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










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Research Article



Integrative taxonomy of *Tethya*: description of four new species based on morphology, phylogeny and microbiome diversity

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The genus *Tethya*, one of the most iconic groups in the phylum Porifera, includes nearly 100 valid species. *Tethya* shows a nearly cosmopolitan distribution, and thus this clade could help elucidate global mechanisms of speciation in sponges. Contrasting with many other marine organisms, *Tethya* peaks in diversity in the temperate region, with only ~30% of diversity occurring in the tropics. This pattern may however be related to a lack of studies in the tropics, masked by dubious taxonomic identifications. To address this question, we studied new collections from the Western Atlantic (Caribbean and Brazil) and the Northeastern Atlantic, as well as museum material from the Indian Ocean. Combining morphological investigation with molecular phylogenetics and the study of the sponge's microbial communities, we conclude that four constitute new species that we describe here: *Tethya martini* Riesgo, Giribet, & Santodomingo, sp. nov.; *Tethya simoni* Santodomingo, Zea, & Riesgo, sp. nov.; *Tethya erici* Díez-Vives, Santodomingo, Moles, & Riesgo, sp. nov.; *Tethya orioni* Kenny, Santodomingo & Riesgo, sp. nov. Some species thought to be widespread (e.g., *T. aurantium* and *T. seychellensis*), each represent multiple species with unique geographic distributions. A phylogenetic analysis of *Tethya* (based on COI and 28S rRNA sequence data) recovered five main clades, which were also characterized by distinct prokaryotic communities. This suggests that microbiomes could be used as a guide for taxonomic identification in *Tethya*. We finally explored the existence of a phyllosymbiotic pattern in sponges at different levels of prokaryotic taxonomic resolution (i.e., at phylum, class, and genus-level analysis of prokaryotes). Remarkably, our analysis revealed high levels of coevolution of *Tethya* and their associated microbial communities, even when microbiomes were analysed at the genus level. Our findings support the use of an integrative approach to better understand the evolutionary history of sponges.

<http://zoobank.org/urn:lsid:zoobank.org:pub:54F80E5E-99A4-43CE-BA3A-BCD01E715595>

Key words: systematics, microbiome, spicules, sponges, phylogeny

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Introduction

Sponges in the genus *Tethya* Lamarck, 1815 are iconic taxa among the phylum Porifera (Sarà, 2002) for being the only metazoans with spherical symmetry (Brusca *et al.*, 2023). Their spherical shape, well-defined cortex, and surface usually covered by tubercles and buds, are characteristics that give *Tethya* the common name of ‘golf balls’, and make this genus one of the most easily recognized groups of species among demosponges (Sarà, 1987). There have been 96 accepted species described worldwide since the original designation of the genus name *Tethya* by Lamarck in 1815 (de Voogd *et al.*, 2024). They have been noted to occur from shallow coastal waters to the deep sea, in all oceans except the poles (Bergquist & Kelly-Borges, 1991; Sarà, 2002).

Some of the first species described for the genus, *Tethya aurantium* (Pallas, 1766), *T. seychellensis* (Wright, 1881), *T. diploderma* Schmidt, 1870, and *T. japonica* Sollas, 1888, were long thought to be cosmopolitan or having a broad distribution range (Sarà, 1987). However, some records remained dubious and were later confirmed to belong to undescribed species (Bergquist & Kelly-Borges, 1991; Ribeiro & Muricy, 2004, 2011). Regional inventories conducted over the past 30 years have uncovered a hidden diversity of *Tethya*, with species described from Australia and New Zealand (Bergquist & Kelly-Borges, 1991; Sarà & Sarà, 2004), Brazil (Ribeiro & Muricy, 2004, 2011), the Mexican Pacific and California (Austin *et al.*, 2014; Sarà Gómez, & Sarà, 2001), as well as Galápagos (Desqueyroux-Fáunderz & van Soest, 1997; Sarà *et al.*, 2000; Sim-Smith *et al.*, 2021). The remaining described *Tethya* species are thought to be widely distributed across the Mediterranean Sea, the North Atlantic Ocean, North Pacific Ocean, with a few species described in the Chilean fjords, the Caribbean Sea, and the Indian Ocean. From all this diversity, over 60% of the *Tethya* species occur in temperate waters, about 30% are from the tropics, and 10% have a widespread distribution in both temperate and tropical waters. Although the current *Tethya* distribution could be a result of an underlying biogeographic pattern, it could also be attributed to a collection bias, as in other poriferan groups (van Soest *et al.*, 2012) and lack of studies in the tropical region, in particular the central Indo-Pacific and the Caribbean. To date, only six species are known from the Caribbean: the imprecise type locality of *T. diploderma* and *T. globum* Duchassaing de Fonbressin & Michelotti, 1864 is Antilles; *Tethya actinia* de Laubenfels & Hindle, 1950 was first described from Bermuda; and *T. maza* Selenka, 1879 was described from southern Brazil. The other two Caribbean species, *T. aurantium* and *T. seychellensis*, have type localities in the Mediterranean Sea

and the Seychelles, respectively. Previous records of *T. aurantium* from California have been reclassified into two new species: *T. californiana* de Laubenfels, 1932 and *T. vacua* Austin *et al.*, 2014. Similarly, former *T. aurantium* specimens from Brazil have been identified as *T. beatrizae* Ribeiro & Muricy, 2011. In consequence, the occurrence of *T. aurantium* in the Caribbean remains uncertain.

The taxonomic work led by Sarà and collaborators on *Tethya* has resulted in the description of 34 species, which comprise over a third of the current valid species (de Voogd *et al.*, 2024). According to Sarà (1994, 2002), the diagnostic characters that distinguish *Tethya* from the other 13 genera in the family Tethyidae include their spherical or hemispherical body, with a well-developed cortex differentiated from the choanosome, and their main skeleton formed by strongyloxea bundles radiating from the centre of the sponge. The spicule complement of the genus is greatly conserved, containing megascleres (mostly strongyloxeas) and euasters (both megasters and micrasters) (Ribeiro & Muricy, 2004; Sarà, 2002; Sarà & Sarà, 2002). Because the terminology for the megasters and micrasters has evolved over the years, Ribeiro and Muricy (2011) re-described the different morphologies and unified the euaster terminology to make comparisons across species from Brazil more detailed and less confusing. This terminology was adopted in later work for a new species from Brazil (Mácola & Menegola, 2018), but it has not been used in subsequent descriptions of new *Tethya* species from other regions (Austin *et al.*, 2014; Hajdu *et al.*, 2013; Sorokin *et al.*, 2019), where the authors still used the classic terms (Boury-Esnault & Rützler, 1997; Sarà, 1994, 2002).

Pioneering work on the evolutionary framework of *Tethya* was also developed by Sarà (1987), who compiled the main character variation within the genus including morphological, ecological, cytological, biochemical, reproductive, and biogeographic traits. This study exposed some of the main gaps of information and envisaged the presence of cryptic species within the widely distributed species *T. aurantium*, *T. seychellensis*, and *T. japonica* Sollas, 1888 (Sarà, 1987). Bergquist and Kelly-Borges (1991) built the first phylogeny of *Tethya* based on morphological characters, mainly with Australian and New Zealand species. Their work highlighted the need to use standard terminology and revealed that well-defined characters such as the arrangement of megascleres, spheraster shape, colour, and the regularity or deformity of micrasters, among others, had high consistency index values, hence being highly informative in phylogenetic analyses. The exercise was repeated by Sarà and Sarà (2004) with a larger

number of species also from the Austral region and showed a correspondent arrangement of species groups within *Tethya*. A similar cladistic approach was used for the reconstruction of a family-level phylogeny, suggesting that the speciose genus *Tethya* diverged early in the evolution of the Tethyidae with successive branching of monospecific or less speciose genera in the family (Sarà & Burlando, 1994).

Subsequent attempts to reconstruct the phylogeny of *Tethya* included the combination of a single molecular marker (*cytochrome c oxidase subunit I*, COI) and morphological characters (Heim et al., 2007), and with few exceptions, the groups recovered by the molecular analyses correspond to those found using morphological characters. Since then, some of the new *Tethya* species have been described including molecular data and, therefore, subsequent reexaminations of the phylogeny have been conducted (Heim & Nickel, 2010; Sorokin et al., 2019). In the most recent study, Sorokin et al. (2019) discussed the possibility of using external colour as a diagnostic character for some clades, and, given the lack of other synapomorphies, they proposed that the exploration of chemical compounds, specialized cells, and associated microbes could provide additional insights on the phylogenetic relationships of the different lineages in the family.

The sponge microbiome is well-known for being species-specific (Thomas et al., 2016, Yang et al., 2019) and even genotype-specific (Griffiths et al., 2019; Díez-Vives et al., 2020; Easson et al., 2020), and also showing strong functional convergence of the microbiome roles (Fan et al., 2012). Two ecological sponge types were described based on their microbial abundance, diversity, and pumping rate (Vacelet & Donadey, 1977). One type, termed ‘Low Microbial Abundance’ (LMA) sponges, has a microbial concentration close to that of seawater, and these sponges rely on heterotrophic feeding on particulate organic matter. The second type, ‘High Microbial Abundance’ (HMA) sponges, can contain microbial communities occupying up to 80% of their tissues in some species, and these sponges rely on this microbiome to acquire nutrients (Hentschel et al., 2006; Weisz et al., 2007; Pankey et al., 2022). While the LMA sponges harbour microbiota of very diverse phylogenetic signatures (Sipkema et al., 2015, Thomas et al., 2016), the HMA sponges are dominated by a few microbial phyla (Moitinho-Silva, Nielsen et al., 2017; Pita et al., 2018). *Tethya* species are consistently considered to be LMA, either by using predictive methodologies (Moitinho-Silva, Steinert et al., 2017; Pankey et al., 2022), by 16S rRNA amplicon sequencing (Thiel et al., 2007; Waterworth et al., 2017), or by direct ultrastructural observation of tissues (Gaino et al., 2006; Gaino &

Sarà, 1994). Microbiomes can be vertically transmitted to the offspring through oocytes, which secure necessary symbionts during the settlement phase, or horizontally acquired from the environment by filtration, or a combination of both (Schmitt et al., 2008; Vrijenhoek, 2010; Ebert, 2013; de Oliveira et al., 2020). Given that several studies have reported vertical transmission of symbionts in Mediterranean species of *Tethya* (Gaino et al., 1987; Gaino & Sarà, 1994; Sciscioli et al., 2002), and following up on the idea by Sorokin et al. (2019), we explored how the microbiome signatures can potentially serve as an additional feature to aid in the systematics of sponges, in the genus *Tethya* in particular.

Here, we describe the morphological features of four new species of the genus *Tethya* from the Caribbean, North and South Atlantic, and the Persian Gulf, and place them in a molecular phylogenetic framework. In addition, we analyse the microbiome of the new species and additional sympatric *Tethya* species to understand phyllosymbiotic patterns in this cosmopolitan sponge genus.

Materials and methods

Several specimens of two morphospecies (preliminarily identified as *Tethya*) were collected on coral reefs in Isla Cristóbal, Bocas del Toro, Panama (9.21685, -82.21370) in March 2010 by scuba diving (Table 1) and three more in the intertidal area of Pituba Beach, Salvador de Bahia, Brazil (-13.0066632, -38.4547663) in January 2018. Three specimens of an undescribed *Tethya* species were also collected in a rocky shore in the ‘corrales’ of Chipiona, Cádiz, Spain (36.74168, -6.43887) and subtidal pools in La Palma, Canary Islands (28.693245, -17.759086) by snorkelling, in October 2014 and September 2022, respectively. A specimen collected in 1998 in the Persian Gulf, Ras Ghumeis, United Arab Emirates (24.380880, 51.576362) and deposited at The Natural History Museum, London, UK (NHMUK) under voucher number 2000.9.14.16 was also examined (Table 1). All newly collected specimens were fixed in 96% ethanol soon after collection. To compare the morphological and molecular features of the new species, several other *Tethya* specimens, including type material from the collections of the NHMUK, were examined (Table 1).

Spicules were prepared by dissociation in sodium hypochlorite, followed by two washes with water, and one with 96% ethanol. Microscope slides of the spicules were prepared and permanently mounted with DPX (distyrene, plasticizer, and xylene) medium. Thick sections for most species were prepared using tissue embedded in paraffin, sectioned with a microtome at 10 µm,

Table 1. Metadata for all specimens considered in the phylogenetic analyses, including species name (and label in the tree), collection localities, voucher and accession numbers for both COI and 28S, and whether microbiome sequencing was included.

Species	Name in trees	Locality	Voucher/Sample name	COI Accession number	28S Accession number	16S amplicon Accession number
<i>Adreus fascicularis</i>	<i>Adreus fascicularis</i> HQ379428 or HQ379314	English Channel, UK	Mc4559	HQ379428	HQ379314	NO
<i>Adreus micraster</i>	<i>Adreus micraster</i> AY626306	Mauritania	POR06837		AY626306	NO
<i>Adreus</i> sp.	<i>Adreus</i> sp. HQ379377	Ireland	BELUM:Mc4982		HQ379377	NO
<i>Axos cliffoni</i>	<i>Axos cliffoni</i> KY947263	Vietnam: Quang Tri	MT5.2015	KY947263		NO
<i>Axos cliffoni</i>	<i>Axos cliffoni</i> AY626308	Australia	G300111		AY626308	NO
<i>Axos cliffoni</i>	<i>Axos cliffoni</i> HQ379380	Australia	BELUM:POR06166		HQ379380	NO
<i>Axos flabelliformis</i>	<i>Axos flabelliformis</i> KC869578	Australia	NCI089		KC869578	NO
<i>Cliona celata</i>	<i>Cliona celata</i> KY492595	Portugal	M23F	KY492595		NO
<i>Jaspis</i> sp.	<i>Jaspis</i> sp. AY561977	Papua New Guinea	UCMPWC1021	AY561977		NO
<i>Adreus</i> sp.	<i>Adreus</i> sp. AY561901	Australia	G315767		AY561901	NO
<i>Laxothelya dampierensis</i>	<i>Laxothelya dampierensis</i> AY561905	Australia	WAMZ11871		AY561905	NO
<i>Liosina blastifera</i>	<i>Liosina blastifera</i> KX866762	Israel: Ma'agan-Mikha'el	Po.25551	KX866762		NO
<i>Liosina paradoxa</i>	<i>Liosina paradoxa</i> AF437303 or AY319318	Sulawesi/Indonesia		AF437303	AY319318	NO
<i>Paratimea hoffmanae</i>	<i>Paratimea hoffmanae</i> KC869429	Norway: Røst reef	ZMBN 125735	KC869429		NO
<i>Placospongia intermedia</i>	<i>Placospongia intermedia</i> JX999089	Panama: Bocas del Toro	MCZ:DNA106621	JX999089		NO
<i>Placospongia</i> sp.	<i>Placospongia</i> sp. KC869430	Panama: Bocas del Toro	PC-BT-18	KC869430		NO
<i>Stellitethya ingens</i>	<i>Stellitethya ingens</i> AY561899	Australia	WAMZ11947		AY561899	NO
<i>Tectitethya keyensis</i>	<i>Tectitethya keyensis</i> KC869616	USA	NCI360		KC869616	NO
<i>Tethya actinia</i>	<i>Tethya actinia</i> AY320033	USA: Zane Grey Creek, Long Key, Florida		AY320033		NO
<i>Tethya actinia</i>	<i>Tethya actinia</i> i1	Panama: Bocas del Toro	MCZ:IZ:135213_2	PP594403		PRJNA970968
<i>Tethya actinia</i>	<i>Tethya actinia</i> i2	Panama: Bocas del Toro	MCZ:IZ:135213_3	PP594404		PRJNA970968
<i>Tethya actinia</i>	<i>Tethya actinia</i> i3	Panama: Bocas del Toro	MCZ:IZ:135213_1	NO		PRJNA970968
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> JX999070	Italy: Bari	MCZ:ICZ:DNA106628	JX999070		NO
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> EF558571	Croatia		EF558571		NO
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> EF093531	Croatia		EF093531		NO

(Continued)

Table 1. Continued.

Species	Name in trees	Locality	Voucher/Sample name	COI Accession number	28S Accession number	16S amplicon Accession number
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> EF093529	Croatia		EF093529		NO
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> EF584565	Elba, Italy		EF584565		NO
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> i6	Italy: Naples	MCZ:ICZ:106628	QQ948196	QQ947274	PRJNA970968
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> i7	Italy: Naples		QQ948197	QQ947273	PRJNA970968
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> i8	Italy: Naples				PRJNA970968
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> i9	Italy: Naples		QQ948198		PRJNA970968
<i>Tethya aurantium</i>	<i>Tethya aurantium</i> i10	Italy: Naples				PRJNA970968
<i>Tethya bergquistae</i>	<i>Tethya bergquistae</i>	New Zealand: Wellington		QQ948203		NO
<i>Tethya burtoni</i>	<i>Tethya burtoni</i>	New Zealand: Wellington		QQ948210		NO
<i>Tethya californiana</i>	<i>Tethya californiana</i> AY561978 or AY561900	USA: Carmel, California		AY561978	AY561900	NO
<i>Tethya californiana</i>	<i>Tethya californiana</i> KJ620403	South Australia		KJ620403		NO
<i>Tethya californiana</i>	<i>Tethya californiana</i> GQ292532	Canada: Vancouver Island	BCA1	GQ292532		NO
<i>Tethya californiana</i>	<i>Tethya californiana</i> GQ292533	Canada: Vancouver Island	BCA2	GQ292533		NO
<i>Tethya cf. aurantium</i>	<i>Tethya cf. aurantium</i> AY552024	France: Marseille			AY552024	NO
<i>Tethya citrina</i>	<i>Tethya citrina</i> EF558570	UK: Northern Ireland	MC3217	EF558570		NO
<i>Tethya citrina</i>	<i>Tethya citrina</i> HQ379427 or HQ379237	UK: Wales	Mc5113	HQ379427	HQ379237	NO
<i>Tethya citrina</i>	<i>Tethya citrina</i> EF558569	Croatia	SPD279	EF558569		NO
<i>Tethya citrina</i>	<i>Tethya citrina</i> i4	Italy: Naples	US113	QQ948205	QQ947272	NO
<i>Tethya citrina</i>	<i>Tethya citrina</i> i5	France: Roscoff	Roscoff 2	QQ947284	QQ947284	PRJNA970968
<i>Tethya citrina</i>	<i>Tethya citrina</i> i6	France: Roscoff	Roscoff 10	QQ947285	QQ947285	PRJNA970968
<i>Tethya citrina</i>	<i>Tethya citrina</i> i7	Spain: Ría de Vigo, Galicia	Gali1	QQ948206		PRJNA970968
<i>Tethya citrina</i>	<i>Tethya citrina</i> i8	Spain: Ría de Vigo, Galicia	Gali2			PRJNA970968
<i>Tethya citrina</i>	<i>Tethya citrina</i> i9	Spain: La Palma, Canary Islands	MNCN 1.01/1041	PP587641		PRJNA970968
<i>Tethya citrina</i>	<i>Tethya citrina</i> i10	Spain: La Palma, Canary Islands	MNCN 1.01/1042	PP587642		NO
<i>Tethya citrina</i>	<i>Tethya citrina</i> i11	Spain: La Palma, Canary Islands	MNCN 1.01/1043	PP587638		PRJNA970968

(Continued)

Table 1. Continued.

Species	Name in trees	Locality	Voucher/Sample name	COI Accession number	28S Accession number	16S amplicon Accession number
<i>Tethya citrina</i>	<i>Tethya citrina</i> i12	Spain: La Palma, Canary Islands	MNCN 1.01/1044	PP587640		PRJNA970968
<i>Tethya citrina</i>	<i>Tethya citrina</i> i13	Spain: La Palma, Canary Islands	MNCN 1.01/1045	PP587639		PRJNA970968
<i>Tethya coccinea</i>	<i>Tethya coccinea</i> JQ034584	Australia: Queensland	G303083	JQ034584		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i1	Spain: Chipiona	MNCN 1.01/1023	OQ948204	OQ947278	PRJNA970968
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i2	Spain: Chipiona	MNCN 1.01/1024			PRJNA970968
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i3	Spain: Chipiona	MNCN 1.01/1025			PRJNA970968
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i4	Spain: La Palma, Canary Islands	MNCN 1.01/1027	PP587635		PRJNA970968
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i5	Spain: La Palma, Canary Islands	MNCN 1.01/1028	PP587627		PRJNA970968
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i6	Spain: La Palma, Canary Islands	MNCN 1.01/1029	PP587633		PRJNA970968
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i7	Spain: La Palma, Canary Islands	MNCN 1.01/1030	PP587634		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i8	Spain: La Palma, Canary Islands	MNCN 1.01/1031	PP587636		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i9	Spain: La Palma, Canary Islands	MNCN 1.01/1032	PP587624		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i10	Spain: La Palma, Canary Islands	MNCN 1.01/1033	PP587626		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i11	Spain: La Palma, Canary Islands	MNCN 1.01/1034	PP587628		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i12	Spain: La Palma, Canary Islands	MNCN 1.01/1035	PP587625		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i13	Spain: La Palma, Canary Islands	MNCN 1.01/1036	PP587629		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i14	Spain: La Palma, Canary Islands	MNCN 1.01/1037	PP587630		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i15	Spain: La Palma, Canary Islands	MNCN 1.01/1038	PP587631		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i16	Spain: La Palma, Canary Islands	MNCN 1.01/1039	PP587632		NO
<i>Tethya erici</i> sp. nov.	<i>Tethya erici</i> sp. nov. i17	Spain: La Palma, Canary Islands	MNCN 1.01/1040	PP587637		NO
<i>Tethya gracilis</i>	<i>Tethya gracilis</i> EF584568	Germany: Aquazoo Loebekke-Museum, Düsseldorf		EF584568		NO
<i>Tethya gracilis</i>	<i>Tethya gracilis</i> i1	Brazil: Salvador de Bahia	UFBAPOP 5021	OQ948213		PRJNA970968
<i>Tethya gracilis</i> Chengue	<i>Tethya gracilis</i> i2	Colombia: Chengue Bay, Colombia	INV-PSM492	OQ948212	OQ947276	PRJNA970968
<i>Tethya hibernica</i>	<i>Tethya hibernica</i> EF558566	UK: Northern Ireland: Rathlin Island	Mc2426	EF558566		NO

(Continued)

Table 1. Continued.

Species	Name in trees	Locality	Voucher/Sample name	COI Accession number	28S Accession number	16S amplicon Accession number
<i>Tethya hibernica</i>	<i>Tethya hibernica</i> EF558567	UK: Northern Ireland: Rathlin Island	Mc3037	EF558567		NO
<i>Tethya hibernica</i>	<i>Tethya hibernica</i> EF558568	UK: Northern Ireland: Rathlin Island	Mc2486	EF558568		NO
<i>Tethya hibernica</i>	<i>Tethya hibernica</i> HQ379238	UK: Northern Ireland: Rathlin Island	Mc4061		HQ379238	NO
<i>Tethya irisaе</i>	<i>Tethya irisaе</i> MHS18072 or MHS11148	South Australia: Great Australian Bight	SAMA S2913	MHS18072	MHS11148	NO
<i>Tethya martini</i> sp. nov.	<i>Tethya martini</i> sp. nov. i1	Panama: Bocas del Toro	MCZ:IZ:133702	OQ948211	OQ947275	PRJNA970968
<i>Tethya martini</i> sp. nov.	<i>Tethya martini</i> sp. nov. i2	Panama: Bocas del Toro	MCZ:IZ:135212_2	PP594405		PRJNA970968
<i>Tethya martini</i> sp. nov.	<i>Tethya martini</i> sp. nov. i3	Panama: Bocas del Toro	MCZ:IZ:135212_1	PP594406		NO
<i>Tethya minuta</i>	<i>Tethya minuta</i> EF584567	Germany: Vivarium of the Museum for Natural History, Karlsruhe		EF584567		NO
<i>Tethya norvegica</i>	<i>Tethya norvegica</i> EF558565	Norway: Trondheimfjord		EF558565		NO
<i>Tethya norvegica</i>	<i>Tethya norvegica</i> i2	UK: Rockall Bank, UK EEZ	Rockall Bank 1 (22) i2 = SPI4348001	OQ948207	OQ947281	PRJNA970968
<i>Tethya norvegica</i>	<i>Tethya norvegica</i> i3	UK: Rockall Bank, UK EEZ	UPSZMC 190662	OQ948208	OQ947282	PRJNA970968
<i>Tethya norvegica</i>	<i>Tethya norvegica</i> i4	UK: Rockall Bank, UK EEZ	UPSZMC 190664	OQ948209	OQ947283	NO
<i>Tethya orioni</i> sp. nov.	<i>Tethya orioni</i> sp. nov. i1	United Arab Emirates: Ras Ghumeis	NHMUK 2000.9.14.16	OQ948214	OQ947277	PRJNA970968
<i>Tethya samaaii</i>	<i>Tethya samaaii</i> KX618511	South Africa	TIC2016_003		KX618511	NO
<i>Tethya seychellensis</i>	<i>Tethya seychellensis</i> EF584569	Vietnam: Halong Bay, Hang Du Lake		EF584569		NO
<i>Tethya seychellensis</i>	<i>Tethya seychellensis</i> KC869475	Panama: Bocas del Toro	P09		KC869475	NO
<i>Tethya simoni</i> sp. nov.	<i>Tethya simoni</i> sp. nov. i1	Solarte Mangroves, Bocas del Toro, Panama			KC869527	NO
<i>Tethya simoni</i> sp. nov.	<i>Tethya simoni</i> sp. nov. i2	Panama: Bocas del Toro	MCZ:IZ:133701	OQ948199	OQ947279	PRJNA970968
<i>Tethya simoni</i> sp. nov.	<i>Tethya simoni</i> sp. nov. i3	Brazil: Salvador de Bahia	UFBAPOR 5019	OQ948200		NO
<i>Tethya simoni</i> sp. nov.	<i>Tethya simoni</i> sp. nov. i4	Brazil: Salvador de Bahia	UFBAPOR 5020	OQ948202	OQ947280	NO
<i>Tethya simoni</i> sp. nov.	<i>Tethya simoni</i> sp. nov. i5	Brazil: Salvador de Bahia	UFBAPOR 5022	OQ948201		PRJNA970968
<i>Tethya simoni</i> sp. nov.	<i>Tethya simoni</i> sp. nov. i5	Panama: Bocas del Toro	MCZ:IZ:135212_3	PP594407		PRJNA970968

(Continued)

Table 1. Continued.

Species	Name in trees	Locality	Voucher/Sample name	COI Accession number	28S Accession number	16S amplicon Accession number
<i>Tethya simoni</i> sp. nov.	<i>Tethya simoni</i> sp. nov. i6	Panama: Bocas del Toro	MCZ:IZ:135212_4	PP594408		PRJNA970968
<i>Tethya</i> sp. 1	MW016333	USA: Hawaii	UF4067		MW016333.1	NO
<i>Tethya</i> sp. 2	MW016337	USA: Hawaii	KBOA04101728		MW016337	NO
<i>Tethya</i> sp. 3	MW016339	USA: Hawaii	UF3981		MW016339.1	NO
<i>Tethya</i> sp. 5	MW016342	USA: Hawaii	KBOA0921711		MW016342.1	NO
<i>Tethya</i> sp. 6	MW016346	USA: Hawaii	UF3842		MW016346.1	NO
<i>Tethya</i> sp. 7	MW016348	USA: Hawaii	UF3919		MW016348.1	NO
<i>Tethya</i> sp. 8	KX866754	Israel: Ma'agan-Mikha'el	Po.25553	KX866754		NO
<i>Tethya</i> sp. 9	KJ620390	Australia	190513-05		KJ620390	NO
<i>Tethya</i> sp. 10	KU060331	Saudi Arabia: Red Sea, JazAir Silla	SNSB-BSPG.GW26962		KU060331	NO
<i>Tethya</i> sp. 11	KU060339	Saudi Arabia: Red Sea, Gulf of Alqaba	BSPG-SNSB.GW26972		KU060339	NO
<i>Tethya</i> sp. 12	KU060339/KU060752	Saudi Arabia: Red Sea, Gulf of Aqaba	BELUM:Mc5113		KU060339	NO
<i>Tethya</i> sp. 13	KU060536	Saudi Arabia: Red Sea, Qunfudhah	BSPG-SNSB.GW6033	KU060752	KU060536	NO
<i>Tethya</i> sp. 14	KU060640	Saudi Arabia: Red Sea, Al Wajh	BSPG-SNSB.GW3401	KU060640		NO
<i>Tethya</i> sp. 15	MZ487256	Caribbean PANKEY	NZCC2	MZ487256		NO
<i>Tethya stellagrandis</i>	<i>Tethya stellagrandis</i> AY561920	Maldives	POR15727		AY561920	NO
<i>Tethya wilhelma</i>	<i>Tethya wilhelma</i> EF584570	Germany: Zoological-Botanical Garden 'Wilhelma', Stuttgart		EF584570	c34318_g4_i4 (from transcriptome)	NO
<i>Tethyidae</i> sp.	AY561904		UCMPWC957		AY561904	NO
<i>Tethyidae</i> sp.	AY561986		UCMPWC1062	AY561986		NO
<i>Tethyidae</i> sp.	AY626310	Australia	G304645		AY626310	NO
<i>Tethytimea carmelita</i>	<i>Tethytimea carmelita</i> KY130425	Mexico: Campeche	LEB-ICML-UNAM-3133		KY130425	NO
<i>Timea cf. centrifera</i>	<i>Timea cf. centrifera</i> AY561906	Australia	WAMZ9801		AY561906	NO
<i>Timea</i> sp.	AY561907		G303973		AY561907	NO

(Continued)

Table 1. Continued.

Species	Name in trees	Locality	Voucher/Sample name	COI Accession number	28S Accession number	16S amplicon Accession number
<i>Timea</i> sp. AY626300	<i>Timea</i> sp. AY626300		UCMPWC957		AY626300	NO
<i>Timea</i> sp. AY626303	<i>Timea</i> sp. AY626303		G313459		AY626303	NO
<i>Timea unistellata</i>	<i>Timea unistellata</i> KC869427	Ireland	BELUM:Mc7300	KC869427		NO
<i>Trachycladus laevispirulifer</i>	<i>Trachycladus laevispirulifer</i> AY626305	Australia	WAMZ1186		AY626305	NO
<i>Trachycladus stylifer</i>	<i>Trachycladus stylifer</i> KC869453	New Zealand	NCI301		KC869453	NO
<i>Xenospongia patelliformis</i>	<i>Xenospongia patelliformis</i> KC869650	Maldives	NCI335		KC869650	NO

stained with haematoxylin and mounted in DPX. Images were taken on an Olympus BX43 light microscope with a UC50 camera. Additional spicules were mounted on SEM stubs, coated with gold, and imaged under a Zeiss UltraPlus scanning electron microscope (SEM) at the NHMUK. Measurements for megascleres were taken from light microscopy images, while measurements for megasters/micrasters were taken from SEM images using the software ImageJ (Schneider et al., 2012). All measurements were obtained from the holotype specimens unless otherwise indicated.

DNA extraction, amplification, and sequencing

Following the protocols of the Sponge Barcoding Project (www.spongebarcoding.org), we sequenced two molecular makers, the standard mitochondrial COI fragment (Folmer et al., 1994) and a fragment of the 28S rRNA ribosomal gene (28S rRNA), using the primer pair Por28S15F—Por28S1520R (Morrow et al., 2012). This encompassed 51 specimens from 11 putative species (Table 1). Small fragments of choanosomal tissue were used for DNA extraction using the Qiagen™ Blood and Tissue extraction kit, following the manufacturer's instructions. Spicules were removed by adding one spinning step after lysis and before transferring the mixture into the Qiagen columns. The quality and quantity of resulting extracts were checked with a NanoDrop spectrophotometer.

The PCR programme for COI was 94 °C for 5 min, 5 cycles (94 °C for 30 s, 45 °C for 1.5 min, 72 °C for 1 min), 35 cycles (94 °C for 30 s, 50 °C for 1.5 min, 72 °C for 1 min) and final extension at 72 °C for 7 min (Cárdenas et al., 2010). The PCR programme for 28S rRNA was 94 °C for 5 min, 30 cycles (94 °C for 30 s, 53 °C for 30 s, 72 °C for 30 s) and final extension at 72 °C for 5 min (Morrow et al., 2012). Amplification of both COI and 28S rRNA markers was performed in 12.5 µl reactions, using 10.5 µl of VWR Red Taq DNA Polymerase 1.1 × Master Mix (VWR International bvba/sprl, Belgium), 0.5 µl of the forward and reverse primers, and 1 µl of DNA template. PCR products were verified by gel electrophoresis on 1.5% agarose. Purification and sequencing of PCR products were conducted at the Molecular Core Labs (Sequencing Facility) of the NHMUK.

Individual reads were assembled and trimmed into consensus sequences using the software Geneious v.2021.1.10 (<https://www.geneious.com>). In a few cases, only forward or reverse sequences were used due to the poor quality of one of the sequencing reactions. Consensus sequences were checked for contamination

using BLAST (Madden, 2002). Available sequences of *Tethya* and Tethyidae were downloaded from GenBank. Alignment was performed with the multiple sequence alignment program, MAFFT v.7.309 (Katoh & Standley, 2013). The alignment for COI contained 658 bp for 84 taxa while the alignment for 28S rRNA contained 1500 characters for 54 taxa. Phylogenetic analyses were conducted using maximum likelihood in RAxML 8.1.22 (Stamatakis, 2014). The best-fit model of evolution was selected using jModelTest (Darriba *et al.*, 2012), resulting in HKY + I + G4 model for COI and GTR + I model for 28S rRNA. Phylogenetic analyses for COI and 28S rRNA were run separately 10 times, with 100 bootstrap replicates. Additional COI sequences of *T. burtoni* Sarà & Sarà, 2004 and *T. bergquistae* Hooper in Hooper & Wiedenmayer, 1994 from Australian waters were also included in the analyses (Table 1), although not directly sequenced in this study. Raw sequences for COI and 28S generated in this study were deposited at GenBank (Table 1).

Microbiome composition and structure

16S rRNA amplification. For 33 specimens of 9 species of *Tethya*, we targeted the V4 hypervariable region of the 16S rRNA gene to study the composition and structure of their microbiome. The V4 region was amplified using general bacterial primers 515 F-Y (Parada *et al.*, 2016) and 806 R (Apprill *et al.*, 2015), with the Illumina adapter overhang sequences in both primers and dual-barcoding approach (Kozich *et al.*, 2013). These primers contain degenerate bases to avoid the previous bias against Crenarchaeota/Thaumarchaeota and the Alphaproteobacterial clade SAR11. We used the PCR BIO HiFi Polymerase (PCR Biosystems Ltd) under the following conditions: 95 °C for 3 min, followed by 28 cycles of 95 °C for 20 s, 60 °C for 20 s and 72 °C for 30 s, after which a final elongation step at 72 °C for 5 min was performed. DNA amplification was done in duplicates, and PCR products were checked on a 1% agarose gel to determine the success of amplification and the relative intensity of bands. PCR products were purified with Agencourt AMPure XP Beads (Beckman Coulter Inc.), and libraries prepared with the Nextera XT DNA Library Preparation Kit (Illumina Inc.). An equimolar pool of DNA was generated by normalizing all samples at 4 nM for sequencing. Next-generation, paired-end sequencing was performed at the NHMUK on an Illumina MiSeq using v3 chemistry (2 × 300 bp). Raw amplicon sequence reads were deposited at the Sequence Read Archive (SRA) with BioProject accession ID PRJNA970968.

Read processing, taxonomic assignment, and core ASVs. Raw paired reads were imported into Mothur (v.1.41.3), and an adaptation of the MiSeq SOP protocol was followed (Kozich *et al.*, 2013). Briefly, primer sequences were removed and sequence contigs were built from overlapping paired reads. The merged amplicon sequence lengths were ca. 298 bp for the V4 region. Sequences with >0 N bases or with >15 homopolymers were discarded. Unique sequences were aligned against the Silva reference data set (release 132), and poorly aligned sequences were removed. Unoise3 (Callahan *et al.*, 2017), implemented within Mothur, was used for denoising (i.e., error correction) of unique aligned sequences, to infer amplicon sequence variants (ASVs), allowing one mismatch per 100 bp (Oksanen *et al.*, 2018). Any singletons remaining at this stage were removed. Reference-based chimaera checking was conducted using UCHIME with the Silva reference data set and parameter $\text{minh} = 0.3$. ASVs were classified using the Silva database v.132, with a cut-off value of 80. ASVs classified as eukaryotic-chloroplast-mitochondria or unknown were discarded, these represented 0.08% of sequences. Community sampling efficiency was examined using rarefaction curves.

Statistical design and analysis. Description of the microbial community was done using the total number of ASVs transformed to relative abundances (RA) for each individual. The core microbiome was determined on the rarefied data as the ASVs that were present in 70% of samples at any abundance, and among the phylogenetic clusters defined using the COI gene. Measures of ASV richness and Shannon index were calculated using rarefied samples in R v.4.0.5. Analysis of variance (ANOVA) was performed to compare alpha diversity among both *Tethya* clades and species, and Tukey's honestly significant difference (HSD) was used for addressing pairwise comparisons. Beta diversity was calculated using the Bray–Curtis dissimilarity coefficient on RA transformed ASVs with a minimum of 0.01% RA across samples (these included 89–98% of the total RA). The relative abundances were log₂ transformed prior to the calculation of Bray–Curtis dissimilarities. These dissimilarity matrices were visualized using Principal coordinates analysis (PcoA) using 'cmdscale' in vegan v. 2.5–7 (De Cáceres & Legendre, 2009). Permutational Analysis of Variance (PERMANOVA), using the *adonis* function of the 'vegan' package, was used to examine differences in microbial composition between *Tethya* clades. Microbial orders that were differentially abundant between the taxonomic clades and between the *Tethya* species were identified in pairwise comparisons using the 'TukeyHSD' package in R.

Orders with an adjusted (i.e., Benjamini–Hochberg corrected) p -value of less than 0.05 were considered differential between groups.

Museum acronyms. SNSB-BSPG, Bayerische Staatssammlung für Paläontologie und Geologie, Munich, Germany; INV, Museo de Historia Natural Marina de Colombia, Invemar, Santa Marta, Colombia; LEB-ICML-UNAM, Colección de Esponjas of the Instituto de Ciencias del Mar y Limnología, Universidad Nacional Autónoma de México, México DF, México; MCZ, Museum of Comparative Zoology, Harvard University, Cambridge, MA, USA; MNRJ, Museu Nacional da Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil; NHMUK, The Natural History Museum, London, UK; MNHN, Muséum national d’Histoire naturelle, Paris, France; MSNG, Museo Civico di Storia Naturale di Genova ‘Giacomo Doria’, Genova, Italy; SAM, South Australian Museum, Adelaide, South Australia, Australia; UCPM, University of California Museum of Paleontology, Berkeley, CA, USA; UPSZMC, Zoology collections of the Museum of Evolution, Uppsala University, Uppsala, Sweden; WAM, Western Australian Museum, Perth, Western Australia, Australia; UFBAPOR, Museu de História Natural da Bahia, Salvador, Bahia, Brazil.

Results

Systematics

Class DEMOSPONGIAE Sollas, 1885

Order TETHYIDA Morrow & Cárdenas, 2015

Family TETHYIDAE Gray, 1848

Genus *Tethya* Lamarck, 1814

Tethya martini Riesgo, Giribet & Santodomingo,
sp. nov.

ZooBank registration:

urn:lsid:zoobank.org:act:412FE7B6-BDE5-4678-8665-B4C11326168A

Fig. 1

Examined material: Holotype MCZ:IZ:133702 (DNA in Biobank with number DNA106632), Bahía de los Delfines (9.21685, –82.21370), Isla Cristóbal, Bocas del Toro, Panama, 5 m deep, leg. A. Riesgo & G. Giribet, 15 Mar 2010. Paratypes MCZ:IZ:135212 1 and 2 (DNA in Biobank with number DNA106830), two specimens, same locality as holotype, leg. G. Giribet, 13 Mar 2012.

Diagnosis. *Tethya* with bright red colour, thin cortex, and no spherasters. Two types of micrasters present: strongylasters and tylasters; spherasters absent.

Description. Body of holotype and paratypes hemispherical to spherical, 6–8 mm in diameter by 5–6 mm high (Fig. 1A and Table 2), attached to substrate by short lateral extensions of the base. External colour *in vivo* bright red (Fig. 1A and Table 2), whitish in ethanol. Cortical surface covered by flattened, polygonal tubercles with regular size, 0.4–0.5 mm in diameter by 0.3–0.4 mm high, homogeneously distributed and separated by reticulate areas. Some buds protrude from tubercles, 100–150 µm in diameter, stalked by thin peduncles 1.1–2.1 mm long (Fig. 1A and Table 2). Oscules 0.5–1 mm in diameter, generally located in the apex or laterally in the sponge body, if more than one; holotype with a single apical oscule and paratype with up to three, one apical and two lateral.

Skeleton. Cortex about 0.4 mm thick (holotype and paratypes) with lacunae. Radial bundles of megascleres, approximately 200 µm thick, ending in fan-shaped expansions of the cortex, subdivided into 2–3 fascicles. Without megasters in the cortex or choanoderm. Tylasters present in both the choanosome and the cortex, and strongylasters more prevalent in the choanoderm (Table 2).

Spicules. Holotype. Megascleres, main strongyloxeas with rounded or hastate ends, straight or slightly bent, 700.5–884.8–1174.5 µm long by 8.2–13.6–18.8 µm wide (Fig. 1A–B, Supplemental Fig. S1A); accessory strongyloxeas with rounded or hastate ends, 407.2–473–592.4 µm long by 5.6–7.3–10 µm wide; no spherasters. Microscleres, strongylasters, and tylasters of similar size (Fig. 1E–G). Strongylasters type 1 and type 2 with similar sizes, 9.5–11.1–12.7 µm in diameter, with small nucleus 2.5–2.9–3.3 µm, and 10–12–15 straight, cylindrical rays spined in their distal 1/2 portion, with rounded and sometimes bifurcated ends (strongylaster type 1, Fig. 1E) or 10–12–15 slightly conical rays (4.3–4.7–5.5 µm in length by 0.7–0.9–1.1 µm in width) with larger, sharper spines, along their 2/3 distal portion (strongylaster type 2, Fig. 1F). Tylasters (Fig. 1G) 8.4–9.7–10.9 µm in diameter, with slightly larger nucleus 2.7–3.2–3.7 µm, and 10–12–15 straight, short, cylindrical rays enlarged at the tips; rays 3.0–3.7–4.3 µm long by 0.9–1.1–1.2 µm wide, bearing high spines with rounded tips along the 3/4 or 1/2 of the rays, although in larger density at 1/4 tips (Table 2).

Distribution and ecology. *Tethya martini* sp. nov. has been found in shallow coral reefs in Bocas del Toro (Panama), in crevices among rubble of the scleractinian *Porites porites* (Pallas, 1766), at a depth of <5 m.

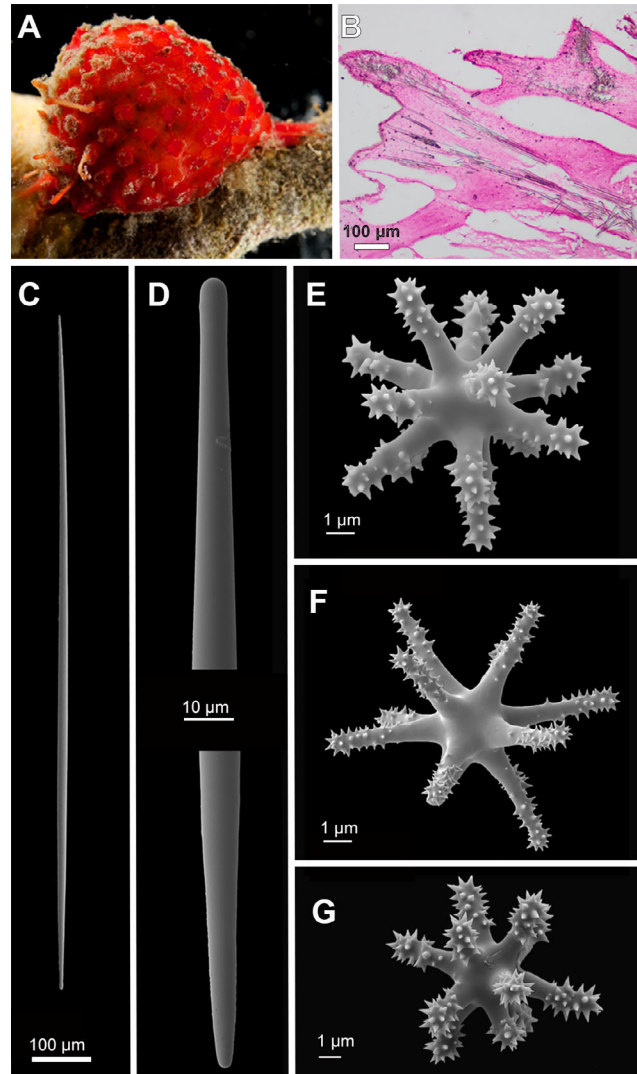


Fig. 1. *Tethya martini* sp. nov. MCZ:IZ:133702. A. Field image of a live specimen. B. Skeletal architecture of cortex and upper choanosome in cross section. C. Strongyloxea, slightly bent. D. Strongyloxea, rounded and hastate tips. E. Strongylaster type 1. F. Strongylaster type 2. G. Tylaster.

Remarks. The lack of spherasters (or megasters) in *T. martini* sp. nov. is a character shared with *T. fastigata* Bergquist & Kelly-Borges, 1991 and *T. amplexa* Bergquist & Kelly-Borges, 1991 from New Zealand (Bergquist & Kelly-Borges, 1991), as well as with the genus *Oxytethya*. With *T. fastigata*, it shares the brick-red colour when alive and a low density of microscleres in the choanosome, but *T. fastigata* differs in having deformed micrasters (~polyrhabs) and abundant oxyasters. *Tethya amplexa* has a distinct yellow colour, and a high density of micrasters and oxyasters, all features different from the new species. *Tethya martini* sp. nov. is not a species of *Oxytethya* (which lacks megasters), since the main feature of the genus *Oxytethya*, and present in the only species described so far, *O. mirabilis* Sarà & Sarà, 2002, is the abundance of oxneas as main

(4000–5000 µm) and auxiliary (180–210 µm) megascleres, and polyrhabs among the micrasters (Sarà & Sarà, 2002), both spicule types absent in *T. martini* sp. nov. In addition, *Oxytethya* is only known from SE Australia. We compared *T. martini* sp. nov. with all other species known to occur in the Caribbean, including *T. diploclerma* Schmidt, 1870 (NHMUK 1870.5.393, holotype, Supplemental Fig. S2) and the NW Atlantic (NHMUK 1938.4.26.14, Supplemental Fig. S3), *T. taboga* (de Laubenfels, 1936) from Panama (USNM 22216), *T. maza* Selenka, 1879 from Brazil (MNRJ 810, neotype), and *T. simoni* sp. nov. (Table 2, Fig. 2), but all of them possess spherasters (Table 2, Supplemental Figs. S2 and S3), making the absence of spherasters in *T. martini* sp. nov. its main diagnostic character among other species in the region.

Table 2. Diagnostic features for all material described and compared herein.

Species	External appearance/attachment	Body diameter	Colour	Skeletal arrangement	Spicules	Type locality	Material	Depth (m)	Reference
Tropical Western Atlantic/Northeastern Atlantic/Mediterranean <i>Tethya simoni</i> sp. nov.	Hemispherical to spherical, cortex (1.4 mm), reticulated surface	1.2 cm	Orange (<i>in vivo</i>), yellow-beige (in ethanol)	Radial bundles of strongyloxeas, 150–250 µm thick, form fan-shaped expansions of the cortex, single or subdivided in 2–3 fascicles	Ms = mstrongyloxeas (1107 µm), accessory strongyloxeas (480 µm), me = spherasters of two sizes (89 and 42 µm) with bifurcated and bent rays, mi = strongylaster with 15 rays (14 µm), tylasters with 14 rays (12 µm), oxyasters with 16 rays (21 µm)	Bocas del Toro, Panama	MCZ:ICZ:DNA106631	5 m	This study
<i>Tethya martini</i> sp. nov.	Hemispherical to spherical, cortex (0.5 mm), reticulated surface, tubercles and buds	8 mm	Red (<i>in vivo</i>), white (in ethanol)	Radial bundles of megascleres, approximately 200 µm thick, ending in fan-shaped expansions of the cortex, subdivided in 2–3 fascicles. Lacunae	Ms = strongyloxeas (884 µm), accessory strongyloxeas (473 µm), mi = strongylasters type 1 and type 2 with 12 spined rays (11 µm) larger in type 1, tylasters with 12 spined rays (9 µm)	Bocas del Toro, Panama	MCZ:IZ:DNA106632	5 m	This study
<i>Tethya erici</i> sp. nov.	Spherical, cortex (0.8–1 mm), flattened tubercles, buds	1 cm	Green-yellow (<i>in vivo</i>), white (in ethanol)	Radial bundles of megascleres, approximately 250 µm thick, ending in fan-shaped expansions of the cortex, not subdivided. Lacunae	Ms = strongyloxeas (860 µm), accessory strongyloxeas (340 µm), me = spherasters of two sizes (40 µm and 15 µm) with 7–13 rays, mi = strongylasters in two types of similar size (9 µm) and 9 cylindrical rays, strongylaster type 1 with short spines, and strongylaster type 2 with larger spines, tylasters (8 µm) and 7–10 rays with sharp, short spines pointing outwards, small oxyasters (9 µm) with thin 6–9 rays	Chiptoma, Spain	MNCN 1.01/1023	2 m	This study
<i>Tethya actinia</i>	Subspherical, verrucose, lumps (3 mm diameter), buds (1–5 mm), cortex 1 mm thick	5 cm	Bright orange (external and internal)	Fascicles of megascleres to the surface	Ms = Strongyloxeas (2000 × 20 µm), me = spherasters (30 µm, cen. 15 µm), chiasters (15 µm, 8 rays), oxyasters (32 µm, 4–7 rays)	Harrington Sound, Bermuda	NHMUK 1948.8.6.48	Littoral	de Laubenfels & Hindle, 1950
<i>Tethya diploderma</i>	Globular, papillose, roots (1–2 cm)	–	–	Fascicles of megascleres to the surface	Ms = strongyloxeas, me = spherasters (30 µm), mi = strongylasters (8.5 µm, 6–9 rays)	Antillen	NHMUK 1870.5.393	–	Schmidt, 1870
<i>Tethya diploderma</i>	Globular, hispid, cortex 0.8 cm, tubercles 0.5 cm	–	Yellow	Ascending tracts of spicules (200 µm)	Ms = tylotes (900–1200 µm), me = spherasters (67 µm),	Panama	USNM 22203	–	de Laubenfels, 1936

(Continued)

Table 2. Continued.

Species	External appearance/attachment	Body diameter	Colour	Skeletal arrangement	Spicules	Type locality	Material	Depth (m)	Reference
<i>Tethya diploderma</i>			Material of Schmidt, 1870		tylasters (10 µm), oxyasphersasters (7 µm), Ms = micronate strongyloxeas, me = spherasters (20–30 µm), mi = strongylasters type 2, tylasters type 1 (20 µm), oxyasters type 2 (14 µm), and oxyasters type 3 (16 µm)	Gulf of Mexico	MNRJ 12573	–	Ribeiro & Muricy, 2011
<i>Tethya diploderma</i>			–	Bundles of megascleres (400 µm), fanning out at the surface	Me = strongyloxeas (800 µm), me = spherasters (30 µm), mi = oxyasters type 2 (30 µm), tylasters type 1 (8 µm)	Bermuda	NHMUK 1938.4.26.14	–	Supplemental Material
<i>Tethya taboga</i>	Subspherical, attached with ribbon-shaped roots (1 cm), tuberculous protrusions (1 mm)	2 cm	Red	Tracts of fascicular megascleres not protruding.	Ms = strongyloxeas or strongyles (600 µm), styles (600 µm), me = spherasters (65 µm), tylasters type I (60 µm), oxyasters type I (60 µm)	Taboga Island, Pacific Panama	USNM 22216	Littoral	de Laubenfels, 1936
<i>Tethya globum</i>	Globular, smooth	2 cm	Bright orange	Not described	Not described	Guadaloupe, St. Thomas, Vieques	Type lost	littoral	Duchassaing de Fontbressin and Michelotti, 1864
<i>Tethya rubra</i>	Spherical or flattened, smooth surface or tuberculate, thin reticulation, cortex 0.5 mm, very low abundance of spherasters in cortex	2–2.3 cm	Red or yellow externally (<i>in vivo</i>), yellow internally	Spicules in poorly organized bundles, sometimes branching out, lacunar cortex	Ms = strongyloxeas (773–960 µm), accessory strongyloxeas (447–519 µm), me = spherasters (32 µm), mi = sinuous, bifurcated or trifurcated oxyasters (25–30 µm), tylasters (9 µm)	Siriba Island, Abrolhos Archipelago, Brazil	MNRJ 5316	intertidal	Ribeiro & Muricy, 2004
<i>Tethya brasiliiana</i>	Spherical or semi-spherical, surface tuberculate, stalked buds, cortex 0.75–2 mm	0.7–2.5 cm	Green or yellow externally (<i>in vivo</i>), yellow internally	Megasclere bundles with terminal fans, branched up to 5 brushes, rare lacunae	Ms = strongyloxeas (852 µm), accessory strongyloxeas (430 µm), me = spherasters bifurcate rays (13–84 µm), mi = strongylasters = 10 µm, microspherasters (10 µm), microoxyasters (10 µm)	Redonda Island, Abrolhos Archipelago, Brazil	UFRJPOR 4670 A	intertidal	Ribeiro & Muricy, 2004
<i>Tethya cyanea</i>	Hemispherical, tuberculate surface, buds, cortex 0.9–1 mm	1.4 cm	Dark blue externally	Megasclere bundles (100–150 µm) with terminal fans, irregular cavities, collagen layer	Ms = strongyloxeas (1026 µm), accessory strongyloxeas (532 µm), me = spherasters (30 µm)	Siriba Island, Abrolhos Archipelago, Brazil	MNRJ 6723	1 m	Ribeiro & Muricy, 2004

(Continued)

Table 2. Continued.

Species	External appearance/attachment	Body diameter	Colour	Skeletal arrangement	Spicules	Type locality	Material	Depth (m)	Reference
<i>Tethya ignis</i>	Hemispherical, smooth or tuberculate surface, buds, cortex 1–2 mm	1.5 cm	Orange (<i>in vivo</i>)	between cortex and choanosome, exogenous particles in the choanosome projecting beyond the surface, regular cavities, spherasters in the cortex	Ms = strongyloxeas (816 um), accessory strongyloxeas (432 um), me = spherasters (53 um), mi = microspined oxyasters (25 um), tylasters (13 um)	Redonda Island, Abrolhos Archipelago, Brazil	MNRJ 5322a	intertidal	Ribeiro & Muricy, 2004
<i>Tethya maza</i>	Spherical to semi-spherical, tuberculate surface separated by reticulate areas, buds, cortex 0.125–1 mm	0.6–3.8 cm	Yellow, white or orange (<i>in vivo</i>), beige or pinkish in ethanol	Bundles of megascleres, intracortical lacunae, abundant tylasters, spherasters in the cortex	Ms = strongyloxeas (1196 um), me = spherasters (51 um, 18–23 rays), mi = strongyloxeas type 1 (20 um, 6–9 rays), tylasters type 1 (11 um, 6–8 rays), oxyasters type 1 (11 um, 7–11 rays), oxyasters type 2 (10 um, 6–7 rays)	Urca Beach, Guanabara Bay, Brazil	MNRJ 810		Ribeiro & Muricy, 2011
<i>Tethya beatrizae</i>	Subspherical, smooth, cortex 0.125–2.125 mm thick without lacunae	1.1 cm	Yellow or whitish green (<i>in vivo</i>)	Strongyloxeas and spherasters in the cortex, bundles of megascleres (125–300 um) with terminal fans (500–875 um) project beyond surface, exogenous particles in cortex	Ms = strongyloxeas (948 um), accessory strongyloxeas (414 um), me = spherasters (42 um, 15–20 rays), mi = strongyloxeas type 2 (12 um, 10–12 rays), oxyasters type 1 (12 um, 8 rays), oxyasters type 2 (8 um, 10–12 rays)	Fernando de Noronha Archipelago, Brazil	MNRJ 7802	1 m	Ribeiro & Muricy, 2011
<i>Tethya nicolae</i>	Spherical, flattened tubercles, cortex 0.1–0.5 mm	1.5 cm	White (in ethanol)	Megasclere tracts protruding beyond surface, ovoid lacunae, cortex with spherasters and oxyasters	Ms = strongyloxeas (1187 um), accessory strongyloxeas (713 um), me = spherasters (48 um, 12–20 rays), mi = strongyloxeas (13 um, 9–11 rays), tylasters type 2 (11 um, 6–8 rays), oxyasters type 2 (12 um, 10 rays)	Pitumbu-Calpso expedition, Station 1	MNHNLBIM.D. NBE.1030	45 m	Ribeiro & Muricy, 2011
<i>Tethya solangeae</i>	Hemispherical, microhispid surface, with buds and tubercles, cortex 825–2125 um	2.5 cm	Yellow	Bundles of megascleres almost cylindrical, not fanning out, sometimes forming the tubercles, cylindrical cavities	Ms = strongyloxeas (1238 um), hastate strongyloxeas (414 um), me = spherasters (46 um), mi = strongyloxeas type II with 10–13 rays (12 um), oxyasters type II with 12 smooth rays (10 um)	Pernambuco state	MNRJ 1488	0.2 m	Ribeiro & Muricy, 2011
<i>Tethya parvula</i>		<1 cm	Yellow				MNRJ 96	3 m	

(Continued)

Table 2. Continued.

Species	External appearance/attachment	Body diameter	Colour	Skeletal arrangement	Spicules	Type locality	Material	Depth (m)	Reference
	Hemispherical, hispid surface, small buds, peduncles and tubercles, resistant cortex (1500 µm)			Bundles of megascleres (100–150 µm), cortical lacunae absent	Ms = stronglyloxeas (818 µm), hastiate stronglyloxeas (400 µm), me = spherasters with 16–20 conical curved rays (37 µm), mi = strongylasters type 2 with 11–18 rays (11 µm), oxyasters type 2 with 10–12 smooth rays (9 µm), oxyasters type 3 with 10–12 spined rays (13 µm)	São Sebastião, São Paulo			Ribeiro & Muricy, 2011
<i>Tethya irregularis</i>	Spherical or subspherical, surface with very small regular cylindrical papillae, cortex 0.5 mm in thickness	4.4 mm	White (dry)	Bundles of stronglyloxeas compact with slight subdivision in the apical tract	Ms = stronglyloxeas (250–1150), not clearly distinct into two categories, me = spherasters and oxypherasters (50–80 µm, R/C 0.2–1.5), in the choanosome also 30 µm, 12–14 rays, many with abnormal rays, bent, shortened, or reduced to hemispherical outgrowths; mi = tylasters 10–15 µm, with a developed centre, 12–14 slender rays ending in a small knob, similar in the cortex and in the choanosome	Lanzarote, Canary Islands	ZMA 1, ZMA 2–4		Sarà & Bavestrello, 1998
<i>Tethya gracilis</i>	Spherical to subspherical, compressible, flattened tubercles, buds, cortex 1.25 mm, lacunar in the outer zone. Lateral stolons	0.6–1.2 cm	White (live and in ethanol)	Radial bundles, 250 µm, fanning and with fascicles. Megasters in 1–2 layers, separated by tissue zone, absent in choanosome, some small megasters upper in choanosome.	Ms = stronglyloxeas often a little curved (900–1800 µm × 11–15 µm), auxiliary stronglyloxeas (275–950 µm × 2–5 µm), me = spherasters (15–45 µm, 16–18 rays), tylasters (6–10 µm, 3–6 rays, often tetroradiate or few anomalous with forked or distorted rays), strongylasters and oxyasters (10–20 µm, 6–8 slender rays), sometimes forked, distorted or with bifid tips	Aquarium of the Aquarium of the Azarzo Løbckke-Museum Düsseldorf	MSNG 50687	Aquaria	Sarà <i>et al.</i> , 2001
<i>Tethya aurantium</i>	Spherical, flattened tubercles, one oscule sometimes visible, thick cortex	1–5 cm	Orange (<i>in vivo</i>)	Radial bundles not forked, spherasters multilayered in cortex, interstitial megascleres in between bundles.	Ms = stronglyloxeas (400–2500 µm × 5–30 µm), me = spherasters (50–80 µm), mi = tylasters (10–15 µm, 9–14 rays), strongylasters (10–15 µm, 9–14 rays), and slender oxyasters (15–25 µm, 12–14 rays)	Bay of Naples, Italy	MSNG 49670	Littoral	Sarà, 2002

(Continued)

Table 2. Continued.

Species	External appearance/ attachment	Body diameter	Colour	Skeletal arrangement	Spicules	Type locality	Material	Depth (m)	Reference
<i>Tethya meloni</i>	Spherical, flattened or scarcely prominent tubercles, one large osculum, thick cortex (0.3–0.5 cm) and may be 1/3 of sponge diameter. Thick collagenous layer	2.9–7.6 cm (5.2 cm)	Cream to pale yellow outside and green-brown inside	Radial bundles wider at cortex, megastyles evenly and densely distributed in cortex, scattered in choanosome; micrasters in a thin external cortical layer and also scattered in choanosome	Ms = large strongyloxeas (730–1890 um × 10–40 um, mean 1310 × 25 um), auxiliary strongyloxeas (340–840 um × 8–10 um, mean 590 × 9 um), me = oxyspherasters slightly curved or bifurcated (65.7–119 um, 12–16 rays), R/C = 1–1.8 (mean 1.4), mi = chiasmers-tylasters and oxyasters (12–16 um, 8–11 rays), smaller in cortex (13.7–14.9 um, mean 14 um), larger in choanosome (15–17.4 um, mean 16 um), small spines usually at tips Ms = strongyloxea (235–1400 um × 7.5–24 um), me = oxyspherasters (18–64.8 um, mean 42.9 um, 10–20 rays, mean 15), R/C = 0.8–1.4 mean 1.1, mi = oxyaster, chiasmer to tylaster (10–15 um, mean 12 um)	Mar Piccolo di Taranto, SW Apulia, Italy	Museum of Zoology, Bari University, no reg. number	1–120 m, rocky shores, seagrass, deep sea coral gardens (Tuscan Archipelago)	Corriero et al., 2015
<i>Tethya citrina</i>	Spherical, smooth or conulose surface, thin cortex (1–1.5 mm, mean 1.2 mm), thin collagenous layer	0.5–2.6 cm (1.5 cm)	Yellowish or greenish	Radial bundles, wider at cortex, not forked, spherasters monolayered in cortex	Ms = strongyloxea (880–1250 um × 8–18 um), auxiliary strongyloxeas (365–820 um × 7–14 um), me = oxyspherasters (31–60 um, 13–18 rays), R/C = 0.4–0.78, mi = oxyspherasters (10–19 um, 10–13 rays), strongylosters (8–18 um, 11–14 rays), tylaster (8–10 um, 7–9 rays) slightly spinulated rays	Stella di Mare, Adriatic Sea, Italy	Museum of Zoology, Bari University, SDM I	0–3 m	Sarà & Melone, 1965
<i>Tethya hibernica</i>	Spherical, surface conulose to papillose, tubercles may have stalked buds, cortex compact (0.8–1.4 mm) relatively few lacunae	0.5–2.5 cm	Ochreous-yellow to yellow. In ethanol, white with greyish core	Radial bundles, dense, forming terminal fans, megastyles evenly and densely scattered in the cortex, micrasters form a discrete layer at exopinacoderm surface and endopinacoderm of lacunae, equally distributed in cortex and choanosome	Ms = strongyloxea (880–1250 um × 8–18 um), auxiliary strongyloxeas (365–820 um × 7–14 um), me = oxyspherasters (31–60 um, 13–18 rays), R/C = 0.4–0.78, mi = oxyspherasters (10–19 um, 10–13 rays), strongylosters (8–18 um, 11–14 rays), tylaster (8–10 um, 7–9 rays) slightly spinulated rays	Rathlin Island, Northern Ireland	Mc3037	Infralittoral to 30 m	Heim et al., 2007
<i>Tethya norvegica</i>	Spherical, contiguous and conulose papillae, consistency resilient, some specimens with	1–2 cm	Ochreous-yellow	Radial bundles, spirally arranged, dense, terminating in a cortical papilla surrounded by a small fan of slender styles. Megastyles in outer	MS = strongyloxeas (1200–1500 × 20 um) subtylostyles, auxiliary slender styles (500–700 um × 5–10 um), me = spherasters (20–60 um, usually 30–50 um, 15	Norway	Syntypes BMNH 1877.5.21.152 and BMNH 1877.5.21.157	10–800 m	Sarà et al., 1992

(Continued)

Table 2. Continued.

Species	External appearance/attachment	Body diameter	Colour	Skeletal arrangement	Spicules	Type locality	Material	Depth (m)	Reference
Somali/Arabian <i>Tethya orioni</i> sp. nov.	pedunculate buds 1–2 mm diameter; cortex thin (1 mm)	2.7 cm	White in ethanol	layer of cortex, sparse and irregular distribution. Microasters in a dense sheet under surface, but also in choanosome, slightly smaller more abundant in cortex	rays) with spines, divided ends and other irregularities, R/C = 0.8–1, mi = oxyasters and rare strongylasters (6–15 um, mainly 8–12 um, 8–12 rays), smooth rays and occasionally developed centre	Abu Dhabi, Persian Gulf	NHMUK 2000.9.14.16		This study
<i>Tethya omanensis</i>	Spherical or subspherical, smooth with flattened tubercles, cortex 1 mm thick	1–1.5 cm	White in ethanol	Radial bundles of strongyloxeas, 280–400 um thick, form fan-shaped expansions of the cortex, generally subdivided into 3–4 fascicles	Ms = strongyloxeas (1349 um), medium strongyloxeas (856 um), small strongyloxeas (466 um), me = spherasters of 2 sizes (53 and 29 um), mi = strongylasters with 7 spined rays (27 um), oxyasters with 6 spined rays (32 um), tyasters with 9 crown spined rays (13 um)	Tiwi Sink Hole, Oman	MSNG 48598	3–9 m	Sarà & Bavestrello, 1995
<i>Tethya robusta</i>	Subspherical, smooth with depressed tubercles	2–5 cm	Light grey (dried)	Multipisculous fascicles, closely compacted, expanding at the distal apices to tubercles	Ms = strongyloxeas (800–900 um), me = spherasters (60 um), mi = strongylasters (10 um), oxyasters (7 um)	Australia	BMNH		Wright, 1879 (measures from present study)
<i>Tethya seychellensis</i> (Mahe)	Spherical	2.5 cm	Brown	Thick bundles of macroscleteres	Ms = strongyloxeas (1000 um), me = spherasters, two sizes (20 and 50 um), mi = strongylasters (10 um), oxyasters (40 um)	Mahé	BMNH	20–25 m	Dendy, 1916
<i>Tethya seychellensis</i> (Sealark)	Subspherical, tessellated, pore-grooves, cortex lacunar	2.5 cm	Grey-yellow (ethanol)	Thick bundles of macroscleteres radiating in stout widely separated fascicles penetrating the cortex	Ms = styles and strongyloxeas, me = two types of spherasters (70 um), oxyasters (40 um, 6 rays)	Praslin Reef, Salomon Islands	BMNH		Dendy, 1916
<i>Tethya stellagrandis</i> (Sealark)	Irregularly spherical, tessellated, cortex 0.7 mm	2.5 cm	Grey-yellow (ethanol)	Thick bundles of macroscleteres radiating in stout widely separated fascicles penetrating the cortex	Ms = styles and strongyloxeas (1900 um), me = spherasters (250 um), mi = chasters (16 um, 10 rays), chasters with slender rays (16 um)	Amirante, Salomon	BMNH	73 m	Dendy, 1916
<i>Tethya ingalli</i> (Sealark)	Subspherical, strongly tessellated in polygonal areas, root-like processes of attachment, cortex 3–4 mm	2.5 cm	Yellowish grey (ethanol)	Spicule bundles penetrate the cortex and spread out into brushes	Ms = strongyloxeas, me = spherasters (80–140 um) R/C = 0.5, mi = cortical tyasters (12–16 um), choanosomal tyaster, strongylaster, oxyasters with short rays (12–20 um)	Praslin Reef, Coctivy, Egmont reef, Salomon Islands	BMNH	18–25 m	Dendy, 1916

(Continued)

Table 2. Continued.

Species	External appearance/attachment	Body diameter	Colour	Skeletal arrangement	Spicules	Type locality	Material	Depth (m)	Reference
<i>Tethya ingalli</i>	Spherical	—	White	—	Ms = strongyloxeas, me = spherasters (55 µm), mi = tylasters (13 µm), oxyasters with 6 rays (5 µm)	Djibouti, Messageries Reef	—	—	Topsent, 1906
<i>Tethya robusta</i> (previously known as <i>Donatia arabica</i>)	Spherical with tubercles, thick cortex, lacunar (not always)	—	Pink, orange	—	Ms = strongyloxeas (1000 µm), tylotes, me = two types of spherasters (75 µm and 15 µm), mi = tylasters and few oxyasters (30 µm)	Bay of Djibouti	—	—	Topsent, 1906

Abbreviations: Megasclere tracts (diameter in µm). Ms, megascleres; me, megasters; mi, micrasters (average size in µm).

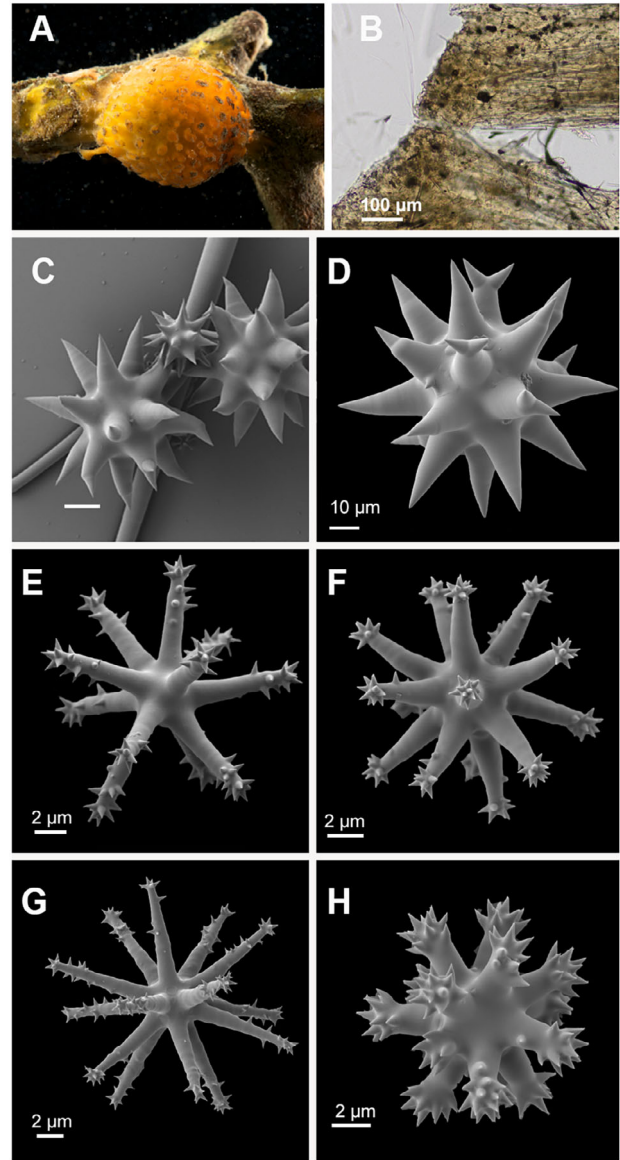


Fig. 2. *Tethya simoni* sp. nov. MCZ:IZ:133701. A. Field image of a live specimen. B. Skeletal architecture of cortex and upper choanosome in cross section. C. Spherasters, large (with bent tips) and small-size classes. D. Common spherasters with bifurcated tips. E. Strongylaster type 1. F. Strongylaster type 2. G. Oxyaster type 1. H. Tylaster type 1.

Etymology. The species is named after Martin Taboada, the firstborn son of Ana Riesgo.

Tethya simoni Santodomingo, Zea & Riesgo, sp. nov.
 ZooBank registration:
 urn:lsid:zoobank.org:act:D9D59AFD-4D20-44C0-8737-6243542CD6B5

Fig. 2

Examined material. Holotype: MCZ:IZ:133701 (ex. DNA106631), Bahía de los Delfines (9.21685, -82.21370), Isla Cristóbal, Bocas del Toro, Panama, 15 Mar 2010, leg. A. Riesgo & G. Giribet, 5 m deep. Paratypes: MCZ:IZ:135212 (spec. 3 and 4), same locality as holotype, leg. G. Giribet, 13 Mar 2012; UFBAPOR 5019, UFBAPOR 5020, UFBAPOR 5021, Intertidal rocks on an arenitic plateau at Praia da Pituba, Salvador de Bahia, Brazil, 25 Jan 2018, leg. E. Lanna & F. Calvacanti.

Diagnosis. *Tethya* with thick cortex, spherasters present, in two sizes; larger spherasters often with bifurcated tips, three types of micrasters: strongylasters = tylasters \ll oxyasters.

Description. Body hemispherical to spherical 0.8–1.2 cm in diameter by 0.9–1.2 cm high (Fig. 2A). Live specimens orange (Fig. 2A), becoming yellow or beige in ethanol. Pinacoderm covered by flattened, circular, irregularly distributed tubercles of variable sizes, 0.2–0.6 mm in diameter with a fiddle-shaped peduncle 0.4–0.7 mm high, separated by reticulate areas. In paratype specimens from Brazil, no tubercles were observed. No oscules evident on holotype nor paratypes.

Skeleton. Cortex 1–1.4 mm thick, with regular lacunae, mainly constituted by abundant spherasters and a thin, outer layer of tylasters. Deposits of collagen form a dense layer between the cortex and choanosome. Main radial bundles of strongyloxeas, 150–250 μ m thick, form fan-shaped expansions of the cortex, single or subdivided in 2–3 fascicles. Tylasters present in both choanosome and cortex. Strongylasters and oxyasters more abundant in choanosome.

Spicules. Holotype. Megascleres, strongyloxeas, and spherasters. Main strongyloxeas with rounded and hastate ends, straight, 810.2–1107.5–1470.5 μ m long by 9.2–19.3–30.3 μ m wide; accessory strongyloxeas with rounded and hastate ends, 210.9–480.3–600.5 μ m long by 2.6–9.9–18.4 μ m wide (Supplemental Fig. S1B). Spherasters in two sizes, large spherasters more abundant, 72.1–89.5–110.1 μ m in diameter, large core 33.3–42.9–53.2 μ m, R/C 0.4–0.6–0.7, with 20–22–26 smooth, conical rays, although mainly with variable shapes, more commonly with bifurcated rays (47%), mammillated (20%), or rays with bent tips (20%) (Fig. 2C–D); small spherasters sparse in the cortex with 22–24–26 smooth, conical, straight rays, 45.1–55.9–62.9 μ m in diameter, large core 20.5–30.7–35.6 μ m, R/C 0.3–0.4–0.6 (Fig. 2C). Microscleres, tylasters, and strongylasters of similar size, larger oxyasters. Strongylasters 11.8–14.7–

17.7 μ m in diameter, with small nucleus 3.2–4.0–4.6 μ m in diameter, and 14–15–16 straight, cylindrical rays with rounded ends, bearing a crown of high, sharp spines, rarely with sparse 1–2 spines at $\frac{1}{3}$ to $\frac{1}{4}$ of distal portion of rays; rays 4.2–5.6–7.2 μ m long by 1.1–1.5–1.9 μ m wide (Fig. 2E–F). Tylasters 10.3–12.5–13.9 μ m in diameter, with large nucleus 3.4–4.6–5.4 μ m, and 12–14–15 straight, short, cylindrical rays enlarged at the tips; short, robust rays 3.7–4.6–5.5 μ m long by 1.7–2.0–2.3 μ m wide, with large, sharp spines concentrated at the $\frac{1}{4}$ ends, mostly pointing upwards; some rays bearing a small spine at their basal part (Fig. 2H). Large oxyasters 18.5–21.7–27.9 μ m in diameter, reduced nucleus 4.0–4.5–5.7 μ m, and long, straight, thin 12–16–19 rays; rays 7.1–8.4–10.6 μ m long by 1.3–1.6–2.0 wide, bearing short, blunt or acerate spines, sparse $\frac{1}{2}$ towards the tips, in more density at the $\frac{1}{4}$ ends of the rays (Fig. 2G); some oxyasters with bifurcated rays at the $\frac{1}{3}$ proximal part.

Distribution and ecology. *Tethya simoni* sp. nov. inhabits the coral reefs of Bocas del Toro (Panama), where it has been found attached to fragments of dead branches of *Porites porites* and other hard corals, around 5 m depth. In Brazil, it was collected under rocks in an intertidal arenitic plateau. This new species is distributed from the southern Caribbean (Panama) to the northeast coast of Brazil.

Remarks. Reports of *T. aurantium* from the Caribbean date back to 1936 in a study of the sponges on both sides of the Panama Canal (de Laubenfels, 1936). Subsequent records include specimens from Puerto Rico and the Dominican Republic (Sarà & Gaino, 1987), from Cuba (Alcolado, 1985), and it is also in a recent checklist from Panama (STRI Research Portal, 2023). These records and some others could be attributed to *T. simoni* sp. nov. (see below). *Tethya aurantium* was described from the Mediterranean Sea, and although we did not have access to the neotype (MSNG 49670, Bay of Naples), *T. aurantium* material studied here was collected in the Bay of Naples. Measurements of our *T. aurantium* specimens from the Mediterranean show a wider and continuous range of strongyloxea size (544.5–1423.2–2366.3 μ m long by 11.8–20.8–29.5 μ m wide) as well as spheraster size (24.3–76.7–105.1 μ m in diameter), which correspond to the neotype description (Sarà, 2002; Table 2). In *T. simoni* sp. nov., strongyloxeas have smaller sizes, and spherasters with bifurcated rays are common, contrary to a unique instance observed in this study on specimen MCZ:IZ:106628 of *T. aurantium* collected in the type locality, in Naples, Italy (Tables 1–2). De Laubenfels (1936) noticed the similarities between his Panamanian specimens and those of *T.*

aurantium from the Mediterranean but remarked the presence of two sizes of spherasters, which are not described for *T. aurantium* but are diagnostic for *T. simoni* sp. nov. (Fig. 2C, Table 2). The molecular affinity of *T. aurantium* and *T. simoni* sp. nov. is clear, as they are sister species in a well-supported clade using two different markers (Figs 3 and 4) and their dominant haplotypes are separated by 8 mutational steps (Fig. 3).

Tethya simoni sp. nov. is constrained to the West Atlantic, and so far only found in the southern Caribbean (Panama) and northeast Brazil. Other examined specimens of the NHMUK collections from the Caribbean and close-by areas (Table 2), included the type of *T. diploderma* NHMUK 1870.5.393 (West Antilles) (Supplemental Fig. S2), USNM 22203 (Panama) and NHMUK 1938.4.26.14 (Bermuda; Supplemental Fig. S3), *T. actinia* BMNH1948.8.6.48 (Bermuda) and *T. taboga* USNM 22216 (Panama). The most important character separating *T. simoni* sp. nov. from these other Caribbean species is the presence of two sizes of spherasters, the most abundant one much larger than those in the other species (Table 2).

Since *T. simoni* sp. nov. also appears in Brazil, we compared it with recently described species from the coast of Brazil, comprehensively reviewed in Ribeiro and Muricy (2011). *Tethya rubra* Samaai & Gibbons,

2005, *T. cyanae* Ribeiro & Muricy, 2004, and *T. ignis* Ribeiro & Muricy, 2004 lack strongylasters, while those are conspicuous in *T. simoni* sp. nov. Diagnostic characteristics of *T. beatrizae* (previously identified as *T. aurantium* sensu Hechtel, 1976 in Ribeiro & Muricy, 2011), such as having a smooth surface (no tubercles) and oxyasters without spines (oxyaster type 2) differ from those of *T. simoni* sp. nov. The main difference from other Brazilian *Tethya* species is the presence of two types of spherasters, one larger than those present in most other Brazilian species, except for *T. brasiliانا* Ribeiro & Muricy, 2004 (Table 2). In this particular case, the character distinguishing *T. brasiliانا* and *T. simoni* sp. nov. is the absence of tylasters in *T. brasiliانا* (Ribeiro & Muricy, 2004). The Brazilian species, *T. nicolae* Ribeiro & Muricy, 2011 and *T. maza* Selenka, 1879, share some characteristics with *T. simoni* sp. nov. *Tethya nicolae* has also spherasters with bifurcated tips, but of much smaller size (37–48–57 µm) and although strongylasters and tylasters have similar shapes and sizes in both species, oxyasters in *T. nicolae* are smaller with smooth rays, contrasting with the typical spinose and sometimes bifurcated in *T. simoni* sp. nov. In comparison with *T. maza*, which also seems to have a wide range of spherasters that vary from a small size (26 µm)

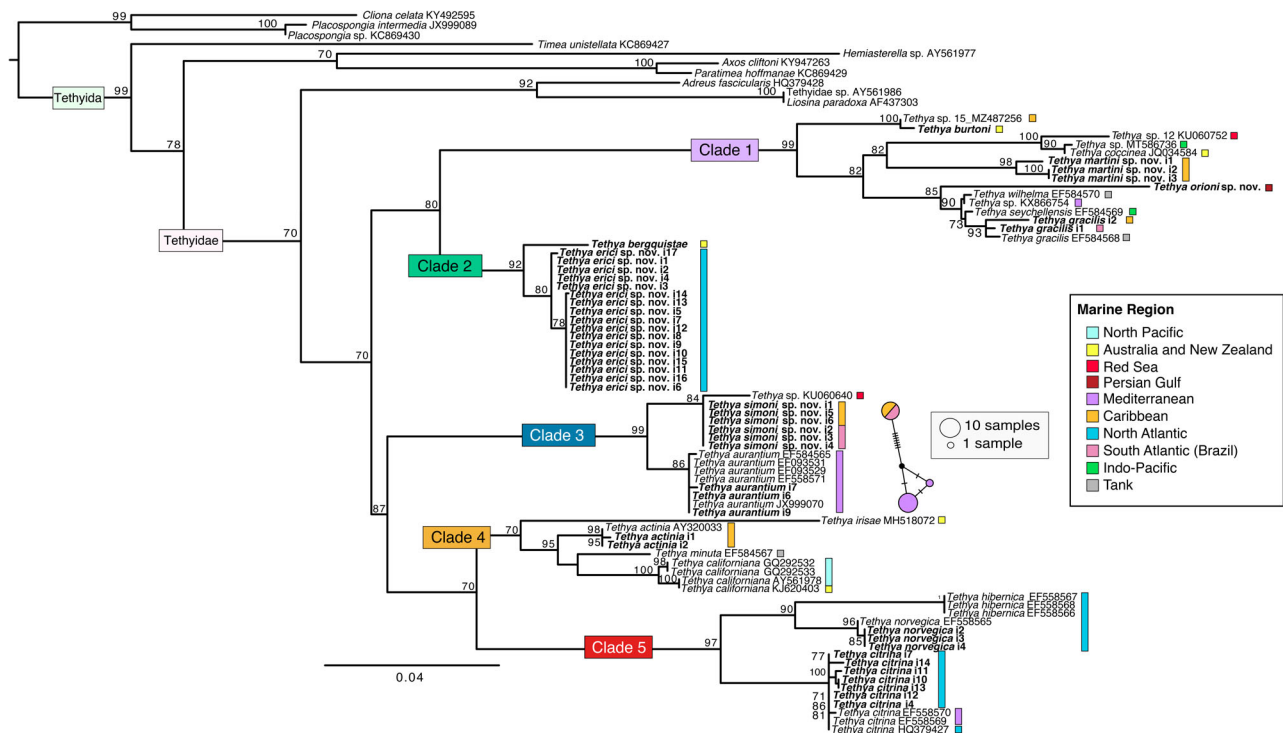


Fig. 3. Maximum likelihood consensus tree for the *cytochrome c oxidase subunit I* (COI) alignment. Only bootstrap support values over 70% are shown. In the haplotype network, each circle is a different haplotype for *Tethya aurantium* and *T. simoni* sp. nov., each colour represents a locality, the size of the circle is proportional to that haplotype frequency, and the length of the line is proportional to the number of mutations. Known geographic regions for species in the tree are shown with coloured squares. Specimens in bold letters were sequenced in this study for the first time.

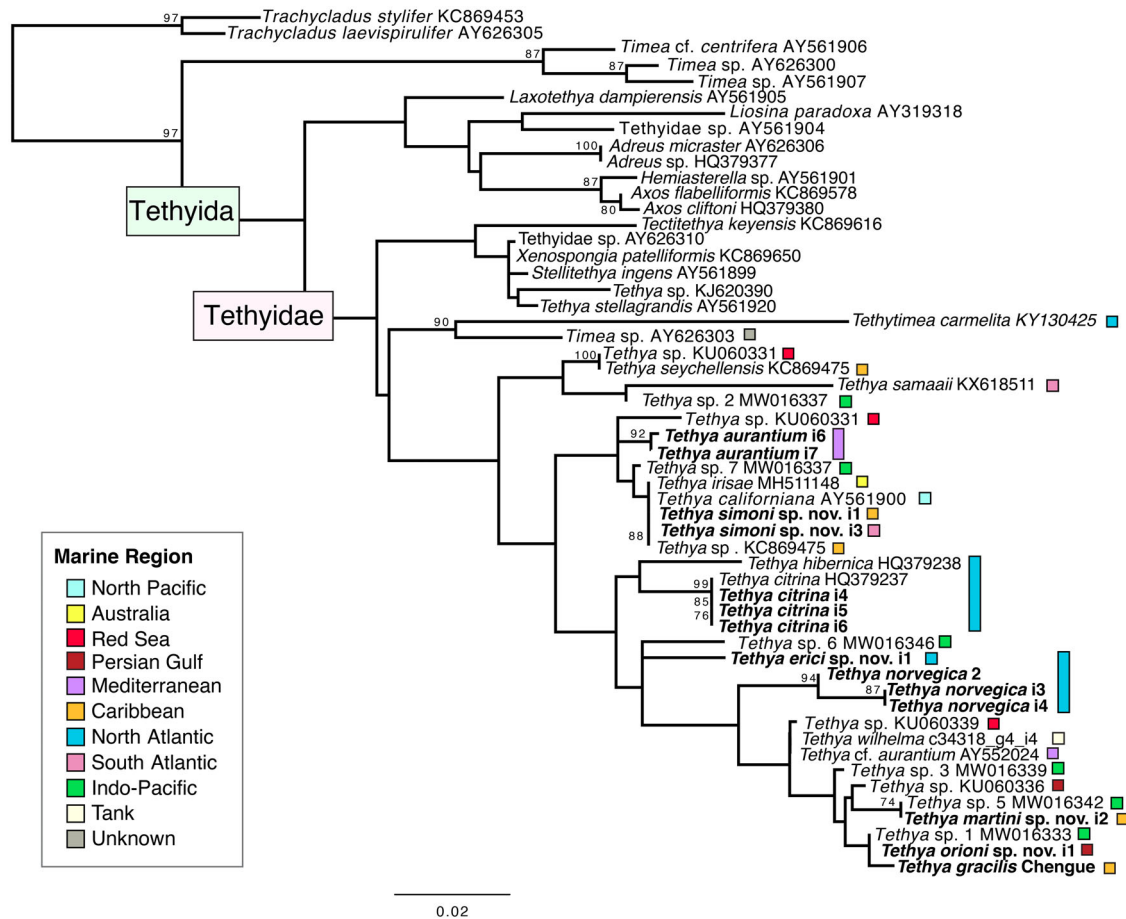


Fig. 4. Maximum likelihood consensus tree for the 28S rRNA alignment. Only bootstrap values over 70% are shown. Known geographic regions for species in the tree are shown with coloured squares. Specimens in bold letters were sequenced in this study for the first time.

to a larger size (100 μ m) and some with bifurcated tips, this species has polyrhabs-like spicules (see fig. 1F in Ribeiro & Muricy, 2011, referred to as oxyasters type 1), which are absent in *T. simoni* sp. nov. The wide distribution of *T. maza* along the Brazilian coast, and similarities in the skeletal features, suggest that this species may be part of the same lineage as *T. aurantium* and *T. simoni* sp. nov.

We also compared *Tethya simoni* sp. nov. with specimens collected in the Bay of Chengue (Colombia) (Supplemental Fig. S4) and in Salvador (Brazil) that belong to *T. gracilis* Sarà (Sarà et al., 2001). While *T. gracilis* was described from aquaria in the Düsseldorf Museum for the first time, it is likely that the species originated in a tropical setting that the authors located in the Indo-Pacific based on the affinities of *T. gracilis* to the previously sequenced *T. seychellensis* (see Sarà et al., 2001). However, the sponges could have also originated in the Caribbean. Although the spicular content is similar in both *T. simoni* sp. nov. and *T. gracilis*, *T.*

simoni sp. nov. has spherasters of two sizes, with the larger class doubling the size of the ones found in *T. gracilis*, and the stronglyloxeas are much smaller in the former species (Table 2).

Etymology. The species is named after the son of Nadia Santodomingo, Simon Peter-Contesse Santodomingo.

Tethya erici Díez-Vives, Santodomingo, Moles, & Riesgo, sp. nov.

ZooBank registration: urn:lsid:zoobank.org:pub:54F80E5E-99A4-43CE-BA3A-BCD01E715595

Fig. 5

Examined material. Holotype: MNCN 1.01/1023, Chipiona ‘Corrales’, Cádiz, Spain (36.74168, -6.43887). Leg. Juan Moles, 2 m depth, Oct 2014. Paratypes: MNCN 1.01/1024 and MNCN 1.01/1025,

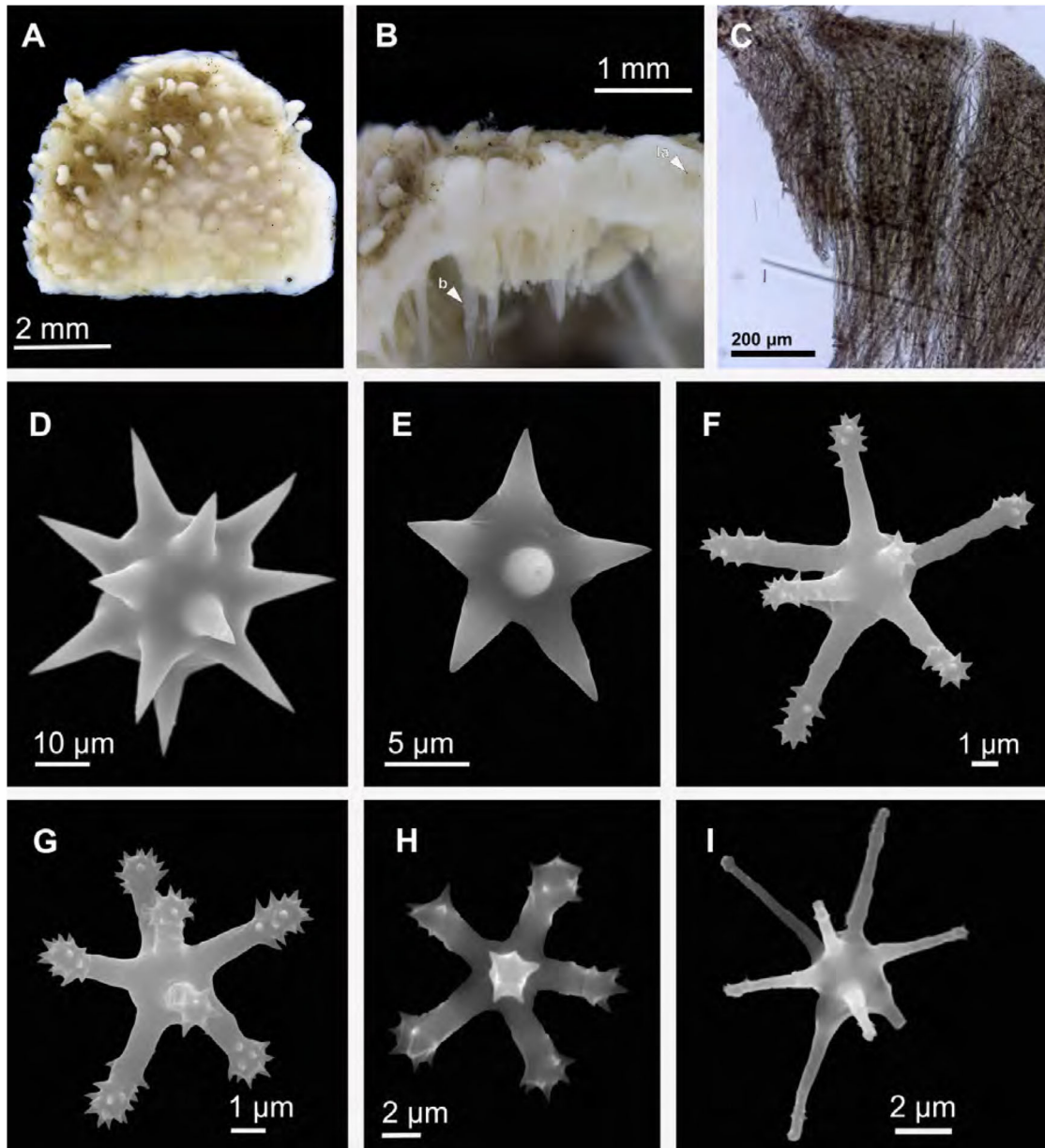


Fig. 5. *Tethya erici* sp. nov. MNCN 1.01/1023. A. Specimen preserved in ethanol. B. Detail of cortex showing bundles (b) and lacunae (la). C. Skeletal architecture. D. Spheraster type 1. E. Spheraster type 2. F. Strongylaster type 1. G. Tyloaster type 1. H. Tyloaster type 2. I. Oxyaster.

collected with the holotype, and 14 specimens MNCN 1.01/1027 to MNCN 1.01/1041 collected in La Palma, Canary Islands, Spain (28.693245, -17.759086), leg. Juan Moles, 2 m depth, Sept 2022.

Diagnosis. *Tethya* with thin cortex, small strongyloxeas in two sizes, spherasters in two sizes, micrasters: strongylasters, tyloasters, and oxyasters, all with similar size ranges, but strongylasters slightly larger and in two types.

Description. Body hemispherical to spherical, 1 cm in diameter; colour green in live specimens, in alcohol white externally, beige internally (Fig. 5A–B). Pinacoderm covered by flattened, plate-like tubercles of variable sizes 0.3–0.4 mm in diameter by 0.1–0.2 mm high, irregularly distributed. Buds with mushroom-like shape, protruding from some tubercles, peduncle 0.4–0.6 mm high and 320–400 µm in diameter (Fig. 5A–B). No oscules evident on holotype nor paratypes. Cortex firm, resistant; choanosome compressible.

Skeleton. Cortex 0.85–0.98 mm thick with regular lacunae (Fig. 5B), mainly formed by abundant spherasters, tylasters, and oxyasters (Fig. 5C). Thin layer of collagen between cortex and choanosome. Main radial bundles of strongyloxeas 250–400 µm thick, forming single fan-shaped expansions of the cortex. Strongylasters mainly present in choanosome, with abundant tylasters and oxyasters, and sparse spherasters (Fig. 5B–C).

Spicules. Holotype. Megascleres, strongyloxeas, and spherasters. Main strongyloxeas with rounded and hastate ends, straight, 808.7–859.1–983.1 µm long by 11.5–14.5–20.4 µm wide; accessory strongyloxeas with rounded and hastate ends, 236.9–340.2–472.86 µm long by 3.6–5.4–7.5 µm wide (Supplemental Fig. S1C). Two sizes of spherasters, small when compared with other *Tethya* species. Spherasters of larger size 15.4–40.1–56.7 µm in diameter, large core 8.2–19.5–25.3 µm, R/C 0.4–0.5–0.7, with 11–13–17 smooth, conical rays (Fig. 5D), and smaller spherasters of 15.2–15.6–16.3 µm, core 6.1–7.1–8.2 µm, R/C 0.5–0.6, with 10–12 conical rays (Fig. 5E). Microscleres, mean size of strongylasters slightly larger than tylasters and oxyasters, but all have similar size range. Strongylasters in two types, 6.9–9.6–11.7 µm in diameter, with small nucleus 1.1–2.6–3.7 µm in diameter, and 7–9–13 straight, cylindrical rays with rounded ends bearing spines at $\frac{1}{3}$ to $\frac{1}{4}$ of distal portion of rays; strongylaster type 1 have enlarged tips with shorter, sparse spines (Fig. 5F); strongylaster type 2 have conical rounded ends, with sharp and larger spines. Tylasters 5.9–8.3–11.8 µm in diameter, with large nucleus 1.7–2.6–3.7 µm, and 7–8–10 straight, short, cylindrical rays slightly enlarged at the tips; cylindrical rays 1.6–3.1–4.2 µm long by 0.7–1.0–1.1 µm wide, sharp, short spines concentrated at the $\frac{1}{4}$ ends, pointing outwards (Fig. 5G–H). Small oxyasters 6.3–8.8–10.8 µm in diameter, reduced nucleus 0.9–1.2–1.6 µm, and long, straight, thin 6–7–9 rays; rays 2.9–3.8–4.4 µm long by 0.3–0.5–0.6 wide, bearing short spines at the $\frac{1}{4}$ ends of the rays (Fig. 5I).

Distribution and ecology. *Tethya erici* sp. nov. occurs under rocks on rocky shores in shallow waters (1 m depth), so far only found in the ‘Corrales’ on the coast of Chipiona (Cadiz, Spain) and La Palma (Canary Islands, Spain) rocky shore pools. The ‘Corrales’ (Spanish plural of corral) are artificial enclosures historically used for centuries for fishing by hand at low tides. These are rocky enclosures with scattered boulders under which *T. erici* sp. nov. was found. This region is located in the southernmost part of the North Atlantic, near the Strait of Gibraltar, which connects the Atlantic and the Mediterranean Sea. The coast is influenced by the sedimentary inputs of the Guadalquivir River and

nearby the large estuarine system of the Doñana National Park. In La Palma, the rocky ecosystems where the sponges were found were mostly intertidal pools under boulders at 2 m depth.

Etymology. The species is named after the son of Cristina Díez-Vives, Eric Nielsen Díez.

Remarks. Among the Atlantic species, *Tethya erici* sp. nov. is most similar to *T. nicoleae* Ribeiro & Muricy, 2011 from Brazil in their spicular content, but the former has smaller strongyloxeas that are in two size classes, spherasters with fewer rays and none with bifurcated tips, micrasters of smaller size and more variability with additional strongylasters type 2 and oxyaster type 3. In comparison with other North Atlantic species *T. erici* sp. nov. falls within the range of distribution of *T. aurantium* and *T. citrina*. Although the spicule types are similar to *T. aurantium*, they differ in having smaller sizes, with strongyloxeas and spherasters up to half of the size. Spherasters in *T. erici* sp. nov. have a larger core (R/C 0.4–0.7) different from the typical oxyspherasters (R/C 0.6–1.4) in *T. citrina*. Concerning *T. hibernica* and *T. norvegica*, spicules of *T. erici* sp. nov. are more similar to *T. hibernica* but the latter has typical tylasters of shorter and thicker rays (Heim, Nickel, Picton, & Brümer, 2007). Finally, in contrast to the recently described *T. meloni* Corriero *et al.*, 2015 from the Mediterranean Sea, *T. erici* sp. nov. lacks oxyspherasters (R/C 1–1.8), and tylasters described in *T. meloni* are strongylaster-like with longer rays and smaller cores (Corriero *et al.*, 2015). *T. erici* shares geographic distribution with *Tethya irregularis* Sarà & Bavestrello, 1998 from the Canary Islands, however it differs in the absence of abnormal megasters and the presence of strongylasters and oxyasters. In the molecular phylogeny, *T. erici* sp. nov. forms a well-supported clade with *T. bergquistae* from Australia and New Zealand, which has larger strongyloxeas and micrasters (Fig. 3).

Tethya orioni Kenny, Santodomingo & Riesgo, sp. nov.
ZooBank registration: urn:lsid:zoobank.org:act:FB56A
581-9311-46EF-A141-8925E0CD58E0

Fig. 6

Examined material. Holotype: NHMUK 2000.9.14.16, tip of causeway, Ras Ghumais, United Arab Emirates, Persian Gulf. Leg. J.D. George, 3–8 m depth, 15 Oct 1998.

Diagnosis. *Tethya* with thick cortex, irregular tubercles, variable spherasters with fewer rays, three types of micrasters: oxyasters = strongylasters \gg tylasters.

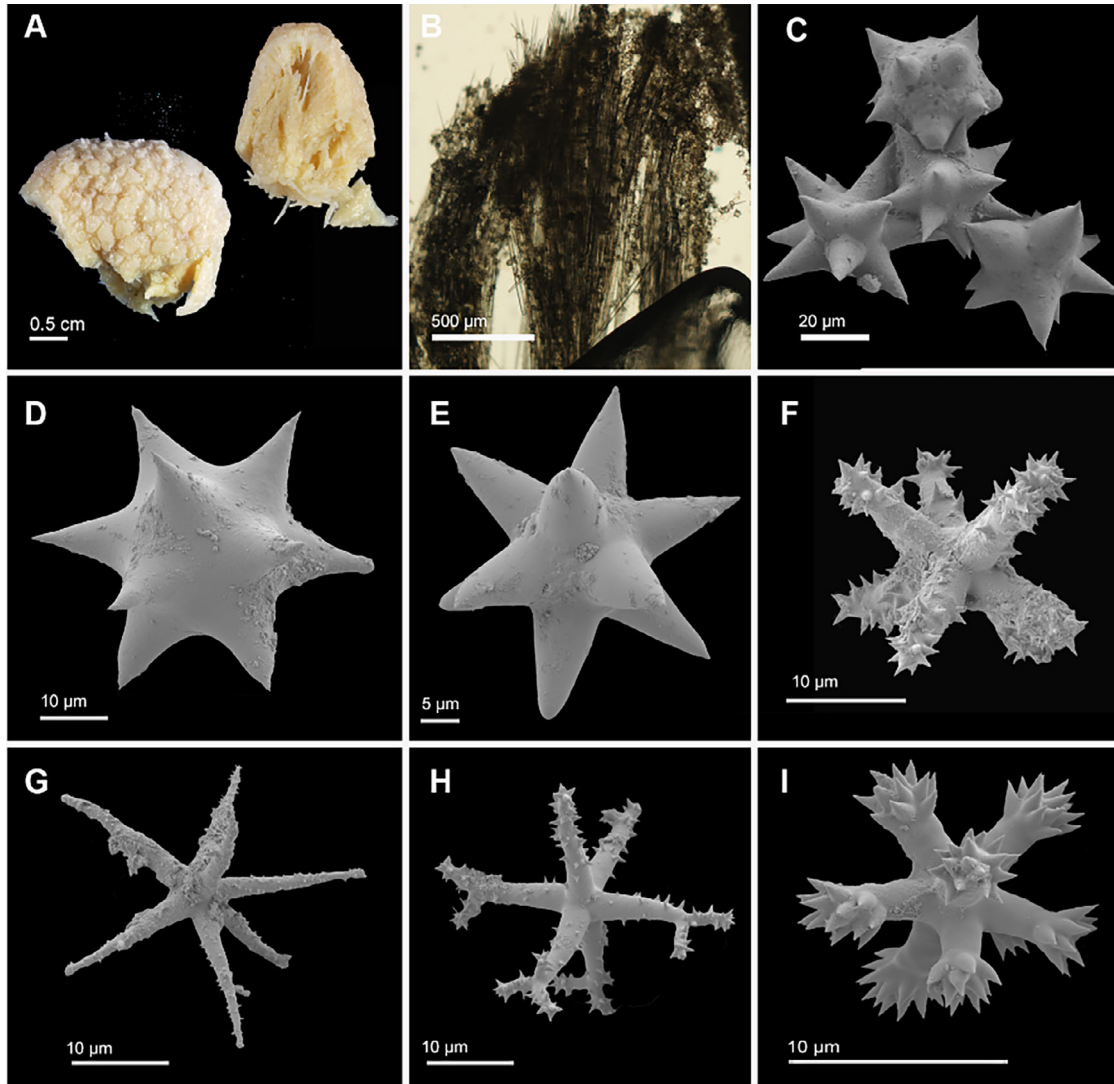


Fig. 6. *Tethya orioni* sp. nov. NHMUK 2000.9.14.16. A. Specimen preserved in ethanol. B. Skeletal architecture of cortex and upper choanosome in cross section. C. Spherasters with abnormal shape and mammillated rays. D. Large spheraster. E. Small spherasters. F. Strongyaster. G. Oxyaster. H. Oxyaster with bifurcated tips. I. Tylaster.

Description. Body spherical 2.7 cm in diameter by 2 cm high (Fig. 6A). Specimen bluish tinge when alive (J.D. George, pers. comm.) and white to light beige in ethanol (Fig. 6A). Pinacoderm covered by tightly arranged tubercles of irregular shape and rugose surface, ranging from 0.5–3.5 mm wide by 0.6–1 mm high. Some buds grow from tubercles, attached through thin peduncles 200–300 μm in diameter and 1.3–2.2 mm long. No oscules visible.

Skeleton. Thick cortex 1–3 mm thick with irregular lacunae, mainly constituted by abundant spherasters and some tylasters (Fig. 6B). Deposits of collagen under the cortex. Main radial bundles of strongyloxeas, 280–

400 μm thick, form fan-shaped expansions of the cortex, generally subdivided into 3–4 fascicles (Fig. 6B). Oxyasters and strongylasters abundant in the choanosome. Tylasters less common in choanosome than in the cortex.

Spicules. Holotype. Megascleres, strongyloxeas in three size classes, all straight with rounded and hastate ends. Main strongyloxeas abundant, conform bundles radiating from the centre of the sponge, 1114.3–1349.1–1598.8 μm long by 16.0–20.5–25.7 μm wide (Fig. 5); medium strongyloxeas, less common, 745.1–856.8–956.4 μm long by 10.7–14.8–17.6 μm wide; small, accessory strongyloxeas, profuse, 337.8–466.3–631.0 μm long by 8.0–12.5–16.8 μm wide (Supplemental Fig. S1D). Megasters,

spherasters in two sizes (Fig. 6C–E), large spherasters abundant in the cortex, 40.1–53.3–68.7 µm in diameter, large core 19.9–26.6–32.9 µm, R/C 0.3–0.6–0.9, with 8–13–16 smooth, conical rays (Fig. 6D), but often in irregular and variable shapes, more commonly with unequal rays, both pointy and rounded (20%), thinner rays with mammillae-like tips (15%), or amorphous shape (15%) (Fig. 6C); small spherasters also in variable shapes, sparse in the cortex, more abundant in the choanosome, with 9–11–14 rays, 22.3–29.7–38.1 µm in diameter, core 11.9–15.9–22.7 µm, R/C 0.4–0.6–0.8 (Fig. 6E). Microscleres, strongylasters and oxyasters of similar size, smaller tylasters. Strongylasters 22.0–27.2–35.0 µm in diameter, with small nucleus 5.1–5.8–7.9 µm in diameter, and 6–7–8 straight, cylindrical rays with rounded ends, with sharp spines sparse thoroughly along rays, concentrated towards $\frac{1}{3}$ of distal portion of rays; rays 9.9–12.5–16.8 µm long by 1.9–2.5–3.1 µm wide (Fig. 6F). Oxyasters 24.2–32.0–37.3 µm in diameter, reduced nucleus 4.1–5.4–6.4 µm, and long, straight, thin 4–6–8 rays; rays 10.6–15.2–18.5 µm long by 1.6–1.8–2.1 wide, bearing short, acerate spines, sparse thoroughly along the rays (Fig. 6G); most oxyasters with bifurcated rays at the tips (Fig. 6H), some oxyasters devoid of spines with straight rays. Tylasters 10.2–13.3–24.7 µm in diameter, with large nucleus 3.1–4.0–6.6 µm, and 8–9–10 straight, short, cylindrical rays enlarged at the tips, 3.6–5.3–10.4 µm long by 1.2–1.9–2.7 µm wide, with a crown of sharp spines concentrated at the $\frac{1}{2}$ ends, mostly pointing upwards (Fig. 6I).

Distribution and ecology. *Tethya orioni* sp. nov. was collected in a sheltered area of a rocky shore, 3–8 m depth.

Remarks. The specimen was deposited in the collections of the NHMUK under the name *T. seychellensis*, identified by Dr Michelle Kelly in 2000. Detailed examination of *T. orioni* sp. nov. in comparison with the *T. seychellensis* holotype NHMUK 1886.10.22.22 (Mahé, Seychelles, Supplemental Fig. S5), showed that both species have similar body size and, despite having densely packed tubercles on the pinacoderm surface, these seem to be more regular in *T. seychellensis*, while very irregular in size and shape in *T. orioni* sp. nov. Spherasters and oxyasters are of similar size in *T. seychellensis* (Table 2), whilst in *T. orioni* sp. nov. oxyasters are smaller than spherasters. Another difference is the presence of strongylasters in *T. orioni* sp. nov., which are absent in *T. seychellensis*. These two species share some features such as the size class of strongyloxeas and tylasters, and by having oxyasters with fewer number of rays and bifurcated tips.

Molecular analyses showed that both species are part of the same main clade (clade 1; Figs 3–4) but *T. orioni* sp. nov. can be distinguished morphologically and genetically from other *Tethya* species within that clade. The irregular shape of spherasters in *T. orioni* sp. nov. is a character also observed in *T. omanensis* Sarà & Bavestrello, 1995, described from an underwater cave in Oman, and *T. irregularis* described from the Canary Islands in the vicinity of a freshwater spring (Sarà & Bavestrello, 1998). It has been suggested that these abnormalities and eroded spicular surface are adaptations to reduced salinity (Sarà & Bavestrello, 1998). *Tethya orioni* sp. nov. was collected in a marine setting with no evident freshwater input. *Tethya omanensis* has shorter strongyloxeas (450–720–980 µm), micrasters with an enlarged nucleus, and spherasters with a higher number of rays (24–28) in comparison to those in *T. orioni* sp. nov. Regarding *T. irregularis*, micrasters have a larger number of rays (10–15). Other species described from adjacent waters in the Indian Ocean are *T. stellagrandis* (Dendy, 1916) and *T. ingalli* (Bowerbank, 1858). *Tethya stellagrandis* differs from other species in the genus in having the largest spherasters recorded (up to 250 µm) (Supplemental Fig. S6A–C). In *T. ingalli* spherasters range between 80–140 µm (Dendy, 1916), hence they are double the size of the ones found in *T. orioni* sp. nov.

Etymology. The species is named after Orion Tamati Kenny, the firstborn son of Nathan J. Kenny.

Tethya phylogeny

Phylogenetic analyses of the individual COI and 28S rRNA sequences (Figs 3–4) recovered a monophyletic order Tethyida with strong support (Figs 3–4). The family Tethyidae was also recovered as a single clade with support over 70 in the COI tree, and low support in the 28S rRNA tree (Figs 3 and 4). In the COI tree, we obtained five main clades: clades 1–3 and 5 highly supported but clade 4 with less support (Fig. 3). In clade 1 (Fig. 3), we found *T. burtoni* (from New Zealand) diverging earlier, and then *T. martini* sp. nov. (from Panama), as sister species to a well-supported clade composed by *T. coccinea* (from Australia) and two unidentified *Tethya* specimens collected in the Indo-Pacific and the Red Sea (Fig. 3). Finally, the last subclade obtained in clade 1 grouped *T. seychellensis* (from Vietnam), *T. wilhelma*, and *T. gracilis* (both described from German aquaria), and two specimens that we collected in the western Atlantic, one from the Colombian Caribbean and another from Salvador de

Bahia in Brazil that were extremely similar to the *T. gracilis* holotype. The morphology and spicule complement of the *T. gracilis* holotype (Sarà et al., 2001) and our specimen identified as *T. cf. gracilis* from the Caribbean locality of Chengue Bay in Colombia were highly similar (Supplemental Fig. S4), despite a 4 bp difference in COI, suggesting that this species may constitute a species complex, or that its potentially invasive nature has allowed it to diverge quickly into different lineages. Clade 2 grouped two species, *T. bergquistae* from Australia and *T. erici* sp. nov. from the North Atlantic. Clade 3 comprised *T. aurantium*, *T. simoni* sp. nov., and an unidentified specimen of *Tethya* from the Red Sea (Fig. 3). All specimens of *T. aurantium* were collected in the Mediterranean, and showed only two haplotypes, with only one mutational step between them (Fig. 3). For *T. simoni* sp. nov., three specimens were collected in Panama and three more in Salvador (Brazil), and all of them showed a single haplotype shared across their distribution range (Fig. 3). Clade 4 is the least supported clade with a bootstrap of 70%, and comprises *T. minuta* Sarà, Sarà, Nickel & Brümmer, 2001, *T. actinia*, *T. californiana*, and *T. irisae* Sorokin et al., 2019. *Tethya californiana* is represented by two haplotypes with a 3 bp difference (Fig. 3). Clade 5 (Fig. 3) comprised the North Atlantic and Mediterranean species *T. citrina*, *T. norvegica*, and *T. hibernica*.

In the 28S rRNA tree, there is a wider coverage of genera within the family Tethyidae thanks to sequences available in GenBank, including species of the genera *Tethytimea*, *Tectitethya*, *Stellitethya*, *Xenospongia*, and *Tethya* which mostly clustered together in an unsupported clade, diverging the closest to the root within Tethyida (Fig. 4). The sister clade to the general *Tethya* clade was composed of *Tethytimea carmelita* Cruz-Barraza, Vega, Ávila & Vázquez-Maldonado, 2017, and an unidentified *Timea* specimen (Fig. 4). The four clades within Tethyidae observed in the COI tree were not recovered with 28S rRNA data, but it is important to note that many species sequenced for COI were not sequenced for 28S rRNA, so these two trees are not directly comparable. Within the genus *Tethya*, generally, the relationships between the species obtained with COI were mirrored with 28S rRNA (Fig. 4), with a few exceptions: *T. hibernica* did not cluster with *T. norvegica* but with *T. citrina* (Fig. 4).

Microbial composition

Diversity descriptors. The microbial communities were studied in 33 samples. The resulting sequencing reads ranged from 38,288 to 155,303, and the average number of unique amplicon sequence variants (ASVs) was

2,723 ± 1382 per sample (Supplemental Table S1). *Tethya citrina* i11 (Supplemental Table S1) included the highest number of ASVs (5552), while *T. orioni* sp. nov. had the lowest number of ASVs (461). The most abundant ASV had 69,754 reads in *T. actinia* i3, but in general, only around 2% of the ASVs showed more than 500 reads per sample. Diversity (Shannon H) of samples was not statistically different for sponge clades in general and in any of the pair-wise comparisons ($p > 0.05$, Supplemental Fig. S7A), but there were differences between some sponge species pairs ($p < 0.05$, Supplemental Table S2). Interestingly, the ‘white’ morphotype of *T. aurantium* i9 from Naples was the most diverse sample (5.32, Supplemental Table S1), even when compared with other specimens from the same species and the same location. As expected, *T. orioni* sp. nov. showed the lowest diversity (1.42), dominated by one ASV that comprised 71.5% RA and was annotated as *Synechococcus* CC9902 (100% similarity). Despite these differences in diversity, all 33 *Tethya* specimens whose microbiome was investigated here can be considered LMA sponges.

The ordination of samples based on Bray–Curtis dissimilarity (Supplemental Fig. S7B), showed a grouping of samples by clade, except for *T. norvegica* specimens that were distant to the clade centroid. These latter samples were collected in deep-sea environments and may explain why they did not cluster with their designated clade. Pairwise PERMANOVAs revealed differences between all pairs of clades, which was the result of both differences in location and dispersion of Bray–Curtis dissimilarities (Supplemental Table S1).

Microbial taxonomy. The samples analysed contained from 11–19 different microbial phyla (Supplemental Table S2). In all species, the dominant phylum was Proteobacteria (from 23.8–98.5% RA), except for *T. orioni* sp. nov., where Proteobacteria accounted only for 10% RA and Cyanobacteria dominated its microbiome (86.9%). Cyanobacteria were also abundant in *T. gracilis* from Brazil (35%) and *T. martini* sp. nov. (27 and 32.7% RA), but were below 12% in the rest of the samples. The archaeal community reached important relative abundance in the species *T. erici* sp. nov. (up to 13.7%) and *T. citrina* (up to 32.4%) (Supplemental Table S3). Also, samples of *T. erici* sp. nov. presented an unusually high abundance of sequences without a confident taxonomic classification beyond the kingdom Bacteria (up to 38.7%), for both ‘la Palma’ and ‘Chipiona’ samples. Since all samples were processed similarly, the reasons for this difference are unclear but could indicate the presence of bacterial taxa that are not well represented in the current databases.

At the deeper level of order, the microbial community of *Tethya simoni* sp. nov. and *T. aurantium* (both comprising clade 3) was mostly formed by Microtrichales (Actinobacteriota), Rhodobacterales (Alphaproteobacteria), UBA10353 marine group (Gammaproteobacteria), and other unclassified Proteobacteria (Supplemental Table S2, Fig. 7A). *Tethya citrina* harboured mostly Burkholderiales (Gammaproteobacteria) and Nitrosopumilales (Crenarchaeota). However, in *T. norvegica*, which also belonged to clade 5, Pirellulales were more abundant (reaching up to 54.4% RA), and Parvibaculares (Alphaproteobacteria, up to 43.9%), while Nitrosopumilales and Burkholderiales were less abundant. The three species belonging to clade 1 were characterized by the dominance of Synechococales (Cyanobacteria), with some differences in other taxonomic orders such as high abundances of

Rhodobacterales (Alphaproteobacteria) in *T. gracilis*, or high abundance of SAR11 clade (Alphaproteobacteria) in *T. martini* sp. nov. Finally, clade 2, represented by *T. erici* sp. nov. in our analysis, presented similar dominant taxonomic orders as the rest of *Tethya* species but with different relative abundances, and particular abundance of Cytophagales (Bacteroidota) (Supplemental Table S2).

The sample clustering based on abundances at ASV level was congruent with the clades of the COI phylogeny (Figs 3 and 7A, Supplemental Fig. S8), although the topology showed small variations in some relationships. For instance, *T. gracilis* clustered with *T. martini* sp. nov., instead of with *T. orioni* sp. nov. in the microbial clustering (Fig. 7A). However, it is important to note here that the relationships within these clusters can slightly differ depending on the taxonomic resolution

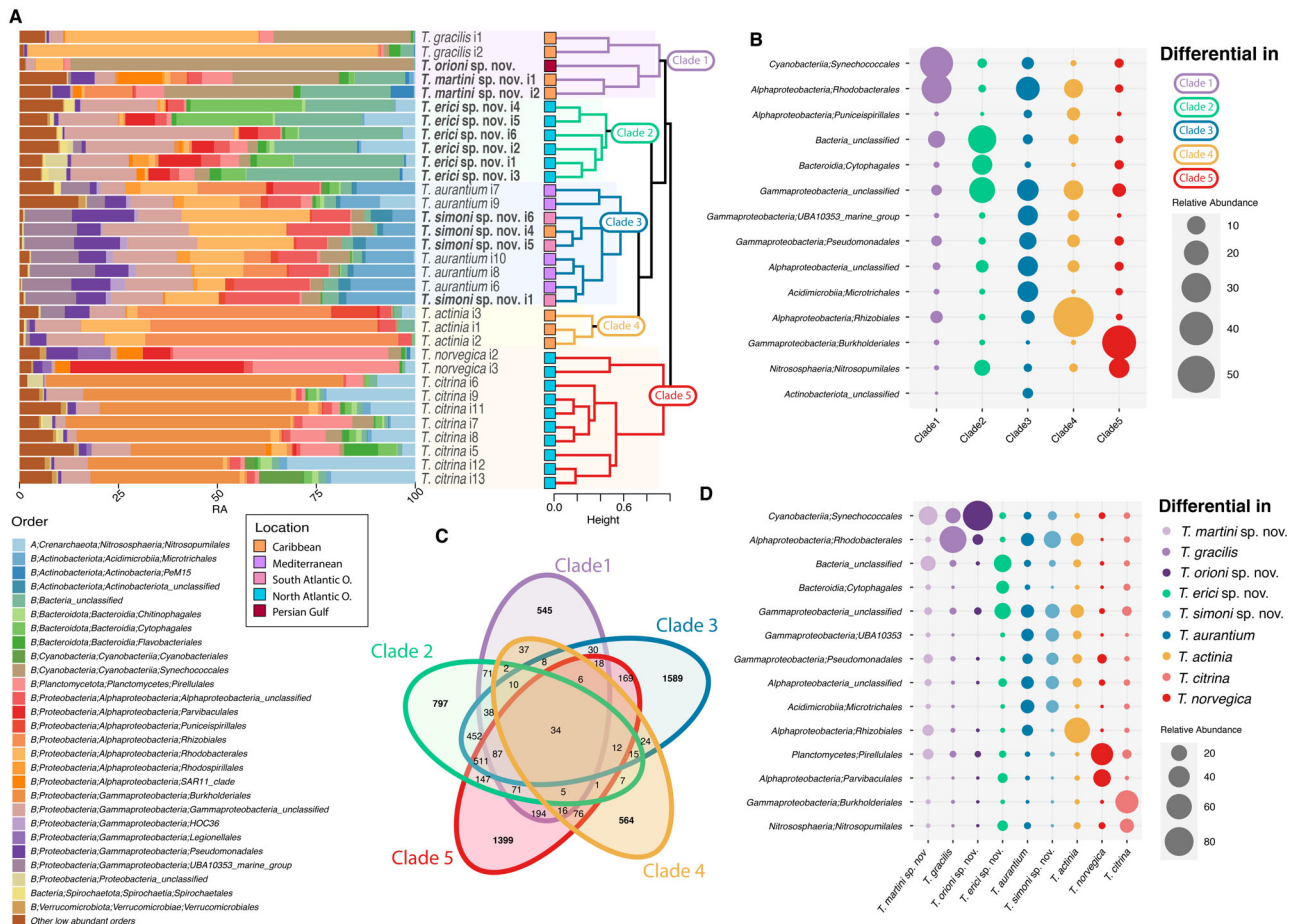


Fig. 7. Microbiome composition for *Tethya* spp. A. Barplot showing the relative abundance of each microbial (Bacteria and Archaea) order with corresponding dendrogram clustering of the samples using Bray-Curtis dissimilarity values. B. Differentially abundant microbial orders between sponge species that represent > 10% RA. Highlighted orders include the ones that are significantly significant in any of the pairwise comparisons (see Supplemental Table S6). C. Venn diagram of shared and unique ASVs for the four clades sequenced. D. Differentially abundant orders that are significant for all pairwise comparisons between one clade and the other three (see Supplemental Table S5). Circles of ‘diagnostic’ microbial taxa show a black stroke in both B and D.

used for the microbial composition. For example, in a dendrogram based on the genus level abundances, clade 2 branches as a sister group of clade 3, a pattern more similar to the COI tree (Supplemental Fig. S8).

Core microbiome and differentially abundant orders.

Only two core ASVs were shared among 70% of the samples: one *Synechococcus* and one *Rubripirellula* (Planctomycetes). This core reached a relative abundance ranging from 0 in *T. norvegica* to 17.8% in *T. gracilis* from Brazil (Supplemental Table S5). In pairwise comparisons, the largest number of shared ASVs occurred intraspecifically in samples of *T. citrina* (up to 1171 ASVs) and *T. erici* sp. nov. individuals (up to 816 ASVs) (Supplemental Table S6).

Across geographic areas, *Tethya simoni* sp. nov. shared 427–624 ASVs in samples from Brazil and Bocas del Toro; *T. citrina* from la Palma, Galicia, and Roscoff shared from 47–463 ASVs; *T. erici* sp. nov. from la Palma shared from 177–503 ASVs with the samples from Chipiona. Across species, *T. erici* sp. nov. shared many ASVs with *T. citrina* samples, with the highest values found among samples of both species collected in the same area (las Palmas) (from 327–726). Interestingly, the *T. aurantium* ‘white’ phenotype also shared high numbers of ASVs with these *T. erici* sp. nov. and *T. citrina* samples (up to 630 ASVs) (Supplemental Table S1).

Among clades (i.e., considering the mean of all microbial sequences present in any sample belonging to the clade), the core microbiome included 37 ASVs, spanning different phyla (Fig. 7C and Supplemental Table S5), and the specific ASVs of each clade ranged between 545 (11.5% of ASVs in clade 1) to 1589 (15.3% of ASVs in clade 3, Fig. 7C). These exclusive ASVs and other taxonomic orders that differed in abundance in one clade compared with the other four clades can be considered indicator taxa of the clade, and are shown in Fig. 7B. For instance, clade 1 presented larger abundances of Synechococcales in all pairwise comparisons, and Rhodobacterales in the comparison between clades 1 and 5. Clade 2 displayed larger abundances of Cytophagales, and, as commented above, a significant proportion of unclassified bacterial sequences. Clade 3 was characterized by Microtrichales, UBA10353 marine group, Pseudomonadales, and unclassified Alphaproteobacteria. Clade 4 distinctively harboured high proportions of Rhizobiales, and clade 5 of Burkholderiales and Nitrosopumilales. Other microbial taxonomic orders that were statistically distinct in some pairwise comparisons among clades can be found in Supplemental Table S7 and Supplemental Fig. 7C. Furthermore, some taxonomic orders were specific to the sponge species but not the

entire clade. This situation is noticeable in clade 5, where Burkholderiales and Nitrosopumilales are representative of *T. citrina*, but Pirellulales and Parvibaculales are characteristic of *T. norvegica* (Fig. 7D). Similarly, Rhodobacterales was abundant in *T. gracilis*, while Synechococcales dominated *T. orioni* sp. nov. (Fig. 7D). Additional orders with less than 10% RA can be found in Supplemental Table S8 and Supplemental Fig. S7D.

Discussion

Phylogeny and biogeography

In this study we describe four new species in the genus *Tethya*, two from the Western Atlantic (southern Caribbean and Brazil), one from the North Atlantic, and one from the Persian Gulf, by integrating morphology, genetics, and their associated microbial communities. We revisited the phylogeny of this group by adding new sequence data from freshly collected material, museum specimens, and unpublished data. Our revised phylogeny retrieved the main clades found in previous studies (Heim et al., 2007; Heim & Nickel, 2010; Sorokin et al., 2019), but incorporated samples representing new biogeographic regions. Interestingly, in our phylogenetic hypothesis, most clades and subclades do not show a clear biogeographic affinity, as they grouped specimens inhabiting disparate regions (Fig. 3). The weak biogeographic concordance uncovered in this study could be explained by an early divergence of Tethyidae and its cosmopolitan genus *Tethya*. According to recent time-calibrated phylogenies, this divergence has been estimated to have occurred during the late Permian/Triassic period (260–210 Ma) (Pankey et al., 2022; Plese et al., 2021), although the only fossil record for *Tethya* is from the Eocene (Łukowiak, 2016). Our results also support the idea of the early radiation in this genus proposed by Sarà and Sarà (2004) but suggest that this may have taken place earlier than previously thought (Heim, Nickel, & Brümmer, 2007). The origin of the main lineages may have occurred in the context of an ancient Palaeo-Tethys fauna (late Permian/Triassic) and subsequent radiation throughout the Mesozoic linked to the break-up of Gondwana. Therefore, the most plausible explanation for the current diversity and distribution of Australian and Caribbean *Tethya* species within the same clades is that they have deep roots and display a relict Tethyan pattern (Heim et al., 2007; Sarà & Sarà, 2004). This hypothesis is further supported by the sympatric occurrence of species belonging to distant evolutionary clades. To the classic example of *T. aurantium* (clade 2) and *T. citrina* (clade 4) from the Mediterranean (Sarà, 1990), we add that the new species

from the southern Caribbean, *T. martini* sp. nov. (clade 1), *T. simoni* sp. nov. (clade 3), and *T. actinia* (clade 4) show also a high genetic distance, belonging to quite divergent clades of COI (Fig. 3).

The early 20th century was pervaded by descriptions of *Tethya* species widely distributed across all oceans, such as *T. aurantium*, *T. diploderma*, or *T. seychellensis*. The cosmopolitanism of these species has been previously refuted (Ribeiro & Muricy, 2011), and our results provide further evidence of this. The extreme similarities in morphology and close molecular affinity of *T. simoni* sp. nov. and *T. aurantium* make them easy to be confused, and thus researchers assigned them to the same species. Our results show, in turn, that the once-thought-widespread *T. aurantium* might be restricted to the Mediterranean Sea or at least the NE Atlantic, as suggested by Heim *et al.* (2007). These species from both sides of the Atlantic have a high genetic distance (two distinct COI haplotypes with eight mutations, Fig. 3), higher than in other sibling Porifera species (e.g., Cárdenas *et al.*, 2013; Duran & Rützler, 2006).

Another interesting finding in our study is that two specimens from the southern Caribbean and the South Atlantic coast in Brazil were identified morphologically and genetically as *T. gracilis*, a species originally described from German aquaria. Our data also suggest a Western Atlantic origin for this species, refuting an Indo-Pacific origin for *Tethya gracilis*, as previously proposed based on a morphological resemblance to *T. seychellensis* (Sarà *et al.*, 2001) and its grouping within the same genetic clade (Heim, Nickel, & Brümmer, 2007). Establishing the origins of the three species described from German aquaria, *T. wilhelma*, *T. minuta*, and *T. gracilis* (see Sarà *et al.*, 2001), is relevant because of their use in experimental studies on kinetics, functional morphology, and biomechanics in clades of early-branching metazoans (Ellwanger & Nickel, 2006; Nickel *et al.*, 2006). The affinity of *T. minuta* with *T. actinia* described from Bermuda (Heim *et al.*, 2007) tilts the balance towards an Atlantic provenance of two of the species of *Tethya* from aquaria, leaving only *T. wilhelma* as the one possibly coming from the Eastern Mediterranean Sea (Sorokin *et al.*, 2019). Indeed, the sequence from *Tethya* sp. (KX866754), originally identified as *T. aurantium* (see Idan *et al.*, 2018) is strikingly similar to that of *T. wilhelma*, with only 1 bp difference in COI, suggesting that the origin of *T. wilhelma* might be the Eastern Mediterranean or the Red Sea. Since the type localities of the three species are three different public aquaria (Stuttgart, Karlsruhe, and Düsseldorf), it is likely that they may have originated from different geographic regions. Although less probable, an alternative scenario to the Atlantic origin for *T. gracilis* could

be its recent introduction from the Indo-Pacific via ballast water or biofouling on ship's hulls (e.g., *Mycale grandis* Shih & Popp, 2020). However, previous records of *T. seychellensis* in the Colombian Caribbean could correspond to *T. gracilis* (e.g. Wintermann-Kilian & Kilian, 1984) and the high diversity of *Tethya* species in Brazil would support that *T. gracilis* is most likely native to these regions.

The addition of species from *Tethya* hotspots such as Australia, Brazil, the Malay Archipelago, and the Galápagos islands in future phylogenetic analyses will bring more insights to the understanding of the evolution of this diverse genus. Although in this study we attempted to include some of the Brazilian species using subsamples from the MNRJ in Brazil, we were unsuccessful in recovering sequences due to low-quality DNA preservation. Further studies may consider the inclusion of museum material through the implementation of mini-barcodes specifically designed to amplify partial COI sequences from degraded DNA (Cárdenas & Moore, 2019) or the use of short-read sequencing technologies, which have proved much more efficient in dealing with degraded museum samples (Derkarabetian *et al.*, 2019; Srivathsan *et al.*, 2021).

Systematics

Taxonomists, systematists, and field ecologists strive for easily observed diagnostic features with high discrimination power. Among the morphological features with the potential to become diagnostic for systematics, colour has been proposed as a reliable character to support some of the clades in *Tethya* (Sorokin *et al.*, 2019). Clades 1 and 2 include red, pink or white species, clade 3 are mainly orange, while clade 4 is composed of mainly yellow species. Our newly described species bring more evidence to this hypothesis, and, as expected, the bright red *T. martini* sp. nov. groups in clade 1 with *T. seychellensis* and *T. coccinea* (all red) and the orange *T. simoni* sp. nov. is part of clade 3 with *T. aurantium* (both orange). However, there are some exceptions to these trends. In fact, monitoring studies of *Tethya* spp. in New Zealand have shown that the same sponge can experience variations in shape and colouration depending on seasonal environmental changes (Shaffer *et al.*, 2020). In any case, the colour-based hypothesis in *Tethya* species demands further investigation as this character could be genetically controlled, since it may be related to the presence of carotenoids (Tanaka *et al.*, 1982; Tanaka & Yamamoto, 1984) produced by the sponges (Liaen-Jensen *et al.*, 1982) or symbionts, unicellular green algae, or Cyanobacteria (Belikov *et al.*, 2019).

The usual morphological diagnostic characters of spicule morphologies in *Tethya* are especially problematic, due to the lack of a homogeneous terminology for the genus. This has been a persistent issue throughout the taxonomic history of *Tethya* (Bergquist & Kelly-Borges, 1991). Therefore, it is important to emphasize that most-needed phylogenetic analyses within this species-rich genus first require the unification of terminology for spicule types based on quantitative morphometric measurements. For instance, an effort to encompass the wide variation of micrasters has resulted in the distinction of three main types, i.e., tylaster, strongylaster, and oxyaster (~previously pooled in chiasters, although most referred to tylasters and strongylasters), and subtypes within (e.g. Ribeiro & Muricy, 2011 used numbers for subtypes, while Hajdu et al., 2013 added the prefix acanthose). Yet their definitions were rather qualitative, leaving room for some intermediate forms of difficult designation. Therefore, the redescription of type material under such a unified morphological framework would facilitate mapping characters onto phylogenetic trees. Finally, identifying new and reliable external characters and from histological sections, e.g., tubercle shape, lacunar structure, and collagenous distribution could help to assign species to their taxonomic units. Although in our study we followed the nomenclature of spicules by Ribeiro and Muricy (2011) to allow comparisons with other Western Atlantic species, we recommend that a comprehensive nomenclature framework should also take into account the definitions proposed by Bergquist and Kelly-Borges (1991), Sarà and Sarà (2004) and Heim, Nickel, and Brümmer (2007), to cover the wide spectra of spicule variation across the different biogeographic regions.

Microbial composition

Integrative taxonomy aims to incorporate several lines of evidence that converge in robust hypotheses for species delimitation. In the field of sponge systematics, integrative taxonomy mainly includes morphological and molecular characters (Dohrmann et al., 2017; Riesgo et al., 2018; Sorokin et al., 2019). In recent years, several authors have tried to also incorporate metabolomics (Reveillaud et al., 2012; Ruiz et al., 2015) and ecological/biogeographic data (e.g., Azevedo et al., 2015). Surprisingly, the microbiome has never been formally used as a taxonomic character, despite its remarkable species-specificity (Thomas et al., 2016) and studies hinting at cophylogeny/coevolution patterns with their hosts (Pankey et al., 2022; Schöttner et al., 2013). Coevolution or phyllosymbiotic patterns in marine invertebrates are especially conspicuous in sponges and

cnidarians, which harbour complex communities of prokaryotic symbionts (O'Brien et al., 2019). In sponges, the host is usually the factor driving most divergences in the coevolutionary patterns (Thomas et al., 2016), with those being more pronounced at high taxonomic levels, such as order, and less clear in LMA than in HMA sponges (Pankey et al., 2022). Our analysis shows high levels of coevolution in the host and microbial communities of LMA sponges, even within a single genus of sponges. Nonetheless, small discrepancies can be observed at different levels of microbial taxonomy, both in our study and across the literature. Thus, the microbiome emerges as an informative intrageneric diagnostic character with enormous potential for taxonomic identification and evolutionary studies, especially for species in which COI is extremely conserved (for instance Riesgo et al., 2016). In fact, in our study, although the clustering analysis of the prokaryotic communities at ASV level only correlated moderately with the phylogenetic hypothesis obtained with COI (Figs 3 and 7A, Supplemental Fig. S8), the distinctive microbial communities for the different clades suggest that these could be considered an additional taxonomic feature to aid in *Tethya* systematics. This high specificity shown by *Tethya* in their microbial communities was previously proposed by Sipkema and Blanch (2010) as an alternative approach to differentiate between species. Interestingly, once *T. norvegica* was removed from the microbial analysis, the correlation between the clustering analysis of the microbiome and the COI tree improved (Supplemental Fig. S8). In fact, the microbiome of *T. norvegica* was very different from that of the rest of specimens in Clade 5 (*T. citrina*) and from those in other clades, likely because it is a deep-sea species, which typically hosts a distinctive microbiome (Díez-Vives & Riesgo, 2024).

The microbiome of the previously studied species of *Tethya* is dominated by temporally stable Proteobacteria species, usually represented by a single, dominant operational taxonomic unit (OTU or ASV in our case), followed by Cyanobacteria and Bacteroidetes (Astudillo-García et al., 2020; Sipkema & Blanch, 2010; Thiel et al., 2007; Waterworth et al., 2017). Although our results can hardly be compared with these previous studies because the primers used for the amplification of the 16S rRNA gene were different, an overall gross comparison also points to the dominance of Proteobacteria (Fig. 7). These phyla are also typically dominant in other LMA sponges (Moitinho-Silva, Steinert, et al., 2017). In our study, Cyanobacteria were present across all species but dominant in clade 1, with the order Synechococcales distinctive of the group. Interestingly, the microbiome of *T. burtoni*, which belongs to clade 1 (Fig. 3), was sequenced

by other researchers and did not show a dominance of Cyanobacteria (Astudillo-García *et al.*, 2020) but, in this case, the primers used could have had an amplification bias different from ours. Several other microbial taxa were characteristic of each of the clades we recovered. For instance, only *Tethya* species from clade 3 have abundant communities of Actinobacteria (in our case of order Microtrichales), which were also recovered from other *T. aurantium* samples (Thiel *et al.*, 2007) and the species *T. californiana* (Sipkema & Blanch, 2010), which clusters also in clade 3 according to our phylogenetic results. Similarly, an unidentified species of *Tethya* collected on Orpheus Island in Australia had Actinobacteria dominating the community, together with Proteobacteria, Chloroflexi, Crenarchaeota, and Verrucomicrobia (Thomas *et al.*, 2016). However, whether this unidentified species belonged to clade 3 is currently unknown. *Tethya citrina* (clade 5) was remarkably dominated by Burkholderiales (Gammaproteobacteria). This group is hard to track in the literature because it has been reorganized differently in the available databases; it was formerly known as class Betaproteobacteria, but can also be found as an order placed within class Betaproteobacteria, or as class Gammaproteobacteria (i.e., NCBI and SILVA taxonomy) (Parks *et al.*, 2018). Recently, Taylor *et al.* (2021) characterized this group classifying it as a new gammaproteobacterial order named *Candidatus Tethybaerales*. Only a Metagenome-Annotated Genome (MAG) annotated as *Beroebacter blanensis*, and assembled from *Crambe crambe* (Schmidt, 1862) from the Mediterranean Sea, could be detected among 14 of our samples covering all species (100% sequence identity), although in low abundances (up to 0.007% RA). In summary, although the use of the microbiome as a diagnostic character has only been tentatively explored here, it holds enormous potential for future taxonomic studies, especially in groups where the morphology and the molecular characters are highly homoplastic.

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Supplemental material

Supplemental material for this article can be accessed here: <http://dx.doi.org/10.1080/14772000.2024.2383341>.

Disclosure statement

No potential conflict of interest was reported by the author(s).










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