difficult, with consequences for many studies in ecology and evolution. Anthozoans are a diverse group of key structural organisms worldwide, but the lack of reliable morphological characters and informative genetic markers hampers our ability to understand species diversification. We investigated population differentiation and species limits in Atlantic (Iberian Peninsula) and Mediterranean lineages of the octocoral genus Paramuricea previously identified as P. clavata. We used a diverse set of molecular markers (microsatellites, RNA-seq derived single-copy orthologues [SCO] and mt-mutS [mitochondria]) at 49 locations. Clear segregation of Atlantic and Mediterranean lineages was found with all markers. Species-tree estimations based on SCO strongly supported these two clades as distinct, recently diverged sister species with incomplete lineage sorting, P. cf. grayi and P. clavata, respectively. Furthermore, a second putative (or ongoing) speciation event was detected in the Atlantic between two P. cf. grayi colour morphotypes (yellow and purple) using SCO and supported by microsatellites. While segregating P. cf. grayi lineages showed considerable geographic structure, dominating circalittoral communities in southern (yellow) and western (purple) Portugal, their occurrence in sympatry at some localities suggests a degree of reproductive isolation. Overall, our results show that previous molecular and morphological studies have underestimated species diversity in Paramuricea occurring in the Iberian Peninsula, which has important implications for conservation planning. Finally, our findings validate the usefulness of phylotranscriptomics for resolving evolutionary relationships in octocorals.
Introduction

The species is the fundamental unit of analysis in ecology and evolution, from field and behavioural ecology to population genetics, phylogeography and systematics. Thus, species delimitation sets the tone for research questions in a myriad of fields, with critical implications for biodiversity assessment and conservation planning. However, identifying species boundaries is challenging in many taxa for several reasons, including phenotypic plasticity; lack of diagnostic morphological characters; homoplasy; low-resolution of DNA barcoding markers; incomplete lineage sorting (ILS) in recently diverged taxa; and introgressive hybridization. The latter phenomenon is generally the result of complex biogeographic histories in which populations that diverged in allopatry come into secondary contact. Identifying barriers to gene flow and understanding how genetic differences accumulate over time is an inherently challenging task, particularly for marine systems in which these barriers are less obvious than for their terrestrial counterparts.

The Atlantic-Mediterranean transition provides an interesting geographic setting to study population differentiation and speciation in the marine realm. This narrow region has a complex biogeographical history, involving dramatic geological events that affected the entire Mediterranean Sea, including the Messinian Salinity Crisis and subsequent Zanclean flood, as well as sea level fluctuations during glacial cycles. These, in turn, have contributed to multiple episodes of species extinction-recolonization or allopatry-secondary contact between the Atlantic and Mediterranean basins, which have shaped contemporary patterns of genetic differentiation across this range. Within the Mediterranean, several oceanographic discontinuities exist, among which is the Almeria-Oran Front (AOF), a large-scale density front at the eastern edge of the East Alboran gyre that extends southwards from Almeria (SE Spain) to Oran (Algeria). The AOF constitutes a barrier to gene flow in many species, often in complex phylogeographical patterns, particularly in taxa with restricted dispersal such as seagrasses, sponges, and corals. Indeed, genetic discontinuities associated with the AOF are even reported for species with high dispersal potential such as the European sea bass (Dicentrarchus labrax) (Duranton et al., 2018; Lemaire, Versini, & Bonhomme, 2005; Naciri, Lemaire, Borsa, & Bonhomme, 1999; Tine et al., 2014).

Branching octocorals of the genus Paramuricea (Octocorallia: Plexauridae) are important constituents of many shallow and deep-sea communities of benthic suspension feeders (e.g., corals, sponges and bivalves) in the Atlantic Ocean and Mediterranean Sea, often forming extensive, dense assemblages supporting high levels of biodiversity. Currently, seven valid species are known to occur along the NE Atlantic and Mediterranean, with a further two putatively new species recently discovered after re-examination of natural history collections. Two of these, Paramuricea clavata and Paramuricea macrospina, were historically regarded as endemic to the Mediterranean Sea despite several recent studies extending the distribution ranges of P. clavata to adjacent Atlantic coastlines and of P. macrospinapotentially to the Cape Verde archipelago.

The red gorgonian Paramuricea clavata is a major structuring species of Mediterranean coraline assemblages, generally restricted to low-light, reaching densities well above 20 colonies m\(^{-2}\). The genetic structure of P. clavata in the Mediterranean shows sharp regional genetic breaks, strong differentiation at 10s to 100s of meters and significant isolation by distance (IBD) both regional and local scales. Spatial genetic structure in P. clavata is congruent with oceanographic barriers to gene flow seasonal cyclonic gyres along the Adriatic, and with the limited dispersal ability inherent to a reproductive strategy of larval brooding on maternal colonies (as observed in other brooding corals). Atlantic populations identified as P. clavata along the Portuguese coast are genetically differentiated from Mediterranean ones and also differentiated between western and southern Portugal. Interestingly, this differentiation between western and southern populations in Portugal appears to coincide with colour morphotypes that dominate circalittoral coral gardens in each region; with a yellow morphotype prevalent in the Algarve (southern Portugal) and a purple morphotype dominating communities in western Portugal. Although both morphotypes can occur in sympathy, our field surveys indicate that the purple morphotype is extremely rare in the south and vice-versa (J. Boavida and M. Coelho, personal observations). Despite these findings, the existence and exact location of the Atlantic-Mediterranean break, and the level of divergence across these regions (i.e., whether it represents within-species population structure or speciation processes) remains unknown in P. clavata due to incomplete
geographic sampling and low marker resolution. Indeed, standard DNA barcoding approaches to species delimitation are poorly suited to resolving closely related or recently diverged octocorals due to slow rates of mitochondrial gene evolution. These issues obscure the relationship between Mediterranean and Atlantic populations. For instance, Mediterranean populations of the threatened precious red coral *Corallium rubrum* typically exhibit strong genetic differentiation at small spatial scales. However, the Atlantic-Mediterranean genetic break associated with the AOF is apparently lacking, presumably due to the extended influence of Atlantic populations in the Mediterranean gene pool, such as those recently rediscovered in SW Portugal.

In addition to limited geographic sampling and marker resolution, the evolutionary relationships between *P. clavata* populations occurring across the Atlantic-Mediterranean transition have also been obscured by the potential occurrence in sympathy of congenic species with intergrading morphologies that are difficult to discriminate. For instance, the southern coast of Portugal is part of the distribution range of *Paramuricea grayi*, a species of deep littoral habitats historically described from the Lusitanian-Mauritanian East Atlantic to the Gulf of Guinea, but which has subsequently been described from Galicia and the Bay of Biscay (northern Iberian Peninsula), among other locations in the NW Atlantic. Importantly, recent phylogenetic reconstructions based on a mitochondrial marker and RAD-sequencing (RAD-seq) loci have shown that *P. grayi* is a sister species of *P. clavata*. The divergence between these two species could be either the result of a vicariance event in an Atlantic-Mediterranean ancestor; or of divergence within the Mediterranean in which *P. clavata* diversified in mesophotic habitats from an ancestor with a broad depth range. Thus, it is possible that the Iberian Peninsula and adjacent areas form a contact zone between two recently diverged sister species (*P. clavata* in the Mediterranean and *P. grayi* in the NE Atlantic). Alternatively, the Atlantic populations of *P. clavata* reported from Portugal could be misidentified *P. grayi*. This highlights the need for further study of *Paramuricea* (and octocorals in general) along the Atlantic-Mediterranean transition to better understand the evolutionary forces governing population differentiation and speciation in this genus across these important biogeographic regions. The delimitation of spatially significant and unique genetic units within a species is also necessary for emerging conservation policies aiming to preserve intraspecific genetic diversity and the associated adaptive potential. *Paramuricea clavata* needs such efforts given its key structuring role in ecosystem functioning across the Mediterranean, particularly with the growing recognition of its vulnerability to human disturbances and environmental changes. Resolving the nature of the divergence (i.e., population- or species-level) across the Atlantic and Mediterranean is a necessary first step.

The aim of this study was to test genetic boundaries and species affinities within the genus *Paramuricea* across the Atlantic-Mediterranean transition zone and assess the genetic diversity of the genus in these biogeographic regions. For this aim, the geographic scope includes the western and southern coasts of Portugal (Northeast Atlantic) and a large part of the species distribution in the Mediterranean Sea, including the Adriatic and Aegean seas. We combined evidence from three types of molecular markers: 1) single-copy orthologs (obtained from RNA-seq data) for species delimitation and screening for cryptic species diversity; 2) microsatellite loci to analyse spatial genetic diversity and structure both across and within the Atlantic and Mediterranean regions; and 3) the mitochondrial gene *mt-muts* (assembled from RNA-seq read data) to extend taxonomic coverage of our phylogenetic analysis for the genus. We present evidence showing that previous assignments of Atlantic populations to *P. clavata* are incongruent with genomic data, and the taxa present along those coasts should instead be regarded as the morphospecies *P. cf. grayi*.

**Material and Methods**

**Phylogenomic inference and species delimitation using RNA-seq data**

**Sample collection**

Gorgonian samples used for the phylogenomic analyses were subsampled from colonies used by Gómez-Gras et al. (*in review*). Our treatment focused on four populations originally studied by these authors: Balun (Kornati, Croatia), Altare (Portofino, Italy) and La Vaca (Medes, Catalonia) for *P. clavata*; and Baleeira (Sagres, Portugal) for *P. cf. grayi* (Figure 1 and Table S1). Tissue from each of 6-8 individuals per population...
was sampled for RNA-seq analysis (Table S1). The samples were preserved in RNA later and stored at -80 ºC until further processing. In addition, four samples of *P. cf. grayi* were collected from two locations in southern (Tavira) and western (P39, Cape Espichel) Portugal. This included two specimens of the yellow colour morph (1 from Tavira and 1 from P39) and two of the purple colour morph from P39 (Table S1). These samples were flash frozen with liquid nitrogen upon collection and stored at -80 ºC until further processing.

**RNA extraction and sequencing**

Total RNA was extracted from a 0.5-1 cm long branchlet piece. We coupled the TRIzol Plus RNA Purification Kit (Invitrogen) or the NZYol reagent (Nzytech) for tissue homogenization with the RNeasy Mini Kit (Qiagen) for RNA purification following the manufacturer’s instructions. Residual DNA was removed using TurboDNase (Invitrogen) or DNase I (Qiagen). The quality, purity and yield of the RNA was assessed using gel electrophoresis and a Qubit Fluorometer and/or Nanodrop spectrophotometer. RNA-seq library preparation and sequencing was outsourced to BGI Tech Solutions in Hong Kong (100 bp paired-end DNBseq). A total of 34 individuals were processed for RNA-seq and deposited on NCBI’s Sequence Read Archive (BioProject ID: PRJNA847883).

**Sequence quality control and filtering**

Raw reads were pre-processed by BGI to trim adaptor sequences and low-quality reads (< Q20). The quality of filtered sequence data was then evaluated with two complementary quality control tools, FastQC v0.11.6 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and FastqPuri. Because these analyses detected the presence of overrepresented sequences corresponding to ribosomal RNAs (rRNAs), contaminant rRNA reads were filtered with FastqPuri using the bloom filter approach implemented in the function trimFilterPE. The input bloom filter was constructed from nuclear rRNA sequences downloaded from NCBI GenBank for the top octocoral blastn hit of the contaminants, which corresponded to *Dendronephthya gigan
tea*.

**De novo transcriptome assembly and completeness**

Reference transcriptomes were assembled *de novo* with rnaSPAdes using default parameters. In total, we performed four independent transcriptome assemblies using the sequence data of all samples from each population, one for samples from Baleeira (*P. cf. grayi*) and three for samples from Balun, Altare or La Vaca (*P. clavata*). The assemblies were subjected to multiple steps of quality control and curation. First, the quality of each assembly was evaluated with TransRate v1.0.1 using both sequence-based and read mapping (onto the assembled contigs) metrics. The read-metrics mode measures the quality of the assembled contigs based on evidence from the reads used to generate the assembly, allowing the filtration of ‘bad’ (i.e., poorly assembled) contigs from the assembly. Second, using the ‘good’ contigs obtained with TransRate, we identified putative open reading frames (ORFs) with FragGeneScan and clustered the ORFs at 97% nucleotide sequence identity using VSEARCH. Finally, the contigs of the clustered assemblies were aligned against NCBI’s nonredundant database for Anthozoa (Taxonomy ID: 6101) using DIAMOND in BLASTX mode. Contigs with a top query hit were retained in the final transcriptome assemblies, with the remainder discarded as potential contaminants (e.g., bacteria).

The completeness of the final, curated transcriptome assemblies was evaluated with BUSCO v2/v3 by screening the recovery of single-copy orthologues (queried as predicted proteins) that are expected to be present across higher taxonomic groups. The analysis was performed against the Metazoa reference database (OrthoDB 9) using the online platform gVolante v1.2.1 available at https://gvolante.riken.jp/. Given that the transcriptome assembly for *P. cf. grayi* had the highest TransRate and BUSCO scores (see Results), all subsequent analyses were conducted using this assembly as the reference.

**Phylogenetic analysis**

The phylogenetic relationships among the samples were explored using 261 complete and unduplicated single-copy orthologues ([?] 180 bp) identified from BUSCO (N = 106) and from a reciprocal BLASTX search between the transcriptome assembly of *P. cf. grayi* and that of the octocoral *Eunicella verrucosa* (Coelho
et al. unpublished data) using DIAMOND (N = 155). For the reciprocal blast we retained contigs with up to five query hits, subsequently selecting only those with a single sequence hit in both results tables (and thus putative single-copy orthologues). Transcript .fasta sequences for orthologues were extracted for each sample from sorted .bam files after phasing (samtools phase) and variant calling (bcftools mpileup). Allele sequences were obtained from the resulting .vcf files with vcfutils.pl and seqkit fq2fa, aligned with mafft and trimmed with Gblocks in TranslatorX using a custom python script (https://github.com/cymon). The final set of loci used contained no gaps or missing data. The input .bam files were generated by mapping reads onto the reference transcriptome using the RSEM (v1.2.31) wrapper script and Bowtie2. Locus alignments were assigned to “pseudoalleles” and analysed by maximum likelihood in IQ-TREE 2. A partitioned analysis was performed by fitting a separate evolutionary model of sequence evolution for each locus using ModelFinder and based on BIC score. Gene and site concordance factor (CF) analysis, which estimates the percentage of gene trees (gCF) or sites (sCF) which agree with the consensus species tree, was performed following and the recommendations at http://www.iqtree.org/doc/.

Species delimitation on the same multi-individual dataset as used for the species tree was performed using Species bOundry Delimitation using Astral method based on quartet tree analysis with ASTRAL-III . The input ML gene trees from IQ-TREE 2 were used, and a species tree was estimated with ASTRAL-III. SODA uses the mapping of quartet tree constellations onto an ASTRAL species tree with all individuals to assess if/where coalescence of the tree topology is completely random. Where the coalescence is not random, species boundaries are defined based on the quartet frequencies .

In addition to SODA species delimitation, we reconstructed the species tree under a ML polymorphism-aware model . PoMo is a statistically consistent method for species tree estimation that builds on nucleotide substitution models to incorporate population-level processes from allele frequency information . PoMo was run in IQ-TREE 2 under the HKY (Hasegawa-Kishino-Yano) substitution model, using individual allele count data from the 261 single-copy orthologue alignments. For the analysis, individual counts were assigned to a total of 8 populations corresponding to sampling location and colour morph. Branch support values were estimated with 1000 ultrafast bootstrap replicates. The species Paramuricea biscaya was used as an outgroup in all the analyses. We downloaded RNA-seq data for 4 samples from NCBI’s SRA (Accessions: SRX4389727, SRX4389728, SRX4389729 and SRX4389730) . Sequence data were filtered and mapped as described above.

Finally, to extend the taxonomic coverage of the analysis to other Paramuricea spp., we conducted a ML tree reconstruction in IQ-TREE 2 with the mitochondrial gene mt-mutS . Sequence data were retrieved from NCBI Genbank and from .bam files generated by mapping the RNA-seq reads onto the reference mitogenome of P. clavata using RSEM (Genbank Accession: NC_034749) for a subset of the samples (for additional information see Fig. S1). New mt-mutS sequences were deposited on NCBI’s GenBank (Accession numbers: provided upon manuscript acceptance).

Population differentiation based on microsatellite loci

Sample collection

Sampling in the Atlantic took place between 2010 and 2013 in Portugal. The occurrence of P. cf. grayi (heretoforeParamuricea aff. clavata ) along the southern coast of Portugal was known from deep divers’ records, from shallow hard bottoms down to 90 m depth . We extended the search along the Portuguese west coast between 30-70 m depth with the support of volunteer divers (through the citizen participation project Deep Reefs). Nine locations were sampled: Berlengas archipelago (2 sites 20 km apart), Cape Espichel (2 sites 400 m apart), Sines (1 site), Sagres (2 sites 300 m apart), Lagos (1 site), and Tavira (1 site) (Table S1 and Figure 1). At each site samples were collected haphazardly taking well-separated colonies (i.e. individuals) of the same height (approx. 30 cm height, adult colonies) to avoid sampling the same colony multiple times and to account for the very low rate of asexual reproduction observed in P. clavata/P. cf. grayi . At each site, sample size varied from 12 (BAL) to 84 (CAT) individuals according to the species’ density and decompression time constraints in deeper dives (Table S1). Samples consisted of 10 cm apical
branches that were preserved in 96% ethanol at room temperature until DNA extraction. Two additional locations in the Mediterranean were sampled (VAC and PZU; Table S1) and combined with 35 existing sampling locations in the Mediterranean Sea described in Mokhtar-Jamai et al. (2011). Overall, the dataset covered 46 locations, 9 from the Atlantic and 37 from the Mediterranean, and included 1,597 individuals (448 newly sampled) (Figure 1 and Table S1).

**DNA extraction, amplification and genotyping**

Genomic DNA of the Atlantic samples was extracted with a cetyltrimethylammonium bromide protocol, with purification by standard chloroform:isoamyl alcohol (24:1) followed by DNA precipitation. All samples (N=384 colonies) were genotyped at six microsatellite loci: Parcla_09, Parcla_10, Parcla_12, Parcla_14, Parcla_17 and Par_d. The dataset from and the additional Mediterranean sites (N=1243 samples) were extracted using a salting out procedure. All samples were genotyped as in Mokhtar-Jamai et al. (2011) with minor modifications of the PCR protocol (for more details see Note S1). After quality filtering the microsatellite dataset kept for downstream analyses included 1,223 samples from 46 sites (filters: 23% of samples excluded due to missing data, sites with <5 samples and identical multi-locus genotypes).

**Analysis of genetic diversity among lineages**

Indices of genetic diversity were compared among the three lineages identified with SODA, *P. clavata* from the Mediterranean and two segregating lineages of *P. cf. grayi* coincident with colour morphotype (yellow or purple) within the Atlantic (see Results). We compared the observed heterozygosity (*H_o*), gene diversity (*H_e*), *F_{IS}*, allelic richness (*Ar_g*) and private allelic richness (*Ap_g*). The *H_o*, *H_e* and *F_{IS}*, were computed for each lineage with GENETIX 4.05 (Belkhir et al., 2004), whereas *Ar_g* and *Ap_g* were estimated using the rarefaction method implemented in ADZE. Statistical support for differences in *Ar_g* and *Ap_g* were tested with a Kruskal-Wallis rank sum test followed by pairwise Wilcoxon rank sum tests in R. Further details about these procedures can be found in Note S1.

**Genetic structure**

A Bayesian clustering analyses was conducted with STRUCTURE 2.3.4 to evaluate the number of genetic clusters (K) from the individual genotypes without assumptions concerning population boundaries. Because of the unbalanced sampling, we considered the admixture model with uncorrelated allele frequencies, a separate ALPHA for each population and an initial ALPHA value = 0.022 (1/46 the number of sampling locations). Null homozygous genotypes were considered as missing data. Ten independent runs were performed for each K ranging between 1 and 30, with a burn-in period of 1,000,000 followed by 250,000 iterations and using the StrAuto pipeline for parallelization. The K value corresponding to the “upper most hierarchical level of structure” was determined using the likelihood of observing the data, (lnP(D)). CLUMPP 1.1 and DISTRUCT 1.1 were used for graphical output.

A discriminant analysis of principal components was conducted in ADEGENET. Data were transformed into principal components and discriminant analyses were used to maximize variation among-groups while minimizing within group variation. Population locations were used as a group prior. Based on the a-score method, the number of principal components was set to 80, while we retained 2 discriminant functions. Hierarchical genetic subdivision (i.e., among lineages, among population and among individuals) was analyzed with an analysis of molecular variance based on the number of different alleles (infinite allele model) as implemented in GenAlex 6.51 and using 999 permutations. Total and pairwise *G_{ST}* s and *D_{EST}* s among lineages and populations were estimated in GenoDive. These two measures provide complementary information about genetic structure. Indeed, the *G_{ST}* can be considered as an estimate of the nearness to fixation while *D_{EST}* provides information about the level of allelic differentiation. Significance was tested with a permutation procedure (n=1000) for the two measures.

The evolutionary relationships among populations were characterized using the chord distance. Allele frequencies were bootstrapped 100 times in SEQBOOT in PHYLIP 3.6 (Felsenstein, 2005). The consensus
Results

Phylogenomic inference and species delimitation using RNA-seq data

Transcriptome sequencing and quality of the assemblies

RNA-sequencing generated a global dataset of 143 Gb high-quality paired-end reads (28.04 ± 0.91 M reads per sample [mean ± SE]), 38 Gb and 105 Gb for P. cf. grayi and P. clavata, respectively (Table S2). Quality assessment of the four de novo transcriptome assemblies obtained for each population showed that those built for P. cf. grayi (Baleeira, Portugal) and the Italian population of P. clavata (Altare) were superior to those obtained for the other two P. clavata populations (La Vaca and Balun) (Table S3). Transrate quality metrics for the two top assemblies were generally similar (average Transrate contig score = 0.36 for P. cf. grayi and 0.33 for P. clavata) (Table S3). BUSCO analysis of metazoan single-copy orthologues indicated that the curated assemblies (i.e., following ORF clustering and filtering of non-Anthozoa sequences) from both the Baleeira and Altare populations were very complete, with 98% and 97.4% of core proteins detected, respectively (Table S3). After curation, a total of 73,871 contigs were assembled in P. cf. grayi (Baleeira population), with an N50 of 1,452 bp (Table 1).

Phylogenetic analysis

Our partitioned ML analysis of 261 single-copy orthologues clearly separated the Atlantic (southern-central Portugal) P. cf. grayi and the Mediterranean P. clavata samples with high (Atlantic clade, 98%) or full (Mediterranean) support (Figure 2). Within the Atlantic samples, the yellow morphotype of P. cf. grayi from 3 distinct locations (including P39 off Cape Espichel where both colour morphs occur in sympathy) formed a monophyletic group to the exclusion of the purple colour morph (Figures 1 and 2). Within the Mediterranean, individuals from Croatia (Adriatic Sea) also formed a fully-supported clade with full support and distinct from the Spanish and Italian samples (Figure 2).

Despite the high bootstrap support for the Atlantic and Mediterranean clades, concordance analysis of the branches supporting these clades revealed very low gCFs of only 1.15% (Atlantic) and 2.3% (Mediterranean). Concordance factors for sites (sCF) were also low to moderate (45.3% and 58.3%), which together with the short branch lengths supporting these clades, suggests recent divergence with incomplete lineage sorting (ILS). We note however, that the branch supporting the yellow morphotype clade within the Atlantic samples was somewhat longer than those supporting the Atlantic and Mediterranean clades, had full bootstrap support, and a fairly robust sCF (62.8%) (Figure 2).

Species delimitation

Species delimitation based on the ASTRAL-III species tree and performed with the SODA software supported the clades formed by samples of P. cf. grayi and P. clavata as separate species (Figure 2). The same split was also evident with the barcode marker mt-mutS for which our analyses recovered two distinct haplotypes, one including all P. clavata samples from the Mediterranean (96% bootstrap support) and another including the samples of P. cf. grayi sequenced here together with P. grayi from the northern Iberian Peninsula (100% bootstrap support; Figure S1). Furthermore, within the Atlantic, the purple and yellow morphotypes of P. cf. grayi were all recovered as two distinct species in the SODA analysis (with the caveat that our dataset only included 2 purple individuals, although they were sympatric with one individual of the yellow morphotype P39). These results were congruent with the PoMo ML species tree, which supported the Mediterranean (P. clavata) and Atlantic (yellow and purple morphotypes of P. cf. grayi) populations as 3 distinct entities with full bootstrap support (Figure 3). Given that our sampling included just a few individuals of the purple morphotype, we conservatively assign both colour morphs of the Atlantic clade to P. cf. grayi, a species first described (as Acanthogorgia grayi) from Madeira, Portugal (Johnson, 1861), but which has also been documented to occur in southern Portugal.
**Population variability**

The observed ($H_0$) and expected ($H_e$) heterozygosity ranged between 0.44 (TAV) and 0.83 (PZO) (mean \(\pm\) SD= 0.66 \(\pm\) 0.08) and between 0.5 (BAL) and 0.8 (PHL) (mean \(\pm\) SD= 0.71 \(\pm\) 0.07), respectively. Multiple FIS were between 0 (e.g., VAC, LAG, BAL) and 0.3 (TAV) (mean \(\pm\) SD= 0.07 \(\pm\) 0.08). Significant heterozygote deficiencies, after FDR correction, were found in 15 of 46 populations (28%), located both in the Atlantic (e.g., TAV, SIN) and in the Mediterranean (e.g., CAS, RIO). Regarding rarefied allelic ($A_{R(18)}$) and private allelic richness ($A_{P(18)}$), we found highly contrasted values among populations. $A_{R(18)}$ ranged between 3.5 (BAL) and 7.92 (PHL) (mean \(\pm\) SD= 6.51 \(\pm\) 1.1), while $A_{P(18)}$ was null for 14 populations (all belonging to the Mediterranean, e.g., CPS, CLB, MES) with a maximum of 1.52 for TAV (mean \(\pm\) SD= 0.17 \(\pm\) 0.3). Seven of the highest $A_{P(18)}$ values were observed for Atlantic populations (LAG, BAL, SEG, SIN, RAB; Table S4). Significant differences among the three lineages were observed for all the genetic diversity parameters ($H_0$, $H_e$, $A_{R(18)}$, $A_{P(18)}$) with the exception of $F_{IS}$. Pairwise Wilcoxon rank tests showed that significant differences were mostly driven by variation between Mediterranean and the Atlantic populations, which included the two segregating lineages identified (see Tables S4-S5 for details). The highest $H_0$, $H_e$ and $A_r$ values were observed in *P. clavata*, whereas the yellow lineage of *P. cf. grayi* harbours the highest $A_p$.

**Population structure**

The pattern of population structuring was largely congruent with the phylogenetic analyses. The lnP(D) parameter increased until $K = 4$, plateaued until $K = 9$ and decreased for higher $K$ values. Considering our focus on the genetic structure among Atlantic and Mediterranean populations, we describe only the results for $K = 2$ and $3$. At $K = 4$, structuring among Mediterranean populations became apparent, which has been previously characterized. For $K = 2$, individuals from the Atlantic (purple and yellow lineages; mean membership coefficient = 0.97) were separated from those from the Mediterranean (mean membership coefficient = 0.96) (Figure 4a), with a signal of admixture between the purple lineage of *P. cf. grayi* and the Mediterranean populations of *P. clavata* close to the Strait of Gibraltar (MIR, TYL and MTH), and extending to southern Spain (CPS) and the Balearic Islands (CAB). The three genetic clusters ($K = 3$) were concordant with the areas corresponding to western (*P. cf. grayi* purple lineage; mean membership coefficient = 0.93) and southern (*P. cf. grayi* yellow lineage; mean membership coefficient = 0.97) Portugal and the Mediterranean (*P. clavata*; mean membership coefficient = 0.92) (Figure 4a).

The DAPC analysis corroborates the results obtained with STRUCTURE. The first two axes represented relatively high levels of the total variation, 40.4 and 10.7%, respectively. The first axis separated *P. clavata* populations from the Mediterranean and *P. cf. grayi* populations from the Atlantic (purple and yellow lineages), while the second axis divided the two lineages of *P. cf. grayi* (Figure 4b). The topology of the tree based on the Chord distance ($D_c$) supported the distinction between the Atlantic and the Mediterranean populations with a bootstrap value of 85 (Figure 4c).

The AMOVA revealed highly significant genetic structuring between the three groups (i.e. lineages), among populations, and among individuals ($p < 0.001$; Table S5). The genetic variation explained by differences among groups was 14%, higher than that explained by differences among populations (10%) or among individuals (6%). The global $G_{ST}$ and $D_{EST}$ were 0.17 and 0.53, respectively ($p < 0.001$). Pairwise $G_{ST}$ among lineages were 0.24, 0.12 and 0.11 for *P. clavata* vs. yellow lineage, yellow vs. purple *P. cf. grayi* lineages and *P. clavata* vs. purple lineage, respectively. Overall, pairwise $G_{ST}$ ranged between 0.01 (PPL vs. MED) and 0.46 (AYV and BAL), with only 31 non-significant comparisons ($< 3\%$ of total; Table S6). Pairwise $D_{EST}$ ranged between 0.01 (PPL vs. MED) and 0.99 (AYV and SEG). All except 33 pairwise comparisons (less than 5%) were significant. Focusing on the Atlantic populations, the values of the pairwise $G_{ST}$ and $D_{EST}$ were within the range of the values observed among Mediterranean populations (0.04 [SEG vs. BAL] - 0.31 [FAR vs. BAL] for $G_{ST}$; 0.06 [SEG vs. BAL] - 0.67 [RAB vs. BAL] for $D_{EST}$). For the two estimators, all but one pairwise comparison (SEG vs. BAL) were significant.
Discussion

This study discovered novel species diversity in the gorgonian genus *Paramuricea* in the Iberian Peninsula, revealing recently diverged species found predominantly in the Mediterranean (*P. clavata*) and in the Atlantic (*P. cf. grayi*), where two colour morphotypes (yellow and purple) remain genetically distinct in sympathy, indicating also speciation within *P. cf. grayi*. These discoveries result from integrating genetic data from three independent types of markers (microsatellites, single-copy orthologues and a mitochondrial gene) to assess the extent of divergence between Mediterranean populations of the gorgonian *P. clavata* and adjacent Atlantic populations occurring along southern and western Portugal, suggested to represent the western limit of the species' distribution range. The three phylotranscriptomic approaches used here (ML phylogenetic inference, SODA species delimitation and PoMo species tree reconstruction) strongly support the Atlantic and Mediterranean clades being two distinct sister species: *P. cf. grayi* in the Atlantic (shared *mt-mutS* haplotype with published *P. grayi*), a species previously described for this region; and *P. clavata* in the Mediterranean. Our phylogenetic results are also congruent with genetic structure at microsatellite loci and with a recent phylogenomic study for *P. grayi*/*P. clavata* collected within the Mediterranean. Although based on a limited number of samples, two statistically consistent species tree methods (SODA, based on the ASTRAL species tree, and PoMo, incorporating within population genetic diversity) revealed the genetic segregation of colour morphotypes within Atlantic *P. cf. grayi*, and thus an additional putative (or ongoing) speciation event between purple and yellow lineages. Although these lineages display considerable geographic structure, with the yellow morphotype largely dominant in the Algarve (south Portugal) and the purple dominating habitats on the west coast of Portugal, they can also be found in sympatry (e.g., site P39).

**Lineage divergence across the Atlantic-Mediterranean transition**

Previous genetic studies of Atlantic *P. cf. grayi* (heretofore *P. clavata*) occurring along Portugal and Mediterranean populations of *P. clavata* identified high levels of population differentiation between the two regions, indicating highly restricted gene flow across the Atlantic-Mediterranean transition. Based on comprehensive sampling, our microsatellite analyses corroborate these early findings, revealing a clear segregation between *P. cf. grayi* and *P. clavata*. The uniqueness of the Atlantic populations of *P. cf. grayi* and the relationship with *P. clavata* was also evident in the STRUCTURE and DAPC analyses (Figure 4), which showed a two-level hierarchical partitioning of genetic variation that agreed with the results of the phylogenetic analyses based on single-copy orthologs alignments (Figures 2 and 3), as well as with the statistically significant differences at most genetic parameters analysed. Private allelic richness (*A_p*), in particular, was substantially higher in *P. cf. grayi*, which included seven of the highest *A_p* values observed across all populations studied. This pattern is more consistent with the long-term persistence of distinct lineages, rather than Atlantic populations being derived as marginal extensions of a Mediterranean expansion. Indeed, all phylogenetic methods, as well as a distance-based approach, indicate that Mediterranean *P. clavata* is more genetically divergent from the adjacent Atlantic populations of *P. cf. grayi* (yellow lineage) than from the more geographically distant populations in western Portugal (purple lineage).

Although our analyses did not consider morphological data, the ML analysis of the *mt-mutS* barcode showed that the samples of the Atlantic lineage(s) identified here have the same haplotype as the morphospecies *P. grayi* from northern Iberia (Galicia and Bay of Biscay; Poliseno et al., 2017) and distinct from that of the sister species *P. clavata* (Figure S1). Thus, while we cannot formally reject the occurrence of *P. clavata* in adjacent Atlantic coral assemblages (e.g., unsampled populations in northern Morocco), our results confirm the identity of all southern and western Iberian populations studied here as *P. cf. grayi*. This finding agrees with previous morphological identifications documenting the presence of *P. grayi* in southern Portugal, particularly in circalittoral rocky habitats along the Algarve and in the Gorringe Bank Monteiro-Marques (1987).

The sister relationship between *P. grayi* and the Mediterranean endemic *P. clavata* has been recently described in two studies. Based on *mt-mutS* sequence data, estimated that the divergence between the two species occurred 2.6 Ma (4-1 Ma) or 4.6 Ma (7-3 Ma) assuming a mutation rate of 0.14% or 0.25% Myr-
noteworthy that our STRUCTURE analysis identified a signal of admixture between the two lineages at preliminary evidence for a diagnostic morphological trait in distinguishing the two Atlantic lineages. It is irrespective of geographic origin, including sympatric individuals (P39 site, Figures 1-3), which is important clavata and P. More recently, Quattrini et al. (2022) used RAD-seq data to estimate the timing of divergence between P. clavata and P. grayi at 2 Ma (3.3-1.0 Ma 95% CI), with Bayesian dispersal- extirpation-cladogenesis modelling suggesting that divergence likely occurred in sympathy (though the analysis only included 3 individuals collected within the Mediterranean). Our STRUCTURE analysis based on microsatellite loci identified a signal of admixture between P.cf. grayi and P. clavata at sites in the Alboran sea (MIR, TYL, MTH, located immediately east of the Strait of Gibraltar). Here and in two additional sites east of the AOF (CPS and CAB), the coefficients of membership to either of the Atlantic/Mediterranean clusters were generally lower (Figure 4a). This suggests that the Alboran sea is an area where the two species come into secondary contact, presumably following vicariant speciation. Interestingly, the admixture signal indicates that introgression occurs predominantly from the Atlantic (i.e., P. cf. grayi ) into the Mediterranean (i.e., P. clavata ) gene pool, and involves alleles found in the purple lineage of P. cf. grayi (Figure 4a). However, this inferred directionality could be a sampling artefact, since Atlantic samples from the adjacent northern Moroccan coastline are currently unavailable. Such a pattern of admixture has been described in other Atlantic-Mediterranean sister taxa/lineages , and can be explained by the inflow of Atlantic surface seawater into the Mediterranean resulting from water balance deficits inherent during interglacial periods, which contributes to the continued penetration of Atlantic fauna and flora . While further research is needed to fully resolve the evolutionary history of P. grayi and P. clavata in the Atlantic-Mediterranean, our findings seem to support suggestions thatP. clavata is a “neoendemic” species that diverged from a subtropical Atlantic ancestor that colonized the Mediterranean following the Zanclean flood . The divergence ages estimated recently suggest that the Alboran sea (through the Strait of Gibraltar and AOF) has acted as a buffer reducing admixture between P. cf. grayi and P. clavata throughout the Quaternary, as documented previously for other taxa as a consequence of the coupling between oceanographic barriers (here AOF) and genetic barriers .

The low and moderate values of concordance factors obtained for genes (gCFs) and sites (sCFs) supporting the Atlantic and Mediterranean clades provide further evidence for a recent divergence between P. clavata and P. cf. grayi (Figure 2). Gene-tree discordance is generally attributed to two processes: 1) incomplete lineage sorting (ILS) of ancestral polymorphisms; and 2) gene flow between taxa via introgressive hybridization . Introgression is considered a pivotal process countering lineage diversification of both plants and animals, which is often observed in rapidly (or recently) diverging lineages of closely related species , including several coral lineages . Disentangling ILS and introgressive hybridization is often difficult because they generate similar genomic signatures . However, the congruency between species tree methods (i.e., concatenation and statistically consistent ASTRAL and PoMo models) and between phylogenies based on nuclear and mitochondrial loci analysed here, as well as evident admixture in the Alboran contact zone suggest that ILS is a more likely explanation for the overall low values of gCFs and sCFs supporting the P. cf. grayi and P. clavata clades.

Genetic segregation within the Atlantic lineages

Our results clearly show lineage divergence within Iberian P. cf. grayi . The microsatellite analysis revealed that populations in southern Portugal (Sagres, Lagos and Tavira) were highly differentiated from those sampled along the western coast (Berlengas, Cape Espichel and Sines), with highly significant pairwise GST s and DEST s and genetic partitioning (Table S6 and Figure 4). This south-west genetic divide is coincident with the distribution of yellow and purple colour morphotypes noted in previous studies examining a subset of the populations . Importantly, our ML and species trees support colour morph segregation in P. cf. grayi irrespective of geographic origin, including sympatric individuals (P39 site, Figures 1-3), which is important preliminary evidence for a diagnostic morphological trait in distinguishing the two Atlantic lineages. It is noteworthy that our STRUCTURE analysis identified a signal of admixture between the two lineages at
sites where field surveys confirmed them to occur in sympatry (CAT and SIN; Figure 4a). Collectively, these findings and observations suggest a high degree of habitat segregation (and partial reproductive isolation) between the two morphotypes of *P. cf. grayi*, calling for additional analyses using genomic data, broader taxon and habitat (e.g., depth) sampling, as well as experimental approaches (e.g., crossing experiments) to clarify the taxonomic status of the two colour morphs.

Within segregating lineages of *P. cf. grayi*, we have identified significant pairwise genetic differentiation between most populations studied, including at local spatial scales, from ~20 kilometres (e.g., BAL vs. LAG, southern Portugal) to less than 400 m (e.g., COR vs. CAT, western Portugal) (Table S6). These results suggest restricted gene flow between populations of both Atlantic lineages, with rare long-distance dispersal. The extent to which such genetic differentiation is driven by seascape (i.e. physical environment), life-history traits of the species or the interplay between both remains speculative at this point. However, these findings raise several important questions about the biology of *P. cf. grayi*. For instance, strong genetic differentiation at small spatial scales and a pattern of isolation by distance are common in many brooding corals in which dispersal potential is generally low given the short duration of the pelagic larval stage and predominantly philopatric recruitment. Interestingly, the spatial genetic patterns described for *P. clavata* in the Mediterranean coincide with the low effective dispersal inherent to surface brooding inferred from field observations and parental genetic analysis, while future studies could indicate whether *P. cf. grayi* is also a brooder. We note, however, that reproductive mode in octocorals is relatively plastic even at the congeneric level (e.g., *Antillogorgia*; and *Aleyonium*;).

**Concluding remarks**

In this study we investigated population differentiation and speciation in key structural species of octocorals in marine ecosystems of the NE Atlantic and Mediterranean Sea. The results presented advance the understanding of lineage diversification in these important biogeographic regions. Our analyses revealed that past species identifications based on morphological characters and a limited number of genetic markers have underestimated species diversity in *Paramuricea* occurring in shallow-water and circalittoral habitats of southern Iberia, which has important implications for conservation planning. Specifically, our results provide compelling evidence defining species boundaries between Atlantic populations, here assigned to the morphospecies *P. cf. grayi* and the Mediterranean *P. clavata*; as well as between two segregating lineages within the *P. cf. grayi* clade that are coincident with coloration of the colonies, thus casting doubts about the taxonomic status of these two entities.

Although the broader species limits between *P. cf. grayi* and *P. clavata* could be resolved with a single mitochondrial barcode (*mt-mutS*), this study emphasizes the need for genome-wide multi-locus datasets to resolve recent lineage splitting events in the face of, e.g., ILS and/or introgression. The marked spatial genetic structure and differentiation observed between the *P. cf.grayi / P. clavata* clades reveal historical footprints of divergence across the Atlantic-Mediterranean transition, the corresponding evolutionary scenarios of which could be tested with further population genomic data. Like recent studies, our results emphasize the urgent need to re-evaluate species diversity and the processes underlying diversification within the genus *Paramuricea*, which despite its ecological importance, likely harbours high levels of cryptic diversity (e.g., several species described as amphi-Atlantic; . Finally, cryptic diversity that went undetected in previous molecular and morphological studies validates the usefulness of phylotranscriptomics for resolving the evolutionary relationships of octocorals, and more generally of diverse lineages of marine taxa, and which can be a reliable (and cheaper) alternative to emerging studies based on genome sequence data.

**Acknowledgements**

This work was funded by program BIOMARES, a Pew Marine Fellowship, and FCT - Foundation for Science and Technology UIDB/04326/2020, UIDP/04326/2020, LA/P/0101/2020, PTDC/BIA-CBI/6515/2020; EU-BiodivERsA BiodivRestore-253 (FCT: DivRestore/0013/2020); EU Horizon 2020 research and innovation programme (MERCES: Grant no. 689518); as well as the Inqua Conservation Fund (Oceanario de Lisboa and National Geographic Channel) and National Geographic Society/Waitt Grant No. W153-11. This ar-
Article is also a product of the French network on Marine Connectivity (GDR MarCo). M.C. was supported by postdoctoral fellowships of projects HABMAR (Grant No. MAR-01.04.02-FEAMP-0018) co-financed by the European Maritime and Fisheries Fund of the Operational Program MAR 2020 for Portugal (Portugal 2020), and BiodivAMP (Grant no. FA_06_2017_045) financed by the Directorate-General for Sea Policy of the Ministry of Economy and Sea for Portugal under the Operational Program Fundo Azul. JBL was funded by assistant researcher contract framework of the RD Unit—UID/Multi/04423/2019 – Interdisciplinary Centre of Marine and Environmental Research—financed by the European Regional Development Fund (ERDF) through COMPETE2020 – Operational Program for Competitiveness and Internationalization (POCI) and national funds through FCT/MCTES (PIDDAC). The authors would like to thank the help during fieldwork by volunteer divers and students of project Deep Reefs, in particular Delfim Machado, Luis Magro, Francisco Fernandes, Joao Rodrigues, Joao Pedro Fonseca, Goncalo Ventura Martins, Bartek Ciperling, Pieter Truter, Ricardo Constantino, Joanna Pylicinska and Nikola Moraj; as well as volunteers of project MERCES for assisting with lab work and field collections, in particular I. Montero-Serra, M. Pages-Escola, S. Ochoa, D. Petricioli and A. Medrano; and N. E. Topcu (Istanbul University) for sample collections in Turkey. M. C. thanks R. Jacinto and C. Paulino for help with the RNA extractions, F. Oliveira for helpful discussions concerning the distribution of *P. cf. grayi* along Portugal and A.C. Ferreira for providing photographs of *P. cf. grayi*. All sampling was conducted in accordance with local legislation.

References

Data availability statement

Raw RNA-seq data and mt-*mutS* sequences have been uploaded to NCBI’s SRA (BioProject ID: PR-610199) and GenBank databases (Accessions numbers: provided upon manuscript acceptance), respectively. The accession numbers will be provided upon manuscript acceptance. Final sequence alignments, transcriptome assembly, and microsatellite data are available on DRYAD (doi provided upon acceptance).

Author Contributions

M.C., G.P., J.B., E.S. and J.-B.L. designed research, performed data analyses and wrote the manuscript. D.A. contributed to the data analyses. M.C., J.B., D.P., D.G.-G., N.B., P.L.-S., C.C., S.K., T.B.-P., E.F., C.L., J.G. and J.-B.L. collected samples. P.L.-S., M.C., J.B. and J.-B.L. conducted laboratory work. E.S., J.B. and J.G. contributed funding. All authors revised the manuscript and gave their approval for publication.

Tables

Table 1 - Summary statistics for the top transcriptome assembly obtained with rnaSPAdes using samples of *P. cf. grayi* from Baleeira, Portugal (NE Atlantic).

<table>
<thead>
<tr>
<th>Metric</th>
<th><em>P. cf. grayi</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Original contigs (rnaSPAdes)</td>
<td>524,051</td>
</tr>
<tr>
<td>Good contigs (Transrate)</td>
<td>482,166</td>
</tr>
<tr>
<td>ORFs (FragGeneScan)</td>
<td>688,540</td>
</tr>
<tr>
<td>Clustered ORFs (VSEARCH)</td>
<td>544,317</td>
</tr>
<tr>
<td>Final contigs (Anthozoa top hits)</td>
<td>73,871</td>
</tr>
<tr>
<td>Longest sequence (nt)</td>
<td>48,483</td>
</tr>
<tr>
<td>Shortest sequence (nt)</td>
<td>108</td>
</tr>
<tr>
<td>Mean sequence length (bp)</td>
<td>969</td>
</tr>
<tr>
<td>Median sequence length (bp)</td>
<td>654</td>
</tr>
<tr>
<td>N50 (bp)</td>
<td>1,452</td>
</tr>
<tr>
<td>Sequences &gt; 1k bp</td>
<td>24,579 [33.3%]</td>
</tr>
<tr>
<td>Sequences &gt; 10k bp</td>
<td>44 [0.1%]</td>
</tr>
<tr>
<td>G-C content (%)</td>
<td>45.4</td>
</tr>
</tbody>
</table>
Figure legends

Figure 1 - Map showing the sampling locations of Paramuricea across the NE Atlantic Ocean and Mediterranean Sea (A). Site codes follow the abbreviations of Table S1. Sites sampled for RNA-seq are underlined, with populations that were also sampled for microsatellite genotyping shown in bold. PHL to GAB - all sites between Pharillons in the South of Marseille and Gabiniere in Port-Cros (France) as listed in Table S1; PZU to GGL – all sites in northern Corsica (France). For a detailed visualization of the location of Mediterranean sites see Figure 1 of Mokhtar-Jamai et al. (2011). AOF - approximate location of the Almeria-Orán Oceanographic Front. (B) Detail of sampling sites along southern Iberia (NE Atlantic) and Alboran Sea (Mediterranean) separated by the Strait of Gibraltar; insets show sites sampled in Cape Espichel (upper) and Sagres (lower) with bathymetric isobaths shown as follows: 50 m, 100–500 m (increments of 100 m) and >500 m (increments of 200 m). (C) Locations sampled in Banyuls-sur-Mer (southern France) and northeastern Spain, including the Medes Islands. (D) Locations sampled along the eastern Adriatic Sea (Croatia). Colours of the markers correspond to species identified by SODA analysis (see Results). Note that both the purple and yellow morphotypes of the P. cf. grayi populations were sampled at P39 (Cape Espichel, western Portugal).

Figure 2 - Phylogenetic relationships in *Paramuricea* determined by ML partition analysis in IQ-TREE 2. The tree was built using an alignment of 261 single-copy orthologues with a separate evolutionary model per gene (see text for details). The colours correspond to species identified by SODA analysis (plus outgroup, *P. biscaya* ); text boxes to the right indicate geographic region from which samples originate. Red arrows indicate selected key branches with ultrafast bootstrap support (1000 replicates): gCF / sCF = gene / site concordance factors supporting the branch indicated. For the NE Atlantic, the species name is given as *P. cf. grayi* with colour morphotypes indicated in yellow and purple. The codes of the sites from which the samples originate are embedded in the taxon names (P39, BAL, TAV, VAC, ALT and BALU) and follow the nomenclature used in Figure 1 and Table S1. Photo credits: A.C. Ferreira (*P. cf. grayi*) and J. Garrabou (*P. clavata*).

Figure 3 - Species tree of *Paramuricea* samples in this study using the ML polymorphism-aware (PoMo) model and HKY substitution model in IQ-TREE 2. The tree was built using allelic counts from 261 single-copy orthologues. Values on the branches indicate ultrafast bootstrap support (1000 replicates). The colours correspond to species identified by SODA analysis (plus outgroup, *P. biscaya* ). For the NE Atlantic, the species name is given as *P. cf. grayi* with colour morphotypes (lineages) indicated in yellow and purple. The codes of the sites from which the samples originate are embedded in the taxon names (P39, BAL, TAV, VAC, ALT and BALU) and follow the nomenclature used in Figure 1 and Table S1. Photo credits: A.C. Ferreira (*P. cf. grayi*) and J. Garrabou (*P. clavata*).

Figure 4 - Spatial genetic structure in *Paramuricea*: (a) Clustering analysis conducted with STRUCTURE considering K=2 and K=3. Each individual is represented by a vertical line partitioned in K-colored segments, which represent the individual membership fraction in K clusters. Thin black vertical lines delineate the different locations of the samples while dashed and thick lines show the segregating lineages of *P. cf. grayi* (PPL for ‘purple’ and YLW for ‘yellow’; see results below) and the two oceanographic basins (Atlantic vs. Mediterranean), respectively. Samples names are shown below the assignment plots (for abbreviations see Table S1). The mean membership coefficient for each cluster is also shown. (b) Scatter plot of the discriminant analysis of principal components (DAPC) based on an a-score of 80 and considering axes 1 and 2. These two axes represented 51.1 % of the total variation in the data. Each dot corresponds to one individual (n=1223) from each of the 46 locations. Note that the colours correspond to the three clusters with the red dots for the Mediterranean samples of *P. clavata* and the yellow and purple dots for the Atlantic samples of *P. cf. grayi*. Only the names of the Atlantic samples are shown. (c) Dendrogram based on the chord distance (Cavalli-Sforza & Edward, 1967) showing the relationships among the 46 populations. Colours are the same as for K=3 while the sample names are shown in Table S1. Only bootstrap values higher than 75 are shown.
Figures

Figure 1
Hosted file

Figure_2_ML_CF_tree.pdf available at https://authorea.com/users/489793/articles/573366-not-out-of-the-mediterranean-atlantic-populations-of-the-gorgonian-paramuricea-clavata-are-a-separate-sister-species-under-further-lineage-diversification

Hosted file

Figure3_PoMo_spTree.pdf available at https://authorea.com/users/489793/articles/573366-not-out-of-the-mediterranean-atlantic-populations-of-the-gorgonian-paramuricea-clavata-are-a-separate-sister-species-under-further-lineage-diversification