

Molecular phylogenies confirm the presence of two cryptic *Hemimycale* species in the Mediterranean and reveal the polyphyly of the genera *Crella* and *Hemimycale* (Demospongiae: Poecilosclerida)

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ABSTRACT

Background: Sponges are particularly prone to hiding cryptic species as their paradigmatic plasticity often favors species phenotypic convergence as a result of adaptation to similar habitat conditions. *Hemimycale* is a sponge genus (Family Hymedesmiidae, Order Poecilosclerida) with four formally described species, from which only *Hemimycale columella* has been recorded in the Atlanto-Mediterranean basin, on shallow to 80 m deep bottoms. Contrasting biological features between shallow and deep individuals of *Hemimycale columella* suggested larger genetic differences than those expected between sponge populations. To assess whether shallow and deep populations indeed belong to different species, we performed a phylogenetic study of *Hemimycale columella* across the Mediterranean. We also included other *Hemimycale* and *Crella* species from the Red Sea, with the additional aim of clarifying the relationships of the genus *Hemimycale*.

Methods: *Hemimycale columella* was sampled across the Mediterranean, and Adriatic Seas. *Hemimycale arabica* and *Crella cyathophora* were collected from the Red Sea and Pacific. From two to three specimens per species and locality were extracted, amplified for Cytochrome C Oxidase I (COI) (M1–M6 partition), 18S rRNA, and 28S (D3–D5 partition) and sequenced. Sequences were aligned using Clustal W v.1.81. Phylogenetic trees were constructed under neighbor joining (NJ), Bayesian inference (BI), and maximum likelihood (ML) criteria as implemented in Geneious software 9.01. Moreover, spicules of the target species were observed through a Scanning Electron microscope.

Results: The several phylogenetic reconstructions retrieved both *Crella* and *Hemimycale* polyphyletic. Strong differences in COI sequences indicated that *C. cyathophora* from the Red Sea might belong in a different genus, closer to *Hemimycale arabica* than to the Atlanto-Mediterranean *Crella* spp. Molecular and external morphological differences between *Hemimycale arabica* and the Atlanto-Mediterranean *Hemimycale* also suggest that *Hemimycale arabica* fit in a separate genus. On the other hand, the Atlanto-Mediterranean Crellidae appeared in 18S and 28S phylogenies as a sister group of the Atlanto-Mediterranean *Hemimycale*. Moreover, what was known up to now as *Hemimycale columella*, is formed by

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two cryptic species with contrasting bathymetric distributions. Some small but consistent morphological differences allow species distinction.

Conclusions: A new family (Hemimycalidae) including the genus *Hemimycale* and the two purported new genera receiving *C. cyathophora* and *Hemimycale arabica* might be proposed according to our phylogenetic results. However, the inclusion of additional Operational Taxonomic Unit (OTUs) appears convenient before taking definite taxonomical decisions. A new cryptic species (*Hemimycale mediterranea* sp. nov.) is described. Morphologically undifferentiated species with contrasting biological traits, as those here reported, confirm that unidentified cryptic species may confound ecological studies.

Subjects Aquaculture, Fisheries and Fish Science, Biodiversity, Marine Biology, Taxonomy, Zoology

Keywords Biodiversity, Molecular taxonomy, Morphological taxonomy, Cryptic species, Marine sponges, Phylogeny, *Hemimycale mediterranea*, *Hemimycale columella*, *Hemimycale arabica*, *Crella cyathophora*

INTRODUCTION

The discovery of cryptic species is continuously improving our knowledge on real ecosystem biodiversity and functioning, which are intimately related (*Frainer, McKie & Malmqvist, 2014*). Unrecognized cryptic diversity may mask biological features such as divergent reproduction patterns, growth dynamics, and inter-species interactions, among others (*Knowlton, 1993; Prada et al., 2014; de Meester et al., 2016; Loreau, 2004*), which may confound conservation studies (*Forsman et al., 2010*) and obscure the introduction pathway of invasive species (*Knapp et al., 2015*).

Molecular tools help to confirm suspected hidden species. However, molecular based identifications alone do not solve the problem of species misidentification, in particular when the cryptic species have overlapping distributions (e.g., *Knowlton & Jackson, 1994; Tarjuelo et al., 2001; De Caralt et al., 2002; Blanquer & Uriz, 2007, 2008; Pérez-Portela et al., 2007*). In these cases, deep studies on their morphology, biology (e.g., life-history traits), and ecology (e.g., growth dynamics) become crucial to understand the mechanisms underlying their coexistence (*López-Legentil et al., 2005; Pérez-Portela et al., 2007; Blanquer, Uriz & Agell, 2008; Payo et al., 2013*).

Sponges are sessile, aquatic filter-feeders that are widespread across oceans, depths, and ecosystems (*Van Soest et al., 2012*), with so far 8,789 accepted species inventoried in 2016 (*Van Soest et al., 2016*) and ca. 29,000 predicted to be discovered in the forthcoming years (*Hooper & Lévi, 1994; Appeltans et al., 2012*), many of which remain currently hidden among supposed widespread morpho-species (*Uriz & Turon, 2012*).

The poor dispersal capacities of sponges prevent in most cases gene flow among populations even at short geographical distances (*Boury-Esnault, Pansini & Uriz, 1993; Uriz et al., 1998; Nichols & Barnes, 2005; Mariani et al., 2006; Uriz, Turon & Mariani, 2008*). Consequently, sponge populations become genetically structured (*Boury-Esnault, Pansini & Uriz, 1993; Duran, Pascual & Turon, 2004; Blanquer, Uriz & Caujapé-Castells, 2009; Guardiola, Frotscher & Uriz, 2012, 2016*), which favors speciation, while the

sponge plasticity fosters phenotypic (morphological) convergence to similar habitats (Blanquer & Uriz, 2008).

Many new cryptic sponge species have been discovered in the last decades thanks to the use of molecular markers (see Uriz & Turon, 2012 for a review until 2012, Knapp et al., 2011; de Paula et al., 2012). However, less often, molecularly discovered new species have also been described morphologically (but see Blanquer & Uriz, 2008; Cárdenas & Rapp, 2012; Reveillaud et al., 2011, 2012), which is necessary if phylogeny is aimed to translate into taxonomy, and the new species are wanted to be considered in ecological studies.

Sponge species can be both morphologically (e.g., Uriz & Turon, 2012) and, more rarely, molecularly (with the markers used) cryptic (Carella et al., 2016; Vargas et al., 2016) but show contrasting biological features. For instance, *Scopalina blanensis* (Blanquer & Uriz, 2008), which is sympatric with *Scopalina lophyropoda*, mainly grows in winter. Conversely, *Scopalina lophyropoda* regresses in winter and grows principally in summer–autumn (Blanquer, Uriz & Agell, 2008), thus indicating temporal niche partition.

The Order Poecilosclerida (Porifera: Demospongiae) harbors the highest number of species within the Class Demospongiae (Systema Porifera) and it is far from being resolved from a phylogenetic point of view (Morrow et al., 2012; Thacker et al., 2013). Within Poecilosclerida, the Family Hymedesmiidae represents a hotchpotch where genera of dubious adscription have been placed (Van Soest, 2002). As expected, this family appeared clearly polyphyletic in a molecular phylogeny of the so-called G4 clade based on 28S rRNA gene (Morrow et al., 2012).

Hymedesmiidae currently contains 10 accepted genera among which, *Hemimycale* Burton, 1934 (Van Soest et al., 2016). The position of genus *Hemimycale*, which shares with *Hymedesmia*, and *Phorbas* (Hymedesmiidae) and with *Crella* (Crellidae), the so-called aerolate areas with an inhaling function, has changed from Hymeniacionidae in Halichondrida (Lévi, 1973) to Hymedesmiidae in Poecilosclerida (Van Soest, 2002). More recently, in 18S phylogenies of Poecilosclerida, *Hemimycale columella* was retrieved within the Crellidae clade, although with low support (Redmond et al., 2013).

Hemimycale harbors only four formally described species (Van Soest et al., 2016): the type species *Hemimycale columella* (Bowerbank, 1874), from Northwestern Atlantic and Mediterranean, *Hemimycale rhodus* (Hentchel, 1929) from the North Sea, *Hemimycale arabica* Illan et al., 2004 from the Red Sea and *Hemimycale insularis* Moraes, 2011 from Brazil. However, the simple spicule complement of the genus, which only consists of strongyles with some occasional styles, may propitiate the existence of morphologically (based on the spicules) cryptic species.

Hemimycale columella, the type species of *Hemimycale*, is widely distributed across the Atlanto-Mediterranean basin, from shallow (ca. 10 m) to deep (ca. 60 m) waters (Uriz, Rossell & Martín, 1992). Assays performed with eight microsatellite loci developed from deep specimens of *Hemimycale columella* (González-Ramos, Agell & Uriz, 2015) failed to amplify a high percentage of the assayed individuals from a shallow population, which suggested larger genetic differences than those expected between intra-species sponge populations.

Furthermore, the life cycle of species has been monitored in a shallow Northwestern Mediterranean population of what was thought to be *Hemimycale columella* (Pérez-Porro, González & Uriz, 2012), where all individuals disappeared after larval release in early November and new individuals arose the forthcoming year but on different rocky sites, which pointed to annual mortality and subsequent recruitment from sexually produced propagula (settling larvae). Conversely, during a study of deeper populations of *Hemimycale columella* (González-Ramos, Agell & Uriz, 2015), we recorded their survival for more than three years. Thus, shallow and deep populations of *Hemimycale columella* seemed to show contrasting life spans, which were thought to be a result of contrasting habitat characteristics. However, a 2-year monitoring of two, some km apart, populations (one deep and one shallow) and the main environmental factors at both locations, confirmed their contrasting life span and growth traits, as well as proved no correlation between biological features and environmental factors (M. J. Uriz, L. Garate & G. Agell, 2013–2014, unpublished data), which rather pointed to population intrinsic (genetic) differences.

To assess whether these two population types with contrasting biological traits but without clearly distinct morphological characters belonged or not to different species, we performed a phylogenetic study of individuals considered as *Hemimycale columella* across the Mediterranean, using three molecular (nuclear and mitochondrial) gene partitions. We incorporated additional species to the analyses to gain knowledge on the relationships between *Hemimycale* species and other genera of families Hymedesmiidae and Crellidae.

MATERIALS AND METHODS

Sampling

Fragments of what a priori was thought to be *Hemimycale columella* were collected by SCUBA diving across the Northwestern, central and eastern Mediterranean, and Adriatic Sea, between 12 and 45 m of depth during several campaigns (Coconet, Benthomics, and MarSymbiOomics projects) (Table 1). Moreover, fragments of *Hemimycale arabica* and *Crella cyathophora* from the Red Sea (Dedalos and Ephistone) and Pacific (Bempton Islands) between 5 and 20 m depth were also collected (Table 1). Individuals were photographed underwater before sampling. Collected fragments were divided into two pieces, one of them was preserved in 100% ethanol, and after three alcohol changes, kept at -20°C until DNA extraction; the other fragment was fixed in 5% formalin in seawater and preserved in 70% ethanol as a voucher for morphological and spicule studies. All vouchers have been deposited at the Sponge collection of the Centre d'Estudis Avançats de Blanes (numbers CEAB.POR.GEN.001 to CEAB.POR.GEN.029).

DNA extraction, amplification, and sequencing

DNA extractions were performed on two to three specimens per species and locality (totaling 18 individuals). *Hemimycale* spp. were extracted with QIAmp DNA stool kit (Qiagen), while *Crella* spp. were extracted with DNeasy Blood & Tissue kit (Qiagen) according to the manufacturer's protocol. Standard primers were used for COI partitions

Table 1 Geographical origin and ecological distribution of the individuals used in the phylogenetic study, with accession numbers.

Species	Sea/Ocean	Locality	Voucher numbers	Accession numbers
<i>Hemimyscale arabica</i> ind. 1	Red Sea	Dedalos, Brother Islands	CEAB.POR.GEN.001	COI: KY002124 18S: KY002171 28S: KY002181
<i>Hemimyscale arabica</i> ind. 2	Red Sea	Elphinstone, Brother Islands	CEAB.POR.GEN.002	COI: KY002125 18S: KY002172 28S: KY002182
<i>Hemimyscale columella</i>	Northeastern Atlantic	Plymouth, Wales, UK		28S: HQ379300.1 18S: KC902127.1
<i>Hemimyscale columella</i> ind. 1	Northwestern Mediterranean	Arenys de Mar, Spain	CEAB.POR.GEN.003	28S: KY002183
<i>Hemimyscale columella</i> ind. 2	Northwestern Mediterranean	Arenys de Mar, Spain	CEAB.POR.GEN.004	28S: KY002184
<i>Hemimyscale columella</i> ind. 3	Northwestern Mediterranean	Arenys de Mar, Spain	CEAB.POR.GEN.005	COI: KY002126
<i>Hemimyscale columella</i> ind.1	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.006	COI: KY002127 18S: KY002160 28S: KY002185
<i>Hemimyscale columella</i> ind. 2	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.007	COI: KY002128 18S: KY002161 28S: KY002186
<i>Hemimyscale columella</i> ind. 3	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.008	COI: KY002129 28S: KY002187
<i>Hemimyscale columella</i> ind. 4	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.009	28S: KY002188
<i>Hemimyscale mediterranea</i> sp. nov. ind. 1	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.010	COI: KY002130 18S: KY002162 28S: KY002189
<i>Hemimyscale mediterranea</i> sp. nov. ind. 2	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.011	18S: KY002163 28S: KY002190
<i>Hemimyscale mediterranea</i> sp. nov. ind. 4	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.012	COI: KY002131
<i>Hemimyscale mediterranea</i> sp. nov. ind. 5	Northwestern Mediterranean	Tossa de Mar, Spain	CEAB.POR.GEN.013	COI: KY002132
<i>H. mediterranea</i> sp. nov. ind. 3	Adriatic Sea	Koznati, Croatia	CEAB.POR.GEN.014	COI: KY002134
<i>H. mediterranea</i> sp. nov. ind. 7	Adriatic Sea	Koznati, Croatia	CEAB.POR.GEN.015	18S: KY002170 28S: KY002193
<i>H. mediterranea</i> sp. nov. ind. 8	Adriatic Sea	Koznati, Croatia	CEAB.POR.GEN.016	28S: KY002194
<i>H. mediterranea</i> sp. nov. ind. 2	Adriatic Sea	Tremity, Italy	CEAB.POR.GEN.017	COI: KY002133
<i>H. mediterranea</i> sp. nov. ind. 11	Adriatic Sea	Tremity, Italy	CEAB.POR.GEN.018	28S: KY002199
<i>H. mediterranea</i> sp. nov. ind. 8	Central Mediterranean	Porto Cesareo, Italy	CEAB.POR.GEN.019	18S: KY002164
<i>H. mediterranea</i> sp. nov. ind. 9	Central Mediterranean	Porto Cesareo, Italy	CEAB.POR.GEN.020	18S: KY002165 28S: KY002197
<i>H. mediterranea</i> sp. nov. ind. 10	Central Mediterranean	Porto Cesareo, Italy	CEAB.POR.GEN.021	28S: KY002198
<i>H. mediterranea</i> nov. sp. ind. 5	Adriatic Sea	Karaburum, Albania	CEAB.POR.GEN.022	18S: KY002166 28S: KY002191
<i>H. mediterranea</i> nov. sp. ind. 6	Adriatic Sea	Karaburum, Albania	CEAB.POR.GEN.023	18S: KY002167 28S: KY002192

(Continued)

Table 1 (continued).

Species	Sea/Ocean	Locality	Voucher numbers	Accession numbers
<i>H. mediterranea</i> sp. nov. ind. 3	Eastern Mediterranean	Othonoi, Greece	CEAB.POR.GEN.024	18S: KY002168 28S: KY002195
<i>H. mediterranea</i> sp. nov. ind. 4	Eastern Mediterranean	Othonoi, Greece	CEAB.POR.GEN.025	18S: KY002169 28S: KY002196
<i>Crella cyatophora</i> ind.1	Red Sea	Dedalos, Brother Islands	CEAB.POR.GEN.026	COI: KY002120 18S: KY002173 28S: KY002177
<i>Crella cyatophora</i> ind. 2	Red Sea	Elphinstone, Brother Islands	CEAB.POR.GEN.027	COI: KY002121 18S: KY002174 28S: KY002178
<i>Crella cyatophora</i> ind. 3	Pacific	Bempton Patch Reef (between New Caledonian and Australia)	CEAB.POR.GEN.028	COI: KY002122 18S: KY002175 28S: KY002179
<i>Crella cyatophora</i> ind. 4	Pacific	Bempton Patch Reef (between New Caledonian and Australia)	CEAB.POR.GEN.029	COI: KY002123 18S: KY002176 28S: KY002180
<i>Crella elegans</i>	Mediterranean	France		18S: KC902282
<i>Crella elegans</i>	Mediterranean	France		18S: AY348882
<i>Crella elegans</i>	Mediterranean	France		28S: HQ393898
<i>Crella plana</i>	Northeastern Atlantic	Northern Ireland		18S: KC9023009
<i>Crella rosea</i>	Northeastern Atlantic	Northern Ireland		28S: HQ379299
<i>Crella rosea</i>	Northeastern Atlantic	Northern Ireland		18S: KC902282
<i>Phorbas bihamiger</i>	Northeastern Atlantic	English Channel		18S: KC901921.1 28S: KC869431
<i>Phorbas punctatus</i>	Northeastern Atlantic	Wales		18S: KC869439.1 28S: KC869439.1
<i>Phorbas dives</i>	Northeastern Atlantic	English Channel		28S: HQ379303
<i>Phorbas fictitoides</i>	North Pacific	–		COI: HE611617.1
<i>Phorbas tenacior</i>	Northeastern Atlantic	–		18S: AY348881
<i>Phorbas glaberrimus</i>	Antarctic	Ross Sea		COI: LN850216.1
<i>Hymedesmia paupertas</i>	Northeastern Atlantic			18S: KC902073.1 28S: KF018118.1
<i>Hymedesmia pansa</i>				18S: KC902027.1
<i>Hymedesmia paupertas</i>	Northeastern Atlantic			28S: KF018118.1
<i>Kirkpatrickia variolosa</i>	Antarctic	Ross Sea		COI: LN850202.1

Note:

Individuals sequenced de novo are in bold.

M1–M6 (Folmer *et al.*, 1994) and 18S rRNA (1F and 1795R, from Medlin *et al.*, 1988), and Porifera primers for the D3–D5 partition of 28S rRNA (Por28S–830F and Por28S–1520R, from Morrow *et al.*, 2012). Different amplification protocols were performed for each gene (Table 2). COI (M1–M6 partition) amplifications were performed in a 50 μ L volume reaction, containing 37.6 μ L H₂O, 5 μ L buffer KCl (BIORON; F Holzinger Sales & Support, Germany), 2 μ L BSA, 2 μ L dNTP (Sigma; Sigma_Aldrich, Germany), 1 μ L of primers, 0.4 μ L Taq (BIORON; F Holzinger Sales & Support, Germany), and 1 μ L of genomic DNA. 18S rRNA amplifications were performed in a 50 μ L volume reaction,

Table 2 PCR conditions for the three partitions used (COI, 28S and 18S).

PCR Stage	COI (M1–M6)	28S (D3–D5)	18S
Denaturalization	94 °C 2 min 35 cycles	94 °C 5 min 35–40 cycles	94 °C 5 min 30 cycles
Denaturalization	94 °C 1 min	94 °C 1 min	94 °C 30 s
Annealing	43 °C 1 min	50–55 °C 1 min	53 °C 30 s
Elongation	72 °C 1 min	72 °C 1 min	72 °C 30 s
Final elongation	72 °C 5 min	72 °C 5 min	72 °C 5 min

containing 36.85 μL H_2O , 5 μL buffer (Invitrogen, Carlsbad, CA, USA), 0.75 μL MgCl (Invitrogen, Carlsbad, CA, USA), 1.2 μL DMSO (dimethyl sulfoxide), 1 μL BSA, 1.5 μL dNTP (Sigma; Sigma_Aldrich, Germany), 1 μL of primers, 0.7 μL Taq (Invitrogen, Carlsbad, CA, USA), and 1 μL of genomic DNA. Finally, partition D3–D5 of 28S rRNA amplifications were performed in a 50 μL volume reaction, containing 36.85 μL H_2O , 5 μL buffer (Invitrogen, Carlsbad, CA, USA), 0.75 μL MgCl (Invitrogen, Carlsbad, CA, USA), 2 μL BSA, 2 μL dNTP (Sigma; Sigma_Aldrich, Germany), 1 μL of primers, 0.4 μL Taq (Invitrogen, Carlsbad, CA, USA), and 1 μL of genomic DNA. Polymerase chain reaction products were purified and sequenced in both directions using Applied Biosystems 3730xl DNA analyzers in Macrogen, Korea.

Sequence alignment and phylogenetic reconstructions

Sequences of COI, 28S, and 18S were aligned using Clustal W v.1.81, once their poriferan origin was verified using BLAST (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>), as implemented in Genieous 9.01 (Kearse et al., 2012). When sequences were identical, only one sequence per locality and species was included in the phylogenetic trees. After alignment, ambiguous regions were determined with Gblocks v.091 b software (Castresana, 2000), which removes from 1 to 4% of poorly aligned positions and divergent regions of an alignment of DNA. Representatives of family Hymedesmiidae (i.e., genera *Phorbas* and *Hymedesmia*) and Crambeidae (i.e., genera *Crambe* and *Monanchora*) were selected as outgroups. The inclusion of Crambeidae as an outgroup was decided because the species *Hemimycala arabica* had been reported to contain similar secondary metabolites (polycyclic guanidine alkaloids) to those of *Crambe* and *Monanchora* (Ilan et al. 2004).

JModelTest 0.1.1 (Posada, 2008) was used to determine the best-fitting evolutionary model for each dataset. The model GTR + I + G was used for both mitochondrial and nuclear genes. Phylogenetic trees were constructed under neighbor joining (NJ) (default parameters), Bayesian inference (BI), and maximum likelihood (ML) using Geneious software 9.01 (Kearse et al., 2012). NJ generates unrooted minimum evolution trees (Gascuel & Steel, 2006). BI analyses were performed with MrBayes 3.2.1 (Ronquist & Huelsenbeck, 2003). Four Markov Chains were run with one million generations sampled every 1,000 generations. The chains converged significantly and the average standard deviation of split frequencies was less than 0.01 at the end of the run. Early tree generations were discarded by default (25%) until the probabilities reached a stable

plateau (burn-in) and the remaining trees were used to generate a 50% majority-rule consensus tree. ML analyses were executed with PhyML v3.0 program (Guindon & Gascuel, 2003; Guindon et al., 2005). The robustness of the tree clades was determined by a nonparametric bootstrap resampling with 1,000 replicates in PhyML. MrBayes and PhyML were downloaded by Genieious.

Incongruence length difference (ILD) test (PAUP 4.0b10) was run (Swofford, 2002) to verify sequence homogeneity or incongruence between the 18S rRNA and COI markers and the 18S and 28S rRNA markers. The ILD test indicated no significant conflict ($p = 0.93$ and $p = 0.91$, respectively) between the marker pairs to be concatenated. Thus, concatenated 18S COI and 18S–28S rRNA datasets were constructed for the species with sequences available for both markers. The phylogeny on the three genes concatenated was not performed due to the few species/individuals for which the three genes were available.

Phenotypic characters

To assess whether molecular differences among the target populations and species (*Hemimycale columella*, *Senso latus*, *Hemimycale arabica*, and *C. cyathophora*) were supported by morphological and spicule traits, the target species were observed both in situ and on recently collected samples. Moreover, spicules of all the species were observed through light and scanning electron microscopes (SEM) after removing the sponge organic matter from small (3 mm³) pieces of each individual by boiling them in 85% Nitric acid in a Pyrex tube and then washed three times with distilled water and dehydrated with ethanol 96% (three changes). A drop of a spicule suspension in ethanol per individual was placed on 5 mm diameter stuffs, air dry, and gold–palladium metalized (Uriz, Turon & Mariani, 2008) in a Sputtering Quorum Q150RS. Observation was performed through a Hitachi M-3000 Scanning Electron Microscope at the Centre d'Estudis Avançats de Blanes.

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RESULTS

18S rRNA phylogeny

The resulting phylogeny using the 18S rRNA partition on 25 sequences (17 new) of 695 nt. (46 variable positions, from which 38 were parsimony informative) was congruent under BI, and ML and just differed in the position of *Hemimycale arabica* which appeared as a sister group of the remaining *Crella* spp. and *Hemimycale* spp. under NJ (Fig. S1). The representatives of the family Crambeidae (*Monanchora*) appeared as outgroups and the genus *Phorbas* was a sister group of the remaining species. In the BI, NJ, and ML trees, the genera *Hemimycale* and *Crella* appeared polyphyletic, with the Red Sea species *Hemimycale arabica* and *C. cyathophora*, far away from the Atlanto-Mediterranean *Hemimycale* and *Crella* species. The Atlanto-Mediterranean *Crella* formed a well-supported clade (1/81/98, posterior probability/bootstrapping values), which was the sister group of the Atlanto-Mediterranean *Hemimycale* (1/97/98). Moreover, the deep *Hemimycale columella* clustered with an Atlantic sequence downloaded from the GenBank (0.89/89/88) forming a separate clade from the also well-supported (1/97/98) group containing the shallow Mediterranean *Hemimycale*. No genetic differences for this partition were found among shallow individuals. In the BI and ML trees, the two individuals of *Hemimycale arabica* appeared in unresolved positions while they formed a poorly supported (75%) clade in the tree under the NJ criterion (not shown).

28S rRNA (D3–D5) phylogeny

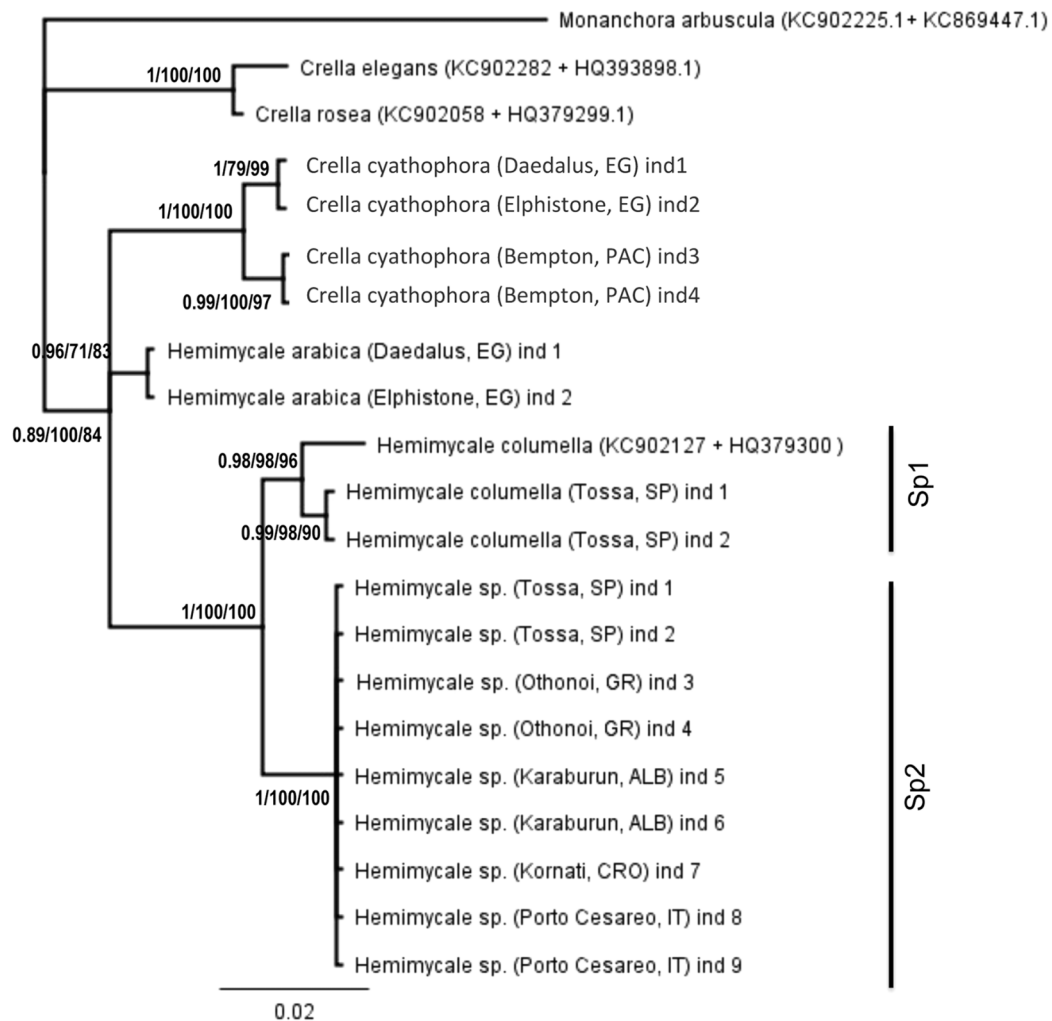
The 28S rRNA (D3–D5) dataset comprised 31 sequences (24 new) of 623 nt. (84 variable positions from which, 60 parsimony informative).

The resulting phylogenies were congruent with the three clustering criteria and matched in most cases the phylogeny based on the 18S rRNA partition, although the supporting values of some clades were in some cases slightly lower (Fig. S2).

The three phylogenies retrieved the representatives of Family Crambeidae (*Monanchora* and *Crambe*) as an outgroup. The monophyly of the in-group containing *Crella* spp. and *Hemimycale* spp. was strongly supported under the BI, NJ, and ML criteria (1/100/100). The genus *Phorbas* was a sister group of the remaining species considered. *Crella* was polyphyletic, with *C. cyathophora* separated from the well-supported clade (1/100/100) encompassing the Atlanto-Mediterranean *Crella*. The latter appeared as a sister clade of a poorly supported group (0.7/77/70) harboring *C. cyathophora* and *Hemimycale* spp. The *Hemimycale* spp. group, although monophyletic, was poorly supported under the NJ and ML criteria (77/70) while the Atlanto-Mediterranean *Hemimycale* clade was well supported under the three clustering criteria (1/92/95).

The deep and shallow Mediterranean populations of *Hemimycale* formed two well-supported monophyletic groups (0.96/87/83 and 0.96/ 100/98, for deep and shallow

18S + 28S



BI/NJ/ML

Figure 1 Phylogenetic tree using concatenated (18S rRNA + COI) partitions. BI, NJ and ML gave the same topologies. Posterior probability, neighbor joining, and maximum likelihood supporting values are at the base of clades.

individuals, respectively), the former containing the Atlantic sequence of *Hemimycale columella*. No genetic differences for this partition were retrieved for shallow individuals despite their spread distribution across the Mediterranean. The individuals of *C. cyathophora* from the Red Sea clustered with those from the Pacific collected between Australia and Nouvelle Calédonie (1/89/76).

COI phylogeny

The COI dataset included 21 sequences (15 new) of 535 nt. (169 variable positions, from which 149 parsimony informative).

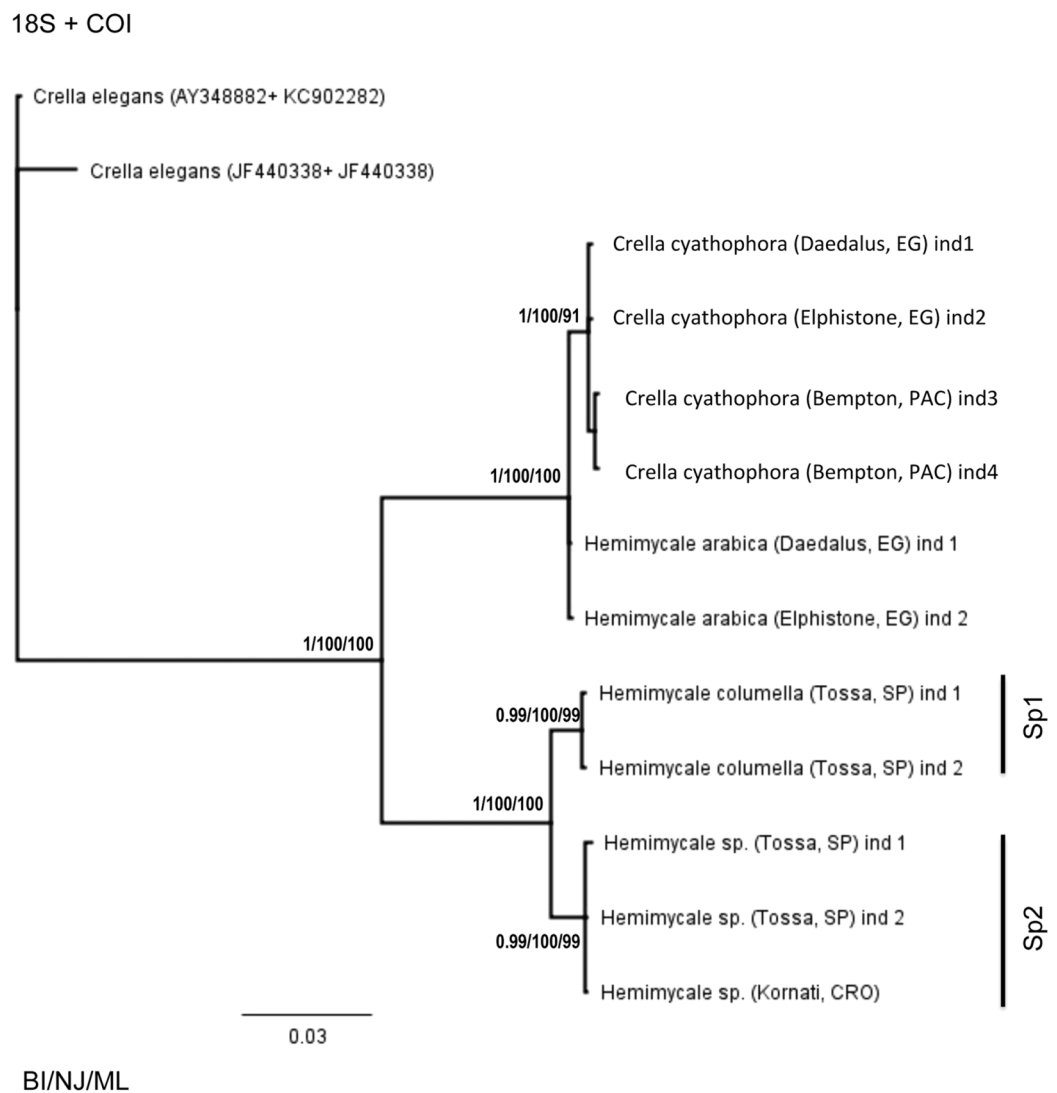


Figure 2 Phylogenetic tree using concatenated (18S + 28S rRNA) partitions. BI, NJ and ML gave the same topologies. Posterior probability, neighbor joining, and maximum likelihood supporting values are at the base of clades.

The COI phylogeny, which was congruent under BI, NJ, and ML, also retrieved the representatives of Crambeidae as outgroups of the group formed by *Crella*, *Phorbas*, and *Hemimyscale*. The genus *Phorbas* clustered with the Atlanto-Mediterranean *Crella* spp. (0.98/100/86) likely because we only included one individual/species of *Phorbas* (Fig. S3).

A clade containing *Hemimyscale* spp. and *C. cyathophora* was well supported (0.94/94/80). The *Hemimyscale* clade was divided into two subclades corresponding to deep and shallow individuals. No genetic differences among shallow individuals were found. A sister, well supported group (1/100/94) contained *C. cyathophora* and *Hemimyscale arabica* representatives with almost no genetic differences between them (Fig. S3).

Concatenated trees

The concatenated 18S + 28S rRNA (Fig. 1) confirmed the outgroup position for the Crambeidae representative (*Monanchora*), the polyphyly of *Crella* with the Red Sea and Pacific species forming a separate clade (1/100/100) from the Atlanto-Mediterranean *Crella*, which appeared in a non-resolved position. *Hemimycale* also appeared polyphyletic, but the position of *Hemimycale arabica* was unresolved. The Atlanto-Mediterranean *Hemimycale* clade was confirmed as well as its division into two subclades: one containing the deep Mediterranean individuals together with two Atlantic sequences of the species and the other one harboring the shallow Mediterranean individuals, which did not show any genetic difference across the Mediterranean and Adriatic Sea.

The concatenated 18S rRNA + COI (Fig. 2) tree contained only 13 sequences and no representative of Crambeidae could be included. The representatives of the Atlanto-Mediterranean *Crella* appeared as outgroups of the remaining target species, which formed two well-supported clades: one containing *C. cyathophora* and *Hemimycale arabica* representatives (1/100/100) and the other with the Atlanto-Mediterranean *Hemimycale* (1/100/100) divided into two monophyletic well-supported groups (deep and shallow individuals).

DISCUSSION

The phylogenetic reconstructions performed with 18S, 28S rRNA and COI, as well as with concatenated genes (18S rRNA + COI and 18S + 28S rRNA) support the polyphyly of *Crella* and *Hemimycale*, under the three clustering criteria used. As although *Hemimycale* was monophyletic with the 28S rRNA (D3–D5) marker, the clade was not statistically supported.

Crella cyathophora sequences differ from those of the Atlanto-Mediterranean *Crella* spp. in 2% (18S rRNA), 2.19% (28S rRNA), and 10.24% (COI). These genetic distances suggest that, despite some spicule similitude (presence of acanthoxeas and smooth diactines with Atlanto-Mediterranean *Crella* spp.), the former species belongs in a different genus, closer to *Hemimycale arabica* (0.71% with 18S rRNA, 1.37% with 28S rRNA, and none with COI) than to the Atlanto-Mediterranean *Crella* spp.

Hemimycale arabica differs from the Atlanto-Mediterranean *Hemimycale* spp. in 1.43–1.86% with 18S rRNA, 1.78–2.19 with 28S rRNA, and in 8.38–8.64% with COI. These strong COI differences and the contrasting morphological traits (blue external color, irregular, rim-free, aerolate areas and abundance of true styles in *Hemimycale arabica* vs. orange–pinkish color, circular, rimmed aerolate areas, and slightly asymmetrical any strongyles almost exclusively in *Hemimycale* spp.) also indicate that *Hemimycale arabica* would belong in a different genus, which might also include *C. cyathophora*, as there are not COI differences between these two species.

Moreover, the Atlanto-Mediterranean Crellidae appeared in 18S and 28S rRNA phylogenies as a sister group of the Atlanto-Mediterranean *Hemimycale*, which suggests higher affinities of this genus with Crellidae than with Hymedesmiidae (its current family). However, more complete analyses including additional Crellidae and Hymedesmiidae OUT's are needed to move *Hemimycale* from Hymedesmiidae to Crellidae.

The phylogenetic trees with any of the three gene partitions either separately or concatenated confirm the presence of two cryptic *Hemimycale* species in the Mediterranean within what was considered until now *Hemimycale columella*. The new species that we name *Hemimycale mediterranean* sp. nov. (see description below) has a shallower distribution across the whole Mediterranean than *Hemimycale columella*, which has Atlantic affinities. *Hemimycale columella* differs from *Hemimycale mediterranea* in 0.85% (18S rRNA), 1.23% (28S rRNA), and in 1–1.2% (COI).

The lack of genetic diversity among the distant populations of *Hemimycale mediterranea* analyzed points to its recent presence in the Mediterranean, which is compatible with a recent introduction. However, the new species has not been recorded out of the Mediterranean, and thus, its origin cannot be established at the present time.

Many cryptic species that were revealed by molecular markers have never been formally described owing to the difficulty of finding diagnostic phenotypic characters. Although after exhaustive observation, only slight, morphological differences have been found to differentiate *Hemimycale mediterranea* sp. nov. from *Hemimycale columella* (see species description below), these phenotypic differences are consistent across individuals and thus, add to molecular differences and biological traits (L. Garate et al., 2013–2014, unpublished data) to consistently differentiate these two species.

Species description

Genus *Hemimycale* Burton, 1934

Sequence accession Numbers GenBank ([Table 1](#))

Type species *Hemimycale columella* (Bowerbank, 1874)

Hemimycale is the only genus of Hymedesmiidae that has smooth diactines and monactines exclusively ([Van Soest, 2002](#)). The genus was described by Burton (1934) as “reduced Mycaleae with skeleton of loose fibers of styli, sometimes modified into anisostrongyles, running vertically to the surface; fibers tending to branch and anastomose; no special dermal skeleton, no microscleres.”

The spicule complement described by Burton; however, seems different from that reported in the several modern redescrptions of *Hemimycale columella* ([Vacelet, Donadey & Froget, 1987](#)), which report predominant straight anisostrongyles with rare or absent styles. Indeed, Burton stated that the Bowerbank representation of *Hemimycale columella* spicules was wrong because it figured anisostrongyles instead of styles, and was precisely the dominance of styles what induced Burton to place the species among the *Mycaleae*. The termination of the diactines either round or pointed ends may be the result of different silica concentration in the water masses, as reported for other siliceous sponge skeletons ([Uriz, 2006](#)), but it cannot be totally discarded that the Burton *Hemimycale columella* belonged in another *Hemimycale* species.

Species: *Hemimycale columella* (Bowerbank, 1874)

Sequence accession numbers GenBank ([Table 1](#))

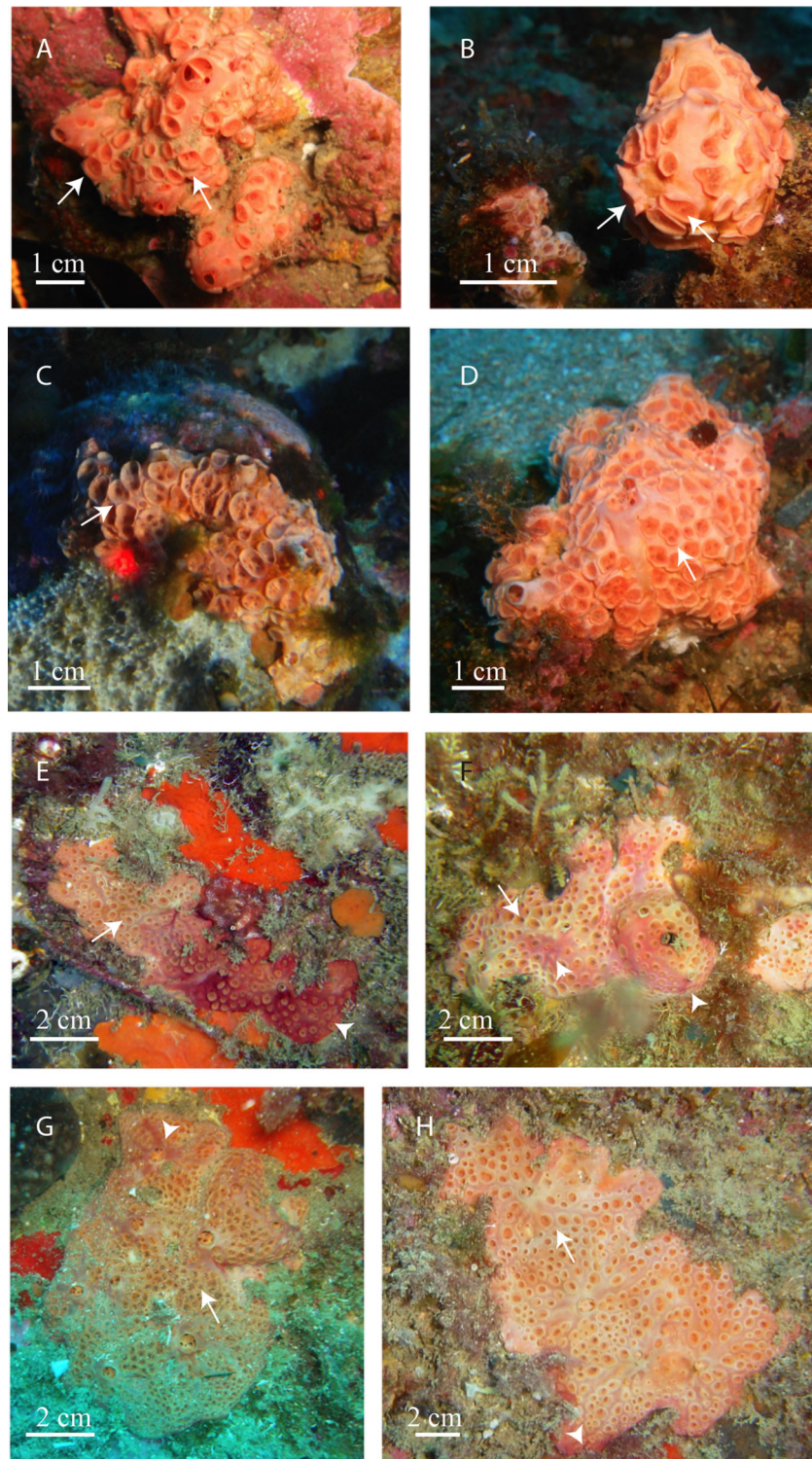


Figure 3 In situ pictures of Atlanto-Mediterranean *Hemimycale* spp. (A, B, C, D) *Hemimycale columella* from 35 to 40 m of depth. (E, F, G, H) *Hemimycale mediterranea* sp. nov. from 12 to 17 m of depth. Whitish tinge is due to calcibacteria accumulation. Red tinges are due to several species of epibiotic cyanobacteria. Arrows point to aerolate inhaling areas; arrowheads indicate the epibiont cyanophyceae on *Hemimycale mediterranea* specimens.

Table 3 Locality and spicule sizes of the studied individuals, and comparison with descriptions by other authors.

Species	Author	Locality	Depth (m)/ Assemblage	Styles	Strongyles (range/mean)	Acanthoxeas
<i>Hemimycale arabica</i> ind. 1	This study	Red Sea (Egypt)	14/coral reef	160–189 (179.6) × 7–8 (7.5)	210–290 (273) × 2.8–4.1 (3.6)	–
<i>Hemimycale arabica</i>	Illan et al. 2004	Red Sea (Egypt)		190–250 (218) × 3.5–5 (4.7)	200–290 (266) × 2.5–4 (3.5)	–
<i>H. mediterranea</i> ind. 7	This study	Adriatic (Croatia)	10–15/rocky sub-horizontal	–	233–330 (274.8) × 3–4.6 (4.0)	–
<i>H. mediterranea</i> ind. 11	This study	Adriatic (Italy)	10–15/rocky sub-horizontal	–	251–300 (276.6) × 2.1–4 (3.0)	–
<i>H. mediterranea</i> ind. 5	This study	Adriatic (Albania)	10–15/rocky sub-horizontal	–	274–317 (296.4) × 2.9–4.5 (4.0)–	–
<i>H. mediterranea</i> ind. 10	This study	Central Med. (Italy)	10–15/rocky sub-horizontal	–	229–328 (291.3) × 2.4–5.2 (3.5)	–
<i>H. mediterranea</i> ind. 3	This study	Eastern Med. (Greece)	10–15/rocky sub-horizontal	–	242–340 (272.7) × 2.6–4 (3.2)	–
<i>H. mediterranea</i> ind. 1	This study	NW Med. (Spain)	12–16/rocky wall	–	261–320 (296.3) × 3.1–3.8 (3.5)	–
<i>Hemimycale columella</i> ind. 1	This study	NW Med. (Spain)	27–29/ coralligenous	–	302–435 (370) × 3–4 (3.7)	–
“ <i>Hemimycale columella</i> ”	Vacelet 1987	NW Med. (France)	–	–	225–310 (285) × 2–4 (3)	–
<i>Hemimycale columella</i>	Vacelet 1987	NW Med. (France)	–	–	320–410 (369) × 2.5–3.8 (3.1)	–
“ <i>Hemimycale columella</i> ”	Vacelet 1987	NW Med. (France)	–	–	220–320 (273) × 2–4 (2.7)	–
<i>Hemimycale columella</i>	Vacelet 1987	North Atlantic (France)	–	–	290–465 (394) × 4–7 (5.1)	–
<i>Hemimycale columella</i>	Topsent 1925	North Atlantic (France)	–	–	400 × 6	–
“ <i>Hemimycale columella</i> ”	Foster 1995	North Atlantic (UK)	–	–	330–420 (373) × 5–6 (5.85)	–
<i>Hemimycale columella</i>	Bowerbank 1874	North Atlantic (UK)	–	–	376 × 7	–
<i>Crella cyatophora</i> ind. 3	This study	Indo-Pacific (Bemptom)	18m/coral reef	–	205–308 (263.9) × 2.2–4.3 (3.4)	92–115 (105.4) × 2–2.3 (2)
<i>C. cyatophora</i> ind. 1	This study	Red Sea (Egypt)	12/coral reef	–	227–293 (267.8) × 2.5–3.9 (3.4)	89–120 (109.4) × 1.8–2.5(2.47)

Description (Figs. 3A–3D): Encrusting to massive sponges. Surface smooth, covered with circular inhaling, areas up to 6 mm in diameter with an up to 3 mm high rim. Morbid and fleshy consistence. Translucent to whitish ectosome, difficult to separate from the choanosome. Thousands of calcareous spherules, 1 μm in diameter formed by intracellular calcifying bacteria (*Uriz et al., 2012*) are spread through the sponge mesohyl and specially accumulated at the sponge periphery of whitish individuals (L. Garate et al., 2013–2014, unpublished data).

Color from pinkish-orange to whitish outside, dark orange inside.

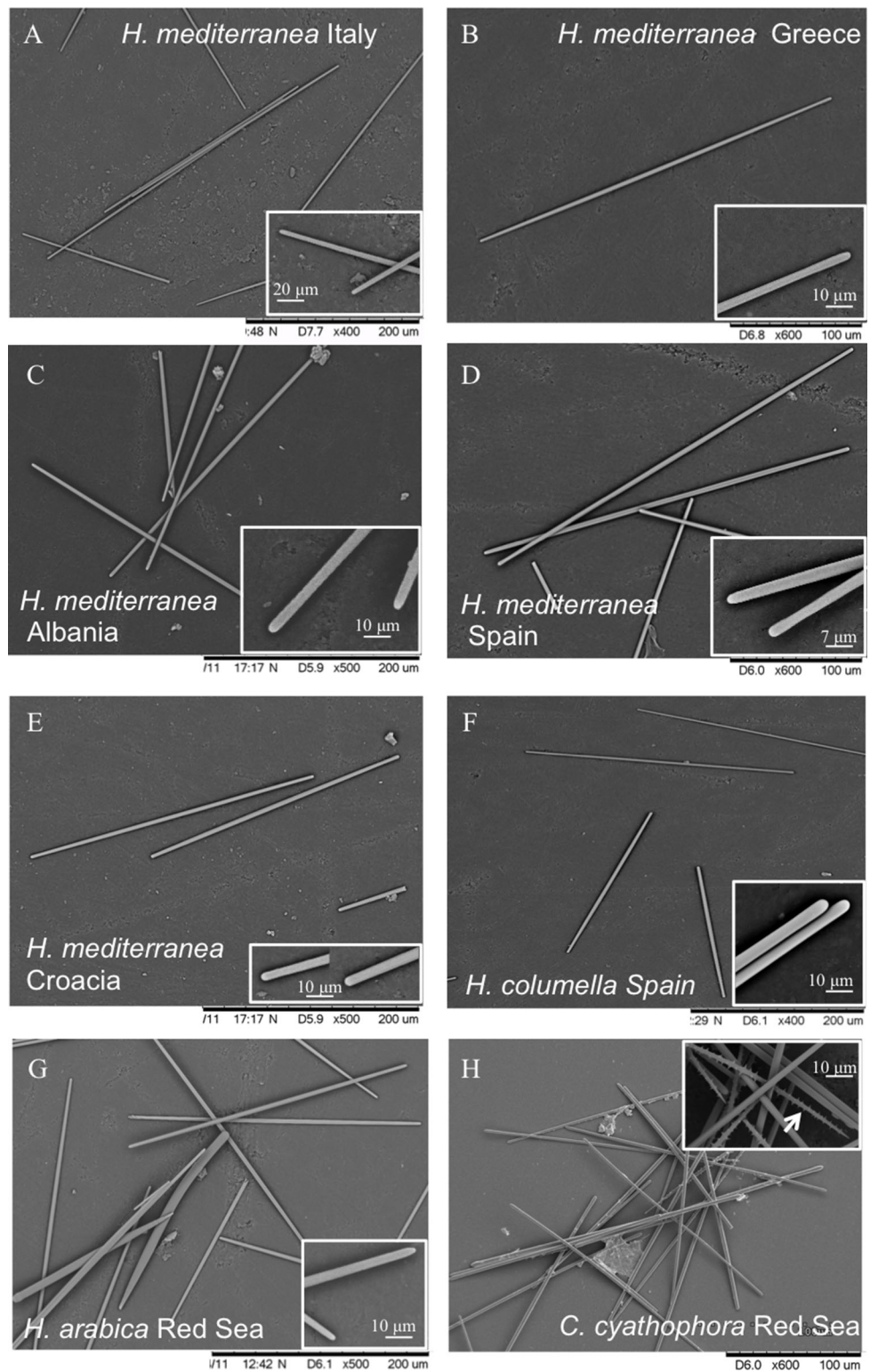


Figure 4 Spicules of *Hemimycale* spp. and *Crella cyathophora* though SEM. (A, B, C, D, E) Anisotrongyles (*Hemimycale mediterranea*). (F) Anisotrongyles (*Hemimycale columella*). (G) Anisotrongyles and one style (*Hemimycale arabica*). (H) Anisotrongyles and acantoxeas (*Crella cyatophora*). Inserts on each picture correspond to magnifications of the spicule ends.

Spicules (Table 3; Fig. 4F): Asymmetric strongyles (anysotrongyles), straight, $302\text{--}435 \times 3\text{--}4 \mu\text{m}$ in size. Styles rare or completely absent from the Mediterranean specimens (this study) and Canary Islands (Cruz, 2002).

Skeletal arrangement: Plumose branching bundles of anysostrongyles together with spread spicules. A palisade of vertical anysotrongyles forms the rim around the inhaling areas.

Distribution: Northeastern Atlantic (United Kingdom and Ireland coasts) Canarias Islands (Cruz, 2002), western Mediterranean: Tossa de Mar, Arenys de Mar, from 28 to 60 m depth (this study). It is not possible to confirm whether previous Mediterranean records of the species (see Vacelet & Donadey, 1977) belonged to *Hemimycale columella* or to *Hemimycale mediterranea*.

Biology: Multiannual life span, ca. 60% survival after two monitoring years; maximum growth in autumn–winter (L. Garate et al., 2013–2014, unpublished data). Larval release occurs at the beginning of November in Mediterranean populations (M. J. Uriz, L. Garate & G. Agell, 2013–2014, unpublished data).

Species: *Hemimycale mediterranea* sp. nov. (Figs. 3E–3H)

Sequence accession numbers GenBank (Table 1)

Description: Thick encrusting sponges with aerolate inhaling areas up to 3 mm in diameter, surrounded by an up to 1.5–2 mm high rim, which in some cases barely surpasses the sponge surface. Thousands of calcareous spherules, $1 \mu\text{m}$ in diameter formed by intracellular calcifying bacteria are spread through the sponge mesohyl and specially accumulated at the sponge periphery (Garate et al., in press).

Ectosome: Firmly attached to the choanosome.

Color: Flesh to clear brownish externally, more or less whitish depending on calcibacteria accumulation at the surface, sometimes partially covered by an epibiotic (reddish or pinkish) cyanobacteria.

Spicules (Table 3; Figs. 4A–4E): Smooth, uniform in size, straight, anysostrongyles, $200\text{--}296 \times 3\text{--}4 \mu\text{m}$ in size. Styles completely absent.

Skeletal arrangement: Plumose undulating bundles of anysostrongyles together with spread spicules. A palisade of vertical anysotrongyles forms the rim around the inhaling areas.

Known distribution: Northwestern Mediterranean, central Mediterranean, Adriatic, eastern Mediterranean (Spain: Cap De Creus, Tossa, Blanes, Arenys, South Italy: Croatia, Tremiti, Turkey, Greece) between 3 and 17 m deep.

Biology: Annual life span, maximum growth rates in summer (M. J. Uriz, L. Garate & G. Agell, 2013–2014, unpublished data). Larval release at the end of September beginning of October (M. J. Uriz, L. Garate & G. Agell, 2013–2014, unpublished data).

In most cases, it is difficult to ascertain whether individuals of *Hemimycale columella* recorded by other authors belong to *Hemimycale columella* or *Hemimycale mediterranea*. The redescription of *Hemimycale columella* by Van Soest (2002) based on the holotype

(from the Atlantic) reported large aerolate porefields with elevated rims, which are shared with the deep Mediterranean specimens of *Hemimycale columella* (Figs. 3A–3D) in contrast to the small, short-rimmed porefields showed by *Hemimycale mediterranea* sp. nov. Both species have mainly straight slightly asymmetric strongyles but the spicule sizes are systematically larger in *Hemimycale columella* (Table 3). However, while styles were rarely present in *Hemimycale columella* individuals, they have not been found in specimens of *Hemimycale mediterranea* sp. nov. The external color also differs between the two species, being orange to pinkish in *Hemimycale columella* and flesh color to brownish *Hemimycale mediterranea* sp. nov. (Figs. 3E–3H). Vacelet & Donadey (1977) reported two different color forms occurring side by side on the littoral of Provence (France), one pink cream and the other one brownish. Likely the second color morph, which besides had smaller strongyles, corresponded to the *Hemimycale mediterranea* sp. nov.

Color has not received much attention as a diagnostic character in sponges because it has been generally considered to be a response to higher or lower light irradiance at the sponge habitat, or to the presence of epibiotic or symbiotic cyanobacteria. However, color has proven to be taxonomically relevant to distinguish other invertebrates such as shrimp species (Knowlton & Mills, 1992) and also sponge species of the genus *Scopalina* (Blanquer & Uriz, 2008), and thus it seems worthy to be taken into account in sponge taxonomy.

The slight phenotypic differences found between the two species appear; however, consistent across individuals and localities within the Atlanto-Mediterranean basin. Moreover, their ecological distribution and bacterial symbionts, strongly differentiate these two cryptic species. For instance, although calcareous spherules produced by intracellular bacteria are present in the two species, the producer bacteria belong in different species (Garate et al., in press), and the respective microbial communities totally differ (Garate et al., in press). Symbionts, as predators do (e.g., Wulff, 2006), often distinguish their target sponge preys or hosts while the species remain morphologically cryptic to taxonomists. Moreover, *Hemimycale mediterranea* sp. nov. shows an annual life span, with individuals disappearing after larval release, while *Hemimycale columella* has a multiannual life span (M. J. Uriz, L. Garate & G. Agell, 2013–2014, unpublished data) and growth dynamics also differs between the two species, as *Hemimycale mediterranea* sp. nov. grows more in summer, while *Hemimycale columella* grows preferentially in autumn–winter (M. J. Uriz, L. Garate & G. Agell, 2013–2014, unpublished data).

The contrasting ecological distribution of these two cryptic species in the Mediterranean helps to predict their identity in the field. *Hemimycale mediterranea* sp. nov. inhabits shallower zones than *Hemimycale columella*. However, it is likely that both species may share occasionally habitats, as the record of the two color morphs side by side (Vacelet & Donadey, 1977) indicate. *Hemimycale mediterranea* sp. nov. seems to be more abundant and widespread in the Mediterranean than *Hemimycale columella*. Molecular differences between groups of individuals of *Hemimycale columella* suggest the possible presence of additional cryptic species among the deep Mediterranean *Hemimycale*.

The presence of two morphologically cryptic *Hemimyscale* species in the Mediterranean, which show contrasting biological traits, reinforces the idea that cryptic species represent something more than wrong taxonomic identifications or biodiversity underestimates. They may feature contrasting biological cycles and life spans, and puzzle biological studies, which may invalidate conservation policies based on those studies.

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ADDITIONAL INFORMATION AND DECLARATIONS

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Competing Interests

The authors declare that they have no competing interests.

Author Contributions

- Maria J. Uriz conceived and designed the experiments, analyzed the data, contributed reagents/materials/analysis tools, wrote the paper, and reviewed drafts of the paper.
- Leire Garate conceived and designed the experiments, performed the experiments, analyzed the data, wrote the paper, prepared figures and/or tables, and reviewed drafts of the paper.
- Gemma Agell performed the experiments, analyzed the data.

DNA Deposition

The following information was supplied regarding the deposition of DNA sequences:

The 18 S, 28 S and COI sequences described here are accessible at GenBank
<https://www.ncbi.nlm.nih.gov/genbank/>.

Accession numbers: [KY002124](#), [KY002171](#), [KY002181](#), [KY002125](#), [KY002172](#), [KY002182](#), [KY002183](#), [KY002184](#), [KY002126](#).

COI: [KY002127](#).

18S: [KY002160](#), [KY002185](#), [KY002128](#), [KY002161](#), [KY002186](#), [KY002129](#), [KY002187](#), [KY002188](#), [KY002130](#), [KY002162](#), [KY002189](#), [KY002163](#), [KY002190](#), [KY002131](#), [KY002132](#), [KY002134](#), [KY002170](#), [KY002193](#), [KY002194](#), [KY002133](#), [KY002199](#), [KY002164](#), [KY002165](#), [KY002197](#), [KY002198](#), [KY002166](#), [KY002191](#), [KY002167](#), [KY002192](#), [KY002168](#), [KY002195](#), [KY002169](#), [KY002196](#), [KY002120](#), [KY002173](#), [KY002177](#), [KY002121](#), [KY002174](#), [KY002178](#), [KY002122](#), [KY002175](#), [KY002179](#), [KY002123](#), [KY002176](#), [KY002180](#).

Data Deposition

The following information was supplied regarding data availability:

CEAB Porifera Collection.

Vouchers: CEAB.POR.GEN.001, CEAB.POR.GEN.002, CEAB.POR.GEN.003, CEAB.POR.GEN.004, CEAB.POR.GEN.005, CEAB.POR.GEN.006, CEAB.POR.GEN.007, CEAB.POR.GEN.008, CEAB.POR.GEN.009, CEAB.POR.GEN.010, CEAB.POR.GEN.011, CEAB.POR.GEN.012, CEAB.POR.GEN.013, CEAB.POR.GEN.014, CEAB.POR.GEN.015, CEAB.POR.GEN.016, CEAB.POR.GEN.017, CEAB.POR.GEN.018, CEAB.POR.GEN.019, CEAB.POR.GEN.020, CEAB.POR.GEN.021, CEAB.POR.GEN.022, CEAB.POR.GEN.023, CEAB.POR.GEN.024, CEAB.POR.GEN.025, CEAB.POR.GEN.026, CEAB.POR.GEN.027, CEAB.POR.GEN.028, CEAB.POR.GEN.029.

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Species name: LSID: <urn:lsid:zoobank.org:act:A90CB361-6EC5-4B94-805B-F5A4C0EDF2F9>.

Supplemental Information

Supplemental information for this article can be found online at <http://dx.doi.org/10.7717/peerj.2958#supplemental-information>.

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