

Molecular phylogenies challenge the classification of Polymastiidae (Porifera, Demospongiae) based on morphology

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Abstract Polymastiidae Gray, 1867 is a worldwide distributed sponge family, which has a great significance for understanding of the demosponge deep phylogeny since the former order Hadromerida Topsent, 1894 has been recently split based on the molecular evidence and a new separate order has been established for the polymastiids. However, molecular data obtained from Polymastiidae so far are scarce, while the phylogenetic reconstruction based on morphology has faced a deficit of characters along with the vagueness of their states. The present study is a phylogenetic reconstruction of Polymastiidae based on novel data on two molecular markers, cytochrome oxidase subunit I and large subunit ribosomal DNA, obtained from a broad set of species. Monophyly of the family and nonmonophyly of four polymastiid genera are revealed, suggesting a high level of homoplasy of morphological characters, which are therefore not an appropriate base for the natural classification of Polymastiidae. Although the presented phylogenies cannot yet provide an alternative

classification scheme, several strongly supported clades, which may be used as reference points in future classification, are recovered and three taxonomic actions are proposed: transfer of one species from *Radiella* to *Polymastia* Bowerbank, 1862; transfer of three species from *Radiella* Schmidt, 1870 to *Spinularia* Gray, 1867; and the consequent abandonment of *Radiella*.

Keywords Phylogeny · Homoplasy · Polymastiidae · CO1 · 28S rDNA

Introduction

Polymastiidae Gray (1867), with its 122 species from 15 genera and a worldwide distribution (Van Soest et al. 2015), is one of the key families in Demospongiae Sollas, 1885, the most diverse class of sponges. At the same time, Polymastiidae is one of the problematic taxa with a controversial classification (Plotkin et al. 2012). Classification of the demsponges has traditionally been based on the shape and arrangement of their skeletal elements, i.e., mineral spicules and organic fibers (Hooper and Van Soest 2002). The polymastiids possess a relatively simple spicule assortment providing a rather scant set of taxonomic characters (Plotkin et al. 2012). Polymastiidae comprises sponges of various body shapes, often bearing papillae and possessing a skeleton mainly composed of smooth monactines (Boury-Esnault 2002; see terminology of the sponge morphology in Boury-Esnault and Rützler 1997). Based on the latter feature, this family was until recently affiliated with the demosponge order Hadromerida Topsent, 1894. For the moment, only one morphological feature delimiting Polymastiidae from other demsponges is usually defined, the presence of a superficial cortical palisade made of spicules differing from those composing the

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choanosomal tracts in size and/or in shape (Boury-Esnault 2002; Plotkin and Janussen 2008). However, this feature is in fact also displayed by some taxa from other families, e.g., by *Aaptos* Gray, 1867 belonging to the Suberitidae Schmidt, 1870 (Plotkin et al. 2012).

Discrimination between polymastiid genera is based on the body shape (e.g., radial body in *Radiella* Schmidt, 1870 and columnar body in *Tentorium* Vosmaer, 1887), the architecture of the choanosomal skeleton (diffuse skeleton in *Quasillina* Norman, 1869 and *Ridleia* Dendy, 1888, reticulate skeleton in *Weberella* Vosmaer, 1885 and radial skeleton in the remaining 12 genera) and the presence of spicules other than the ordinary smooth monactines in the choanosome (in four genera), or in the cortex (in five genera) (Boury-Esnault 2002). However, in some cases, these characters are inconsistent. For example, *Polymastia* Bowerbank, 1862 is usually defined as sponges with a radial choanosomal skeleton and smooth monactines constituting both the choanosomal and cortical skeleton (Boury-Esnault 2002) even though several species traditionally affiliated with *Polymastia* display a reticulate skeleton (Plotkin et al. 2012) or extraordinary spicules in the choanosome or in the cortex (Kelly-Borges and Bergquist 1997). Other characters used in the taxonomy of polymastiids include the number of size categories of the ordinary monactines and the minute differences in their shape, the presence and architecture of additional cortical layers, and the anatomy of the papillae (Boury-Esnault 1987, 2002; Kelly-Borges and Bergquist 1997; Morrow and Boury-Esnault 2000; Plotkin and Janussen 2008). These characters are often unstable and provide poor taxonomic information (Plotkin et al. 2012). Particularly, they fail to discriminate between some morphologically similar polymastiids, which inhabit the polar and temperate waters of the northern and southern hemispheres, but do not occur in the tropics, and consequently, a bipolar distribution is presumed for these species (e.g., for *Tentorium semisuberites* (Schmidt 1870) and *Radiella sarsi* (Ridley & Dendy, 1886)).

Phylogenetic reconstruction of Polymastiidae based on 25 binary morphological characters (Plotkin et al. 2012) questioned the monophyly of the family, with *Pseudotrachya hystrix* (Topsent, 1890) not grouping with any other polymastiid and one of the outgroup species, *Aaptos papillata* (Keller, 1880), joining the main polymastiid clade, and demonstrated that *Polymastia* is polyphyletic. At the same time, molecular phylogenies of Polymastiidae have been never properly reconstructed. Until now, common phylogenetic markers as the barcoding regions of cytochrome oxidase subunit I (CO1) and the partial RNA from the large and small ribosomal subunits (28S and 18S) have only been obtained for a small number of polymastiid species, aiming to resolve a deep phylogeny of the class Demospongiae instead of addressing the relationships within Polymastiidae (Nichols 2005; Morrow et al. 2012, 2013; Redmond et al. 2013; Vargas

et al. 2015). In all phylogenies resulting from these studies, *Polymastia* was nonmonophyletic, while the family Polymastiidae was monophyletic excluding two species from Nichols (2005). Furthermore, in all molecular phylogenies of Demospongiae, Polymastiidae and other hadromerid families appeared in remote clades that seriously contradicted the traditional classification based on their morphological similarities. Very recently, based on the molecular data, Morrow and Cárdenas (2015) proposed abandoning the order Hadromerida and establishing five new orders for the former hadromerids, with the order Polymastiida including only one family, the Polymastiidae. This proposal highlights the importance of the polymastiids in the context of the deep phylogeny of demosponges.

The purpose of the present study was to reconstruct the phylogeny of the family Polymastiidae based on two broadly used molecular markers, the 5'-end barcoding region of CO1 (Folmer et al. 1994) and a large region of 28S rRNA (helix B10 to helix E19, numeration of the helices according to De Rijk et al. (1999, 2000) and Wuyts et al. (2001)), employing a much larger set of polymastiid species than ever studied before. We also tested the monophyly of the family as well as the monophyly of its genera and traced the evolution of morphological characters along the branches of the consensus molecular tree.

Material and methods

Sampling and taxonomic identification

Eighty-seven polymastiid individuals were collected for our study and deposited in the natural history collections of four museums (see Table 1 for details). Both the individuals in toto assigned for morphological examination and the choanosomal pieces of about 1 cm³ for DNA extraction were fixed in 95–100 % ethanol. Sponge anatomy was examined under a light microscope on 500–700- μ m-thick sections prepared using a precise saw with a diamond wafering blade after embedding of tissue pieces in epoxy resin. Isolated spicules were examined under a light microscope and SEM. Preparations and subsequent taxonomic identification followed well-known routines for polymastiids (Boury-Esnault 1987; Boury-Esnault and Bézac 2007; Plotkin and Janussen 2007, 2008).

Taxonomic scope

In our study, we included genetic data on 24 unambiguously identified species and ten operational taxonomic units (OTUs), of which four were identified to species level with some uncertainty and six could not be referred to any known species and were therefore only identified to genus level (Table 1). These species and OTUs belonged to seven

Table 1 List of specimens used in this study with museum voucher numbers, GenBank accession numbers and localities. New sequences from our study are highlighted in italics. Asterisks before accession numbers indicate the sequences resulting from cloning of the PCR products. When all clones from one product were identical, only one sequence was submitted. A range of accession numbers in one cell indicates a library of nonidentical clones. Abbreviations of the museums: BELUM Ulster Museum, Belfast, UK; GNM Gothenburg Natural History Museum, Sweden; SMF Senckenberg Naturmuseum, Frankfurt am Main, Germany; UCMPPWC Museum of Paleontology, University of California, USA; ZMBN Natural History Collections, University Museum of Bergen, Norway

Species	Type species	Museum voucher	28S: B10-C1	28S: D1-D19	28S: D20-E19	COI	Locality	Type locality
<i>Polymastia andrica</i> de Laubenfels, 1949	ZMBN 98055	LN873411		*LN873461	LN873424	LN606449	Arctic Ocean, Norway, Svalbard	
<i>Polymastia andrica</i> de Laubenfels, 1949	ZMBN 98057	HG423736		HG423766	HG423796	HG423707	North Sea, Norway, Hordaland	
<i>Polymastia andrica</i> de Laubenfels, 1949	ZMBN 98074	LN873415		*LN873473- LN873477	LN873428	LN606453	Norwegian Sea, Norway, Troms	
<i>Polymastia andrica</i> de Laubenfels, 1949	ZMBN 98102	-		-	-	LN606454	Atlantic Ocean, Canada, Newfoundland	
<i>Polymastia andrica</i> de Laubenfels, 1949	ZMBN 98108	-		-	-	LN606455	Norwegian Sea, Norway, offshore	
<i>Polymastia arctica</i> (Merejkowsky, 1878)	ZMBN 98060	LN873412		*LN873465- LN873468	LN873425	LN606450	White Sea, Russia, Karelia	Yes
<i>Polymastia arctica</i> (Merejkowsky, 1878)	ZMBN 98062	HG423734		HG423764	HG423794	HG423705	White Sea, Russia, Karelia	Yes
<i>Polymastia arctica</i> (Merejkowsky, 1878)	ZMBN 98063	LN873413		*LN873469- LN873470	LN873426	LN606451	White Sea, Russia, Karelia	Yes
<i>Polymastia arctica</i> (Merejkowsky, 1878)	ZMBN 98065	LN873414		*LN873471- LN873472	LN873427	LN606452	Norwegian Sea, Norway, Finnmark	
<i>Polymastia arctica</i> (Merejkowsky, 1878)	ZMBN 98068	HG423735		HG423765	HG423795	HG423706	Norwegian Sea, Norway, Finnmark	
<i>Polymastia bartletti</i> de Laubenfels, 1942	ZMBN 98111	LN606505		LN606535	LN606565	LN606468	Atlantic Ocean, Canada, Nova Scotia	
<i>Polymastia cf. bartletti</i> de Laubenfels, 1942	GNM 904:1	HG423738		HG423768	HG423798	*LN606467	North Sea, Sweden, Kattegat	
<i>Polymastia boletiformis</i> (Lamarck, 1815)	BELUM: MC5014	HQ379232		HQ379306	HQ379372	-	N/A	
<i>Polymastia boletiformis</i> (Lamarck, 1815)	GNM 901:1	LN606496		LN606526	LN606556	-	North Sea, Sweden, Skagerrak	
<i>Polymastia boletiformis</i> (Lamarck, 1815)	ZMBN 98047	LN606491		LN606521	LN606551	*HG423708	North Sea, Norway, Hordaland	
<i>Polymastia boletiformis</i> (Lamarck, 1815)	ZMBN 98048	LN606492		LN606522	LN606552	-	North Sea, Norway, Hordaland	
<i>Polymastia boletiformis</i> (Lamarck, 1815)	ZMBN 98081	LN606493		LN606523	LN606553	-	North Sea, Norway, Hordaland	
<i>Polymastia boletiformis</i> (Lamarck, 1815)	ZMBN 98088	LN606494		LN606524	LN606554	-	North Sea, Norway, Vest-Agder	
<i>Polymastia boletiformis</i> (Lamarck, 1815)	ZMBN 98089	LN606495		LN606525	LN606555	*HG423709	North Sea, Norway, Vest-Agder	
<i>Polymastia cf. conigera</i> Bowerbank, 1874	BELUM: MC3722	HG423828		HG423828	HG423830	HG423827	Celtic Sea, Ireland, Co. Cork	
<i>Polymastia coriicata</i> Ridley & Dendy, 1886	ZMBN 98097	*LN873417		LN873445	LN873430	LN606456	Atlantic Ocean, Canada, Newfoundland	
<i>Polymastia coriicata</i> Ridley & Dendy, 1886	ZMBN 98104	*LN873452		HG423825	HG423826	HG423824	Atlantic Ocean, Canada, Newfoundland	
<i>Polymastia coriicata</i> Ridley & Dendy, 1886	ZMBN 98105	LN873418		LN873446	LN873431	LN873450	Atlantic Ocean, Portugal, Azores	
<i>Polymastia euplectella</i> Rezvoj, 1927	ZMBN 98044	HG423767		HG423767	HG423797	HG423710	Barents Sea, Norway, Finnmark	
<i>Polymastia euplectella</i> Rezvoj, 1927	ZMBN 98085	LN606497		LN606527	LN606557	LN606457	North Sea, Norway, Vest-Agder	
<i>Polymastia euplectella</i> Rezvoj, 1927	ZMBN 98086	LN606498		LN606528	LN606558	LN606458	North Sea, Norway, Vest-Agder	
<i>Polymastia euplectella</i> Rezvoj, 1927	ZMBN 98087	LN606499		LN606529	LN606559	-	North Sea, Norway, Vest-Agder	
<i>Polymastia grimaldii</i> (Topse, 1913)	ZMBN 98064	LN873419		*LN873478- LN873482	LN873432	LN606459	Barents Sea, Norway, Finnmark	
<i>Polymastia grimaldii</i> (Topse, 1913)	ZMBN 98110	-		-	-	LN606460	Atlantic Ocean, Canada, Newfoundland	
<i>Polymastia grimaldii</i> (Topse, 1913)	ZMBN 98112	-		-	-	LN606461	Barents Sea, Norway, offshore	
<i>Polymastia invaginata</i> Kirkpatrick, 1907	ZMBN 98046	HG423740		HG423770	HG423800	HG423712	Bellinghousen Sea, Antarctic Peninsula	
<i>Polymastia invaginata</i> Kirkpatrick, 1907	ZMBN 98093	LN606500		LN606530	LN606560	LN606462	Weddell Sea, Antarctic, South Shetland Islands	
<i>Polymastia invaginata</i> Kirkpatrick, 1907	ZMBN 98094	HG423739		HG423769	HG423799	HG423711	Weddell Sea, Antarctic, South Shetland Islands	
<i>Polymastia littoralis</i> Stephens, 1915	N/A	-		-	-	KJ129611	N/A	
<i>Polymastia littoralis</i> Stephens, 1915	N/A	-		-	-	NC 023834	N/A	
<i>Polymastia mamillaris</i> (Müller, 1806)	ZMBN 98078	HG423741		HG423771	HG423801	HG423713	North Sea, Norway, Hordaland	
<i>Polymastia mamillaris</i> (Müller, 1806)	ZMBN 98083	*LN873453		*LN873462	LN873439	LN606463	North Sea, Norway, Vest-Agder	
<i>Polymastia pachymastia</i> de Laubenfels, 1932	UCMPWC 932	AY561924		-	-	-	N/A	
<i>Polymastia penicillus</i> (Montagu, 1818)	BELUM: MC5284	HQ393895		HQ393899	HQ393903	-	N/A	
<i>Polymastia penicillus</i> (Montagu, 1818)	BELUM: MC6505	LN606501		LN606531	LN606561	LN606464	UK, Northern Ireland	
<i>Polymastia penicillus</i> (Montagu, 1818)	GNM 460:1	LN606502		LN606532	LN606562	-	North Sea, Sweden, Skagerrak	

Table 1 (continued)

Species	Type species	Museum voucher	28S: B10-C1	28S: D1-D19	28S: D20-E19	COI	Locality	Type locality
<i>Polymastia penicillus</i> (Montagu, 1818)		GNM 460-2	LN606503	LN606533	LN606563	—	North Sea, Sweden, Skagerrak	
<i>Polymastia thielei</i> Koltun, 1964		ZMBN 98052	LN873420	*LN873463	LN873433	LN606469	Arctic Ocean, Norway, Svalbard	Yes
<i>Polymastia thielei</i> Koltun, 1964		ZMBN 98053	—	—	—	LN873451	Arctic Ocean, Norway, Svalbard	Yes
<i>Polymastia thielei</i> Koltun, 1964		ZMBN 98070	—	—	—	LN606470	Norwegian Sea, NE Iceland	
<i>Polymastia thielei</i> Koltun, 1964		ZMBN 98107	—	—	—	LN606471	Norwegian Sea, East Greenland	
<i>Polymastia thielei</i> Koltun, 1964		ZMBN 98109	—	—	—	LN606472	Denmark Strait, East Greenland	
<i>Polymastia uberrima</i> (Schmidt, 1870)		ZMBN 98066	HG423744	*LN873483-873484	HG423804	HG423716	Norwegian Sea, Norway, Bjørnøya	
<i>Polymastia uberrima</i> (Schmidt, 1870)		ZMBN 98073	*LN873456-873458	HG423775	HG423805	HG423717	Norwegian Sea, Norway, Troms	
<i>Polymastia</i> sp. 1		ZMBN 98091	HG423742	HG423772	HG423802	HG423714	North Sea, Norway, Rogaland	
<i>Polymastia</i> sp. 1		ZMBN 98092	LN606504	LN606534	LN606564	LN606465	North Sea, Norway, Rogaland	
<i>Polymastia</i> sp. 2		ZMBN 98080	HG423743	HG423773	HG423803	HG423715	North Sea, Norway, Hordaland	
<i>Polymastia</i> sp. 3		ZMBN 98106	—	—	—	LN606466	Atlantic Ocean, Canada, Newfoundland	
<i>Quasillina brevis</i> (Bowerbank, 1862)	Yes	BELUM: MC6569	LN606510	LN606540	LN606570	—	Irish Sea, UK, England	
<i>Quasillina brevis</i> (Bowerbank, 1862)	Yes	ZMBN 98049	LN606506	LN606536	LN606566	LN606473	North Sea, Norway, Sognefjorden	
<i>Quasillina brevis</i> (Bowerbank, 1862)	Yes	ZMBN 98067	LN606507	LN606537	LN606567	LN606474	Denmark Strait, West Iceland	
<i>Quasillina brevis</i> (Bowerbank, 1862)	Yes	ZMBN 98082	LN606508	LN606538	LN606568	LN606475	North Sea, Norway, Rogaland	
<i>Quasillina brevis</i> (Bowerbank, 1862)	Yes	ZMBN 98084	HG423746	HG423776	HG423806	HG423718	North Sea, Norway, Vest-Agder	
<i>Quasillina brevis</i> (Bowerbank, 1862)	Yes	ZMBN 98090	LN606509	LN606539	LN606569	LN606476	North Sea, Norway, Vest-Agder	
<i>Radiella hemisphaerica</i> (Sars, 1872)		ZMBN 98043	HG423747	HG423777	HG423807	HG423719	North Sea, Norway, Hordaland	
<i>Radiella hemisphaerica</i> (Sars, 1872)		ZMBN 98056	HG423748	HG423778	HG423808	HG423720	North Sea, Norway, Hordaland	
<i>Radiella hemisphaerica</i> (Sars, 1872)		ZMBN 98058	—	—	—	LN606477	North Sea, Norway, Hordaland	
<i>Radiella hemisphaerica</i> (Sars, 1872)		ZMBN 98069	—	—	—	LN606478	Imvinger Basin, South Iceland	
<i>Radiella hemisphaerica</i> (Sars, 1872)		ZMBN 98071	—	—	—	LN606479	Barents Sea, Norway, Finnmark	
<i>Radiella hemisphaerica</i> (Sars, 1872)		ZMBN 98077	—	—	—	LN606480	North Sea, Norway, Hordaland	
<i>Radiella sarsi</i> (Ridley & Dendy, 1886)		ZMBN 98039	HG423749	*HG423779	HG423809	HG423721	Norwegian Sea, Norway, offshore	
<i>Radiella sarsi</i> (Ridley & Dendy, 1886)		ZMBN 98098	LN606512	LN606542	LN606572	LN606481	Norwegian Sea, Norway, offshore	
<i>Radiella cf. sarsi</i> (Ridley & Dendy, 1886)		ZMBN 98103	*LN873459- LN873460	*HG423780	*HG423810	*HG423722	Indian Ocean, Mozambique, Mozambique Channel	
<i>Radiella</i> sp.		ZMBN 98038	HG423751	HG423781	HG423812	HG423723	Norwegian Sea, Norway, offshore	
<i>Radiella</i> sp.		ZMBN 98040	LN606513	LN606543	LN606573	LN606482	Norwegian Sea, Norway, offshore	
<i>Radiella</i> sp.		ZMBN 98041	LN606514	LN606544	LN606574	LN606483	Norwegian Sea, Norway, offshore	
<i>Sphaeropylus antarcticus</i> Kirkpatrick, 1907		POR 21125	—	—	—	KC869424	N/A	
<i>Sphaeropylus antarcticus</i> Kirkpatrick, 1907		ZMBN 98045	HG423752	HG423782	HG423812	HG423724	Bellingshausen Sea, Antarctic, Antarctic Peninsula	Yes
<i>Sphaeropylus borealis</i> (Svarzewsky, 1906)		ZMBN 98036	HG423753	*HG423783	HG423813	—	White Sea, Russia, Karelia	Yes
<i>Sphaeropylus borealis</i> (Svarzewsky, 1906)		ZMBN 98059	HG423754	HG423784	HG423814	HG423725	White Sea, Russia, Karelia	Yes
<i>Sphaeropylus borealis</i> (Svarzewsky, 1906)		ZMBN 98061	LN606515	LN606545	LN606575	LN606484	White Sea, Russia, Karelia	Yes
<i>Sphaeropylus capitatus</i> (Vosmaer, 1885)	Yes	GNM 902	—	—	—	LN606486	North Sea, Sweden, Skagerrak	
<i>Sphaeropylus capitatus</i> (Vosmaer, 1885)	Yes	ZMBN 98042	*LN873454	LN873448	LN873443	LN606485	North Sea, Norway, Hordaland	
<i>Sphaeropylus capitatus</i> (Vosmaer, 1885)	Yes	ZMBN 98075	HG423755	HG423785	HG423815	HG423726	Norwegian Sea, Norway, Troms	
<i>Sphaeropylus</i> sp. 1		BELUM: MC4236	HQ379233	HQ379307	HQ379373	HQ379425	Atlantic Ocean, UK, Scotland	
<i>Sphaeropylus</i> sp. 2		BELUM: MC5015	*LN873455	HG423831	HG423832	*LN606487	Celtic Sea, Ireland, Co. Cork	
<i>Spinularia spinularia</i> (Bowerbank, 1866)	Yes	GNM 792-1	LN606517	LN606547	LN606577	—	North Sea, Sweden, Skagerrak	
<i>Spinularia spinularia</i> (Bowerbank, 1866)	Yes	ZMBN 98037	HG423756	HG423786	HG423816	HG423727	North Sea, Norway, Hordaland	
<i>Spinularia spinularia</i> (Bowerbank, 1866)	Yes	ZMBN 98050	LN606516	LN606546	LN606576	LN606488	North Sea, Norway, Sognefjorden	
<i>Spinularia spinularia</i> (Bowerbank, 1866)	Yes	ZMBN 98076	LN606518	LN606548	LN606578	LN606489	North Sea, Norway, Hordaland	
<i>Spinularia spinularia</i> (Bowerbank, 1866)	Yes	ZMBN 98079	HG423757	HG423787	HG423817	HG423728	North Sea, Norway, Hordaland	
<i>Tentorium papillatum</i> (Kirkpatrick, 1908)		SMF 10571	HG423758	*HG423788	HG423818	HG423729	Weddell Sea, Antarctic, Cape Norvegia	
<i>Tentorium papillatum</i> (Kirkpatrick, 1908)		ZMBN 98095	LN606519	LN606549	LN606579	—	Weddell Sea, Antarctic, South Shetland Islands	

Table 1 (continued)

Species	Type species	Museum voucher	28S: B10–C1	28S: D1–D19	28S: D20–E19	COI	Locality	Type locality
<i>Tentorium papillatum</i> (Kirkpatrick, 1908)		ZMBN 98096	HG423759	HG423789	HG423819	HG423730	Weddell Sea, Antarctic, South Shetland Islands	
<i>Tentorium semisuberites</i> (Schmidt, 1870)	Yes	ZMBN 98054	HG423760	HG423790	HG423820	HG423731	Arctic Ocean, Norway, Svalbard	
<i>Tentorium semisuberites</i> (Schmidt, 1870)	Yes	ZMBN 98099	HG423761	HG423791	HG423821	HG423732	North Sea, Norway, Hordaland	
<i>Tentorium</i> cf. <i>semisuberites</i> (Schmidt, 1870)		SMF 10575	HG423762	HG423792	HG423822	–	Weddell Sea, Antarctic, East for Antarctic Peninsula	
<i>Weberella bursa</i> (Müller, 1806)	Yes	ZMBN 98051	HG423763	HG423793	HG423823	HG423733	Arctic Ocean, Norway, Svalbard	
<i>Weberella bursa</i> (Müller, 1806)	Yes	ZMBN 98072	LN606520	LN606550	LN606580	LN606490	Norwegian Sea, Norway, Troms	
Outgroups								
Family Suberitidae								
<i>Suberites ficus</i> (Johnston, 1842)		BELUM: MC4322	HQ379247	HQ379322	HQ379389	HQ379429	Atlantic Ocean, UK, Scotland	
Family Tethyidae								
<i>Tethya citrina</i> Sarà & Melone, 1965		BELUM: MC5113	HQ379237	HQ379312	HQ379378	HQ379427	N/A	

polymastiid genera. Each genus was represented at least by the type species except for *Radiella* Schmidt, 1870, the type species of which, *Radiella sol* Schmidt, 1870, was unavailable and had an ambiguous status (see “Discussion”). Sequences from 19 species and nine OTUs were novel. Data on two species and one OTU were taken from GenBank and sequences from three species were both obtained by us and taken from GenBank. Two species were chosen as outgroups, the suberitid *Suberites ficus* (Johnston, 1842) and the tethyid *Tethya citrina* Sarà & Melone, 1965. Data on both species were taken from GenBank. This selection was based on the substantial morphological affinities between Suberitidae, Tethyidae Gray, 1848, and Polymastiidae and on the former affiliation of these three families with the order Hadromerida.

DNA extraction, amplification, and sequencing

DNA extractions were made with Qiagen DNeasy Blood and Tissue Minikits or DNeasy Plant Minikits following the manufacturer’s protocols (the latter was found to yield DNA of higher quantity and purity).

CO1 barcoding regions were amplified with the Ex Taq polymerase (TaKaRa) and the amplification program from Vargas et al. (2012). For most species and OTUs, we used the primers dgLCO1490/dgHCO2198 (Meyer 2003), which are slight modifications of the universal primers LCO1490/HCO2198 (Folmer et al. 1994). For one species, *Polymastia corticata* Rildey & Dendy, 1886, CO1 sequences of satisfactory quality could be obtained only with the primers jgLCO1490/jgHCO2198 (Geller et al. 2013).

Amplification of the partial 28S ribosomal DNA (rDNA) was performed with three pairs of primers designed by Morrow et al. (2012): Por28S-15F/Por28S-878R for sequencing of the region coding RNA from helix B10 to helix C1, Por28S-830F/Por28S-1520R for the region coding from D1 to D19, and Por28S-1490F/Por28S-2170R for the region coding from D20 to E18–E19. For most species, the amplification of this DNA piece succeeded with the Ex Taq polymerase and a “touchdown” program reported by Morrow et al. (2012), which was optimized by the doubling of the sequence extension time. In the cases of *Polymastia thielei* Koltun, 1964 and *Polymastia uberrima* (Schmidt, 1870), the amplification succeeded only with the LA Taq (TaKaRa) and the following protocol: 94 °C for 3 min (94 °C for 30 s, 50 °C for 45 s, 72 °C for 1 min) × 35 cycles, 72 °C for 7 min. Quality and quantity of the PCR products were estimated by agarose gel electrophoresis. The PCR products were purified with exonuclease 1 and shrimp alkaline phosphatase as described by Eilertsen and Malaquias (2013) and used for sequencing reactions with BigDye terminator 3.1 (Applied Biosystems, Waltham, MA, USA) following

the protocol of the producer. Subsequent sequence reads were performed with an automated ABI 3730XL DNA Analyser (Applied Biosystems) in the University of Bergen.

Contigs were assembled with the application SeqMan of DNASTAR Lasergene 8.0 and manually checked for sequencing errors. The consensus sequences of contigs were trimmed to remove primer residuals and checked by nucleotide BLAST search (Altschul et al. 1990) against GenBank sequences to verify their polymastiid origin. Where BLAST searches revealed epi- or endofaunal contaminations, the PCR products were cloned using StrataClone PCR cloning kit (Agilent Technologies, StrataGene Products Division, Santa Clara, CA, USA) according to the manufacturer's instructions, and 10–20 clones per product were sequenced by LGC Genomics GmbH (Berlin, Germany). If the direct sequence reads displayed double signals, extractions from the respective samples were repeated once again and the PCRs were repeated two or more times in order to exclude eventual cross-contamination and PCR errors. If these repetitive procedures confirmed the double signals in the reads, PCR products were cloned following the same protocol as used for the separation of the native and contaminating DNA fragments. The resulting clones were checked for errors, e.g., as those reported by Speksnijder et al. (2001) and Acinas et al. (2005), against the alignment of the approved direct sequences. Clones with unique nucleotides or gaps in the conservative sites were disregarded. Polymorphism in the remaining clones was regarded as natural. Strict consensus of the clones with the polymorphic sites encoded with IUPAC symbols were employed in the main phylogenetic analyses. Altogether we submitted 75 CO1 sequences and 236 sequences with the three regions of 28S rDNA including clone libraries (different versions from the same individuals) to GenBank (Table 1).

Alignments

All alignments were performed in SeaView 4.3.4 (Galtier et al. 1996; Gouy et al. 2010). CO1 sequences comprising exactly 658 bp each were aligned manually. The sequences of 28S rDNA varied in length. B10–C1 regions were 799–822 bp long, D1–D19 649–653 bp, and D20–E19 646–648 bp. Sequences of different regions were concatenated with 19 overlapping nucleotides, and their preliminary alignment was performed with the MUSCLE algorithm (Edgar 2004) implemented in SeaView. The initial alignment was further refined manually under consideration of the RNA secondary structure (Erpenbeck et al. 2007a, b, 2008). The GenBank sequence of *Polymastia pachymastia* de Laubenfels,

1932 (accession AY561924), being the longest (3550 bp) polymastiid sequence of 28S rDNA published so far, was used as a template for adapting the secondary structures reconstructed from other families to Polymastiidae. A 90 % consensus of all sequences employed in our study was then adjusted to this template. The resulting alignment was 2155 sites long. Because no secondary structure was proposed for the highly variable C-region (Erpenbeck et al. 2007a, b, 2008) comprising positions 406–813 in our alignment, we treated all sites flanked by the C1-helix strands as single stranded. Main length variation occurred in the C-region (from 366 to 396 bp) and in the terminal loop on helix D5 (alignment positions 940–948, variation from 4 to 8 bp). Search for unambiguously aligned sites was initially performed in GBlocks 0.91b (Castresana 2000) as implemented in SeaView. That excluded 51 sites. However, the resulting set was manually extended to exclude in total only 43 sites, because some obviously homologous sites were neglected by the algorithm. All alignments (CO1 matrix, 28S rDNA complete matrix, and 28S rDNA matrix reduced by 43 sites) were deposited at TreeBase and are available at <http://purl.org/phylo/treebase/phyloids/study/TB2:S18487>. Some descriptive statistics of the alignments was explored with MEGA7 (Nei and Kumar 2000; Kumar et al. 2016).

Selection of substitution models and phylogenetic analyses

For all computing procedures, identical sequences were collapsed into one sequence that is indicated by the sequence labels of the taxa represented in the trees (Figs. 1, 2, and 3). Five datasets, CO1 data alone (35 unique polymastiid sequences), two 28S rDNA matrices (49 sequences), the complete one and the matrix reduced by 43 ambiguously aligned sites, and two corresponding concatenated matrices (47 sequences), CO1 + complete 28S rDNA and CO1 + reduced 28S rDNA, were analyzed. Search for the best fitting substitution model for the CO1 dataset carried out with both hierarchical likelihood ratio tests and Akaike information criterion (AIC) in MrModeltest 2.0 (Nylander 2004) selected GTR+G+I. In all analyses, the CO1 data were split into two partitions, codon positions 1+2 and codon position 3. For RNA-specific models for the 28S rDNA datasets, we applied the model selection procedure implemented in PHASE 3.0 (Allen and Whelan 2014), a recent modification of PHASE 2.0 (Gowri-Shankar and Jow 2006), that included running the script “model_selection.pl” (Allen and Whelan 2014). As an input tree topology for this procedure, we used a ML tree calculated in PhyML (Guindon et al. 2010) under the best fitting standard model, GTR+G, determined with AIC in

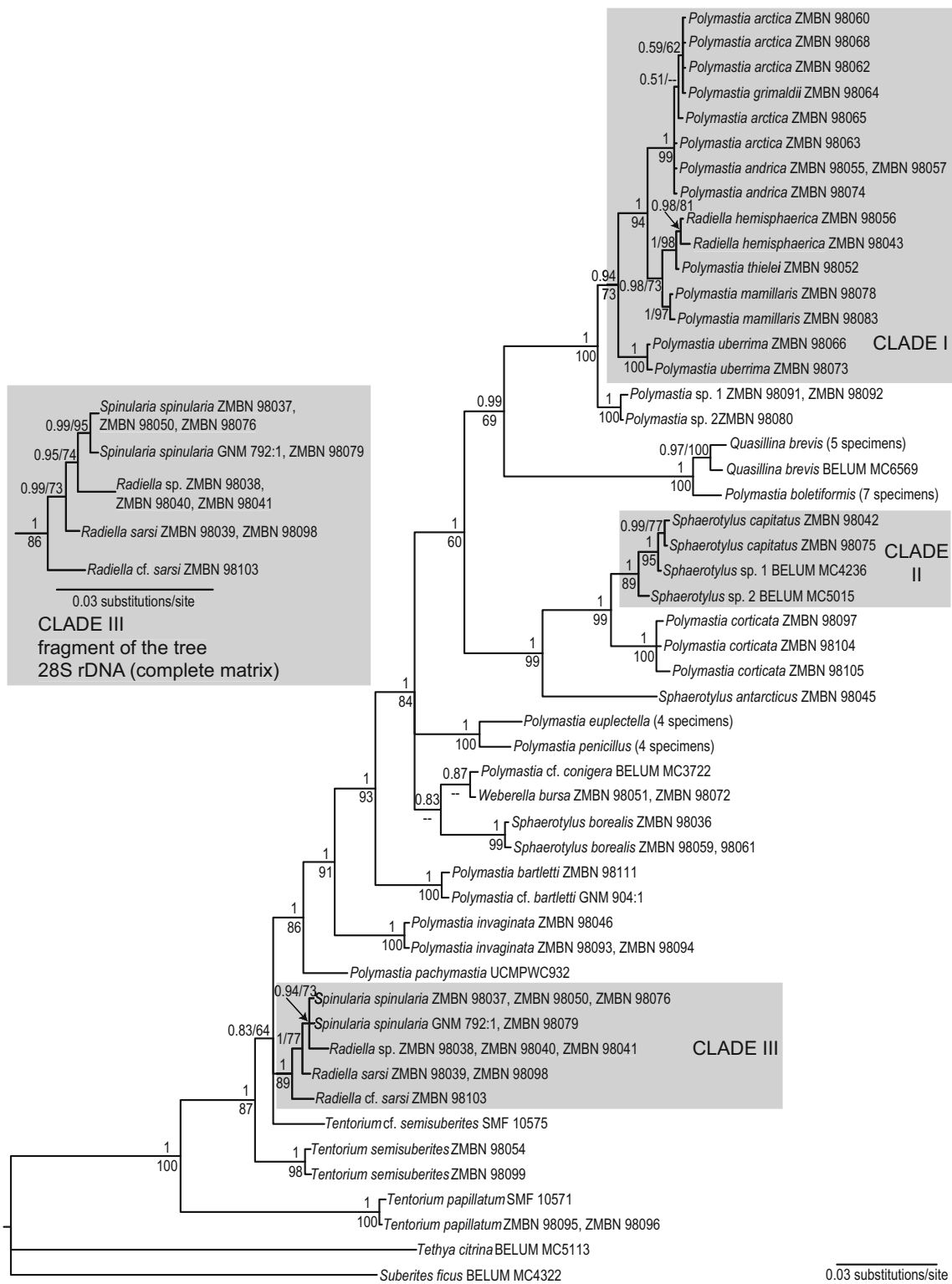
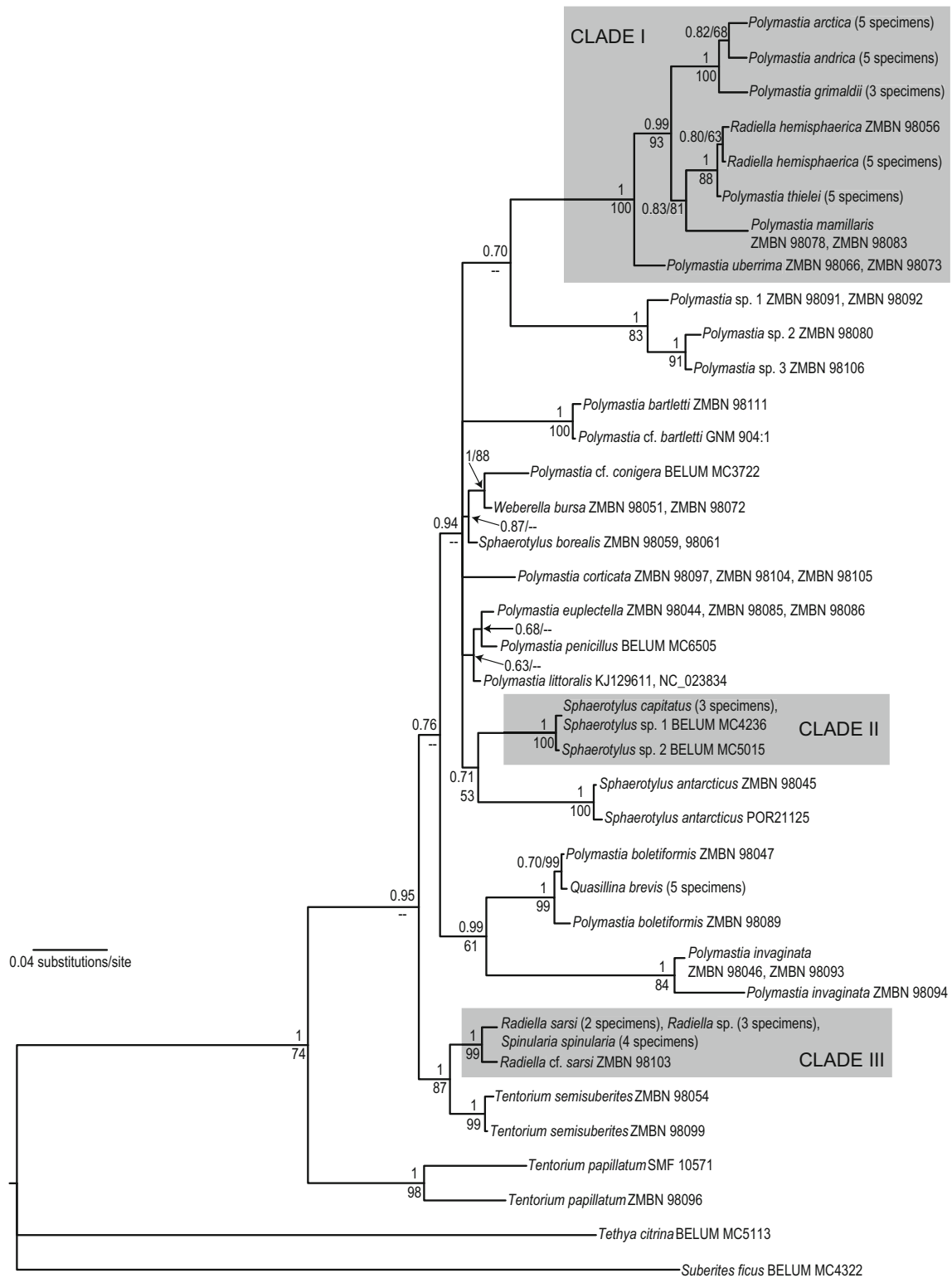


Fig. 1 Bayesian consensus tree reconstructed from the reduced 28S rDNA matrix. *Inset*: a fragment of the comparable Bayesian consensus tree reconstructed from the complete 28S rDNA matrix, displaying better resolution inside Clade III. Nodal supports: upper values—Bayesian posterior probabilities, lower values—ML bootstrap supports in percents. Expansion of the branch labels denoting multiple specimens: *Quasillina brevis* (five specimens)—ZMBN 98049, ZMBN 98067,

ZMBN 98082, ZMBN 98084, ZMBN 98090; *Polymastia boletiformis* (seven specimens)—BELUM: MC5014, GNM 901:1, ZMBN 98047, ZMBN 98048, ZMBN 98081, ZMBN 98088, ZMBN 98089; *Polymastia euplectella* (four specimens)—ZMBN 98044, ZMBN 98085, ZMBN 98086, ZMBN 98087; *Polymastia penicillus* (four specimens)—BELUM: MC5284, BELUM: MC6505, GNM 460:1, GNM 460:2



JModelTest 2.1.6 (Guindon and Gascuel 2003; Darriba et al. 2012). The mixed model RNA16C+G for helix positions and REV+G for loop positions was selected as the best fit. Analyses of the concatenated datasets CO1 + 28S rDNA were run under the mixed model

comprising the models selected for the single gene matrices. All datasets were analyzed in a Bayesian inference framework, with MrBayes 3.2 (Ronquist et al. 2011) for the CO1 matrix and with PHASE 3.0 (Allen and Whelan 2014) for the 28S rDNA matrices and the concatenated

◀ **Fig. 2** Bayesian consensus tree reconstructed from the CO1 matrix. Nodal supports: upper values—Bayesian posterior probabilities, lower values—ML bootstrap supports in percents. Expansion of the branch labels denoting multiple specimens: *Polymastia arctica* (five specimens)—ZMBN 98060, ZMBN 98062, ZMBN 98063, ZMBN 98065, ZMBN 98068; *Polymastia andrica* (five specimens)—ZMBN 98055, ZMBN 98057, ZMBN 98074, ZMBN 98102, ZMBN 98108; *Polymastia grimaldii* (three specimens)—ZMBN 98064, ZMBN 98110, ZMBN 98112; *Radiella hemisphaerica* (five specimens)—ZMBN 98043, ZMBN 98058, ZMBN 98069, ZMBN 98071, ZMBN 98077; *Polymastia thielei* (five specimens)—ZMBN 98052, ZMBN 98053, ZMBN 98070, ZMBN 98107, ZMBN 98109; *Sphaerotylus capitatus* (three specimens)—GNM 902, ZMBN 98042, ZMBN 98075; *Quasillina brevis* (five specimens)—ZMBN 98049, ZMBN 98067, ZMBN 98082, ZMBN 98084, ZMBN 98090; *Radiella sarsi* (two specimens)—ZMBN 98039, ZMBN 98098; *Radiella* sp. (three specimens)—ZMBN 98038, ZMBN 98040, ZMBN 98041; *Spinularia spinularia* (four specimens)—ZMBN 98037, ZMBN 98050, ZMBN 98076, ZMBN 98079

matrices CO1 + 28S rDNA, and in a maximum likelihood framework (ML) with RAxML 8.1.24 (Stamatakis 2014).

MrBayes 3.2 was run on the CIPRES (Cyberinfrastructure for Phylogenetic Research) Science Gateway V. 3.3 (<https://www.phylo.org/>) and on the Lifeportal at the University of Oslo using the high-performance computing cluster Abel (<https://lifeportal.uio.no/>). In the MrBayes 3.2 session, the model parameters were optimized independently for each partition. Two runs with eight chains each were launched under the default chain “temperatures” and flat Dirichlet distributions for the model parameter priors. The chains were sampled each 100 generations. The initial 2.5 million of the samples were disregarded in the burn-in. The convergence of the runs was controlled with the average standard deviation of split frequencies in MrBayes 3.2, while the sufficiency of the number of generations was estimated with the effective sample size (ESS) for all parameters in Tracer 1.5 (Rambaut and Drummond 2009). The convergence was reached and the ESSs exceeded 200 after ten million generations had been run.

PHASE analyses were performed on a desktop computer. Ten million iterations with sampling period 200 iterations after a burn-in of one million iterations were initially run. Each analysis was repeated twice, specifying a different random seed. After the output files had been transformed with the Perl script phase2tracer.pl (Voigt et al. 2012, modified from the script of Matt Yoder (<https://github.com/mjy/phase-utils/blob/master/phase2tracer.pl>)), the stabilization of all parameters was monitored in Tracer 1.5. If stabilization had not been achieved, the computations were repeated under optimized settings and with extra 5–30 million iterations.

RAxML 8.1.24 was run on the CIPRES. Search for the best scoring ML-tree along with rapid bootstrapping (1000 replicates) was performed. Because the model RNA16C is not

implemented in RAxML 8.1.24, the more exhaustive model RNA16A was invoked for helix positions of the 28S rDNA data.

Bayesian analyses of the single-gene datasets revealed some incongruence between the CO1 and 28S rDNA phylogenies. To illustrate the conflicts, we repeated the analyses on the matrices with the identical set of 47 taxa for both CO1 and 28S rDNA (reduced matrix) and, based on the resulting consensus trees, computed a rooted galled network (Huson et al. 2009) with Dendroscope 3 (Huson and Scornavacca 2012) (Fig. 4). To explore these conflicts, we performed an incongruence length difference test (ILD, Farris et al. 1994) on the concatenated dataset CO1 + reduced 28S rDNA with all parsimony uninformative sites excluded running 500 replicates in PAUP* 4.0b10 (Swofford 2002). Furthermore, we used Bayes factor comparisons of the model likelihoods to test the conflicting topological hypotheses on the single-gene datasets following Kass and Raftery (1995). To obtain more accurate model likelihoods, stepping-stone samplings were performed in MrBayes 3.2. The monophyly of the congruent clades was constrained as recommended by Bergsten et al. (2013). Two runs with two chains each were launched. Four million generations on the CO1 data and ten million generations on the 28S rDNA were run to reach the convergence of the runs.

Additionally, in order to examine the intragenomic polymorphism of the D1–D19 region of 28S rDNA, a dataset comprising all versions of this region in three species of *Polymastia*, *Polymastia andrica* de Laubenfels, 1949, *Polymastia arctica* (Merejkowsky, 1878), and *Polymastia grimaldii* (Topsent, 1913), with *Polymastia mamillaris* (Müller, 1806) as the outgroup taxon, was analyzed with Minimum-spanning network algorithm (Bandelt et al. 1999) implemented in PopArt 1.7 (<http://popart.otago.ac.nz>) and in a ML framework with PhyML (Guindon et al. 2010). Consensus trees resulting from the Bayesian analyses along with the ML-tree illustrating the intragenomic polymorphism were deposited at TreeBase and are available at <http://purl.org/phylo/treebase/phyloids/study/TB2:S18487>.

Tracing of the evolution of morphological characters

The consensus tree resulting from the Bayesian analysis of the concatenated dataset CO1 + reduced 28S rDNA was chosen for tracing of the morphological evolution. Branches corresponding to different individuals of the same species or OTU were collapsed. A matrix with 21 morphological characters of the respective 30 polymastiid taxa and two outgroup taxa was built based on the dataset for phylogenetic scenario 3 from Plotkin et al. (2012), but with two emendations: the multistate character “Growth pattern” was assigned by the amalgamation of four binary characters and the character “Longitudinal tracts of principal monactines in the

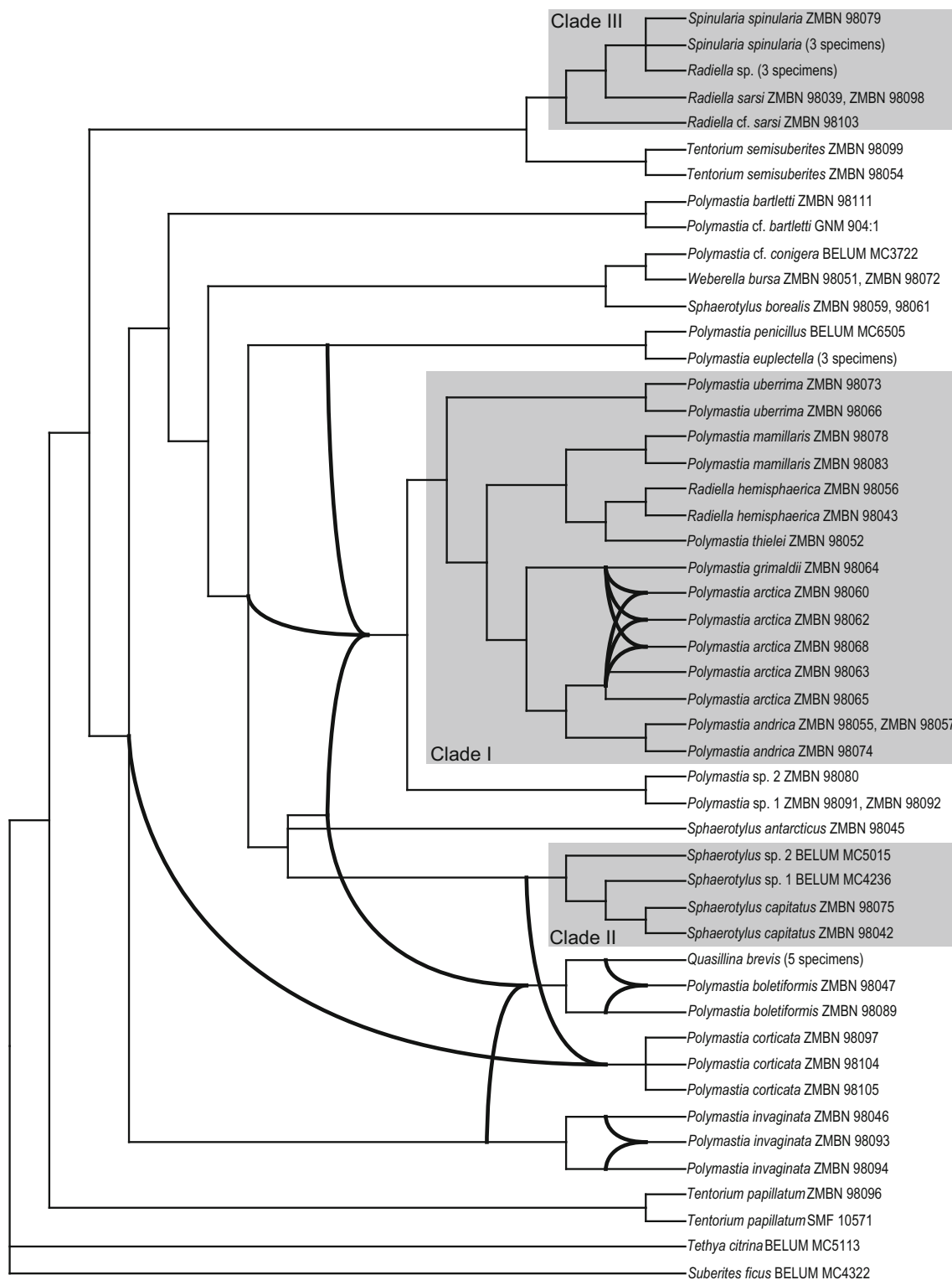


Fig. 4 Rooted galled network compiled from the Bayesian consensus trees reconstructed from CO1 alone and 28S rDNA alone with identical sets of taxa. **Bold curves** indicate discrepancies in the topology. Expansion of the branch labels denoting multiple specimens: *Spinularia spinularia* (three specimens)—ZMBN 98037, ZMBN 98050, ZMBN

98076; *Radiella* sp. (three specimens)—ZMBN 98038, ZMBN 98040, ZMBN 98041; *Polymastia euplectella* (three specimens)—ZMBN 98044, ZMBN 98085, ZMBN 98086; *Quasillina brevis* (five specimens)—ZMBN 98049, ZMBN 98067, ZMBN 98082, ZMBN 98084, ZMBN 98090

reconstruction with parsimony criterion for each character was performed in Mesquite 3.04 (Maddison and

Maddison 2015), while the consistency indices were computed in PAUP* 4.0b10 (Swofford 2002).

Results

Statistics of alignments

Parsimony-informative sites comprised almost 25 % of all sites in the CO1 matrix and about 15 % of all sites in the 28S rDNA matrices. Sites with intragenomic polymorphisms comprised ca. 1 and 0.7 % of all sites in the 28S rDNA complete matrix and reduced matrix, respectively (Table 2).

Congruent aspects of the CO1 and 28S rDNA phylogenies

Separate analyses of CO1 and 28S rDNA resulted in similar overall phylogenies (Figs. 1 and 2) except for few conflicts (see the section “[Incongruence between the CO1 and 28S rDNA phylogenies](#)” below). The phylogenies were not affected by intraspecific or intragenomic polymorphisms except for the relationships within three small terminal subclades (see the section “[Intraspecific and intragenomic polymorphism](#)” below). All analyses supported the monophyly of the polymastiids against the two outgroups (Figs. 1, 2, and 3). At the same time, the polymastiid genera *Polymastia*, *Radiella*, *Sphaerotylus* Topsent, 1898, and *Tentorium* Vosmaer, 1887 were nonmonophyletic. *Polymastia* spp. were scattered over different clades, *Radiella hemisphaerica* (Sars, 1872) fell distantly from other *Radiella* spp., *Sphaerotylus borealis* (Swarzewsky, 1906) lay remotely from its congeners, and *Tentorium papillatum* (Kirkpatrick, 1908) fell on a long branch as the sister group to a clade of the remaining polymastiids. Moreover, in the 28S rDNA tree, the type species of *Tentorium*, *Ten. semisuberites*, and *Ten. cf. semisuberites* did not group together, although the support for their nonmonophyly was very weak (Fig. 1). Unfortunately, no CO1 data from *Ten. cf. semisuberites* were obtained. Three clades of species (highlighted in Figs. 1, 2, 3, and 4) were recovered by all analyses.

Clade I comprised *Polymastia andrica*, *P. arctica*, *P. grimaldii*, *P. mamillaris* (type species of *Polymastia*), *P. thielei*, *P. uberrima*, and *Radiella hemisphaerica*. The support for this clade in the 28S rDNA tree was slightly weaker (Fig. 1) than in the CO1 tree (Fig. 2). Analyses of the 28S rDNA alone and the concatenated datasets CO1 + 28S rDNA supported the sister relationships between the pair *Polymastia* sp. 1 + *Polymastia* sp. 2 and Clade I (Figs. 1 and

3). In the CO1 tree, *Polymastia* sp. 1 and *Polymastia* sp. 2 grouped with *Polymastia* sp. 3, and the position of this trio as the sister to Clade I had very weak support (Fig. 2). Unfortunately, no 28S rDNA was obtained from *Polymastia* sp. 3. Inside Clade I, *P. uberrima* was the sister to the subclade of the remaining six species strongly supported by all analyses (Figs. 1, 2, and 3). In its turn, this subclade split up into two groupings, *P. mamillaris* + *P. thielei* + *R. hemisphaerica* and *P. andrica* + *P. arctica* + *P. grimaldii*. The pair *P. thielei* + *R. hemisphaerica* was supported by all analyses (Figs. 1, 2, and 3). But the grouping of *P. mamillaris* as the sister to this pair was strongly supported only by the analyses of 28S rDNA alone (Fig. 1) and the concatenated datasets (Fig. 3), while the analysis of CO1 alone demonstrated just a very weak support for this relationship (Fig. 2). The grouping *P. andrica* + *P. arctica* + *P. grimaldii* was supported in all analyses (Figs. 1, 2, and 3), but the relationships between these three species were unresolved in the 28S rDNA tree (Fig. 1) because of the intraspecific and intragenomic polymorphism (see the respective section below) and resolved with just a low support for *P. andrica* + *P. arctica* in the CO1 tree (Fig. 2).

Clade II comprised *Sphaerotylus capitatus* (Vosmaer, 1885) (type species of *Sphaerotylus*), *Sphaerotylus* sp. 1 and *Sphaerotylus* sp. 2. This clade was strongly supported in all trees (Figs. 1, 2, and 3). *Sph. capitatus* and *Sphaerotylus* sp. 1 had identical CO1 and were sisters in the 28S rDNA tree with strong support (Fig. 1).

Clade III comprised *Radiella sarsi*, *Radiella cf. sarsi*, *Radiella* sp., and *Spinularia spinularia* (Bowerbank, 1866) (type species of *Spinularia* Gray, 1867). This clade was strongly supported in all trees (Figs. 1, 2, and 3). In the CO1 tree, Clade III and *Tentorium semisuberites* were sisters with strong support (Fig. 2), but this was not confirmed by the analyses of 28S rDNA alone (Fig. 1). *R. sarsi*, *Radiella* sp., and *Spi. spinularia* had identical CO1 and formed a strongly supported subclade in the 28S rDNA tree (Fig. 1). In the same tree, there was also some support for the sister relationships between *Radiella* sp. and *Spi. spinularia*. *Spi. spinularia* was represented by two groups of individuals that differed from each other by two nucleotides in 28S rDNA and each of them differed from *Radiella* sp. (identical sequences from three individuals) by 11 nucleotides. Three of these 11 nucleotides were located in the 43 sites excluded as ambiguously aligned in the matrix as a whole. The exclusion of these sites from the

Table 2 Basic statistics of the analyzed alignments

Alignment	Length	Variable sites	Parsimony-informative sites	Sites with intragenomic polymorphism	Empirical nucleotide frequencies, % for T, C, A, G	Average p-distance (SE)
CO1	658	201	163	0	36.3, 15.8, 25.7, 22.2	0.080 (0.006)
LSU, complete matrix	2155	363	326	21	20.7, 23.7, 23.4, 32.2	0.047 (0.003)
LSU, reduced matrix	2112	347	310	14	20.7, 23.6, 23.6, 32.2	0.045 (0.003)

analyses led to a polytomy formed by *Radiella* sp. and the two groups of individuals of *Spi. spinularia* (main trees in Figs. 1 and 3). However, within Clade III, these excluded sites could be aligned unambiguously and provided a sufficient phylogenetic signal to resolve the polytomy (insets in Figs. 1 and 3).

Furthermore, all analyses strongly supported the pair *Polymastia boletiformis* (Lamarck, 1815) + *Q. brevis* (type species of *Quasillina*) (Figs. 1, 2, and 3) and also revealed the grouping *Sphaerotylus borealis* + *Polymastia* cf. *conigera* Bowerbank, 1874 + *Weberella bursa* (Müller, 1806), although the latter was strongly supported only by the analyses of the concatenated data CO1 + 28S rDNA (Fig. 3), while its support in the single-gene trees was very weak (Figs. 1 and 2). Within this grouping, *P. cf. conigera* and *W. bursa* were sisters with a strong support in the CO1 tree (Fig. 2), but a much weaker support in the 28S rDNA tree (Fig. 1). At the same time, the analyses of the 28S rDNA alone strongly supported the pair *Polymastia euplectella* Rezvoj, 1927 + *Polymastia penicillus* (Montagu, 1818) (Fig. 1), while the support for this pair in the CO1 tree was negligible (Fig. 2).

Incongruence between the CO1 and 28S rDNA phylogenies

ILD test of the concatenated dataset CO1 + 28S rDNA rejected the hypothesis of congruent data with a *p* value of 0.002. The conflicts between the single gene phylogenies are visualized as reticulations in the galled network (Fig. 4). One conflict concerned dissimilarity in the relationships between *Polymastia boletiformis* + *Quasillina brevis* and other taxa. In the CO1 tree, this pair was the sister to *Polymastia invaginata* Kirkpatrick, 1907 (Fig. 2), whereas in the 28S rDNA tree, it was the sister to the grouping Clade I + *Polymastia* sp. 1 + *Polymastia* sp. 2 (Fig. 1). Bayesian support for the indicated relationships was strong in each gene tree, and they were not affected by polymorphism in any of the species. Bayes factor tests revealed no support for the alternative hypothesis in either of the two topologies (Table 3). Two conflicts were due to the low resolution of the CO1 tree. In this tree, five clades formed an unresolved polytomy with *Polymastia corticata*, while *Sphaerotylus antarcticus* Kirkpatrick, 1907 was the sister to Clade II, although with very weak Bayesian support (Fig. 2). Conversely, in the 28S rDNA tree, *P. corticata* was the sister to Clade II, and *Sph. antarcticus* in its turn was the sister to *P. corticata* + Clade II with strong support (Fig. 1). The conflicts between the CO1 and 28S rDNA phylogenies caused by the gene polymorphism were revealed within three small terminal subclades, the trio *Polymastia andrica* + *P. arctica* + *P. grimaldii* in Clade I, the pair *P. boletiformis* + *Q. brevis*, and the group of three individuals of *P. invaginata* (Fig. 4). These conflicts are considered below.

Intraspecific and intragenomic polymorphism

The most conspicuous intraspecific polymorphism was revealed in four sites of the B10–C1 region (positions 578–580 and 583 in the complete matrix) and in seven sites of the D1–D19 region (positions 941–943, 947–948, and 1294–1295) of 28S rDNA in *Polymastia andrica*, *P. arctica*, and *P. grimaldii*. The variation within B10–C1 was estimated on the direct sequences. The sequences of this region from three *P. andrica* were identical, while *P. arctica* displayed a polymorphism—individual ZMBN 98063 differed from *P. andrica* just by one ambiguity, individual ZMBN 98068 by three nucleotides, and two individuals, ZMBN 98060 and ZMBN 98062, by four nucleotides. *P. grimaldii* ZMBN 98064 differed from the latter two individuals of *P. arctica* just by one ambiguity. The variation within D1–D19 was estimated on four direct sequences (two from *P. andrica* and two from *P. arctica*) and 18 clones from five individuals (one of *P. andrica*, three of *P. arctica*, and one of *P. grimaldii*). Of the seven polymorphic sites, five were parsimony-informative when the sequences were aligned with the corresponding DNA region of *P. mamillaris* as the outgroup (see the respective alignment at <http://purl.org/phylo/treebase/phyloWS/study/TB2:S18487>). Here an intragenomic polymorphism was discovered. Seven versions of D1–D19 were spread among the individuals of different species (Fig. 5):

- Version 1 (in three *P. andrica*, three *P. arctica*, and the only *P. grimaldii* employed in this analysis)
- Version 2 (in one *P. andrica* and one *P. arctica*)
- Version 3 (in one *P. arctica* and *P. grimaldii*)
- Version 4 (only in *P. grimaldii*)
- Version 5 (in one *P. andrica*, two *P. arctica*, and *P. grimaldii*)
- Version 6 (in one *P. andrica*, two *P. arctica*, and *P. grimaldii*)
- Version 7 (in one *P. andrica* and one *P. arctica*)

While the 28S rDNA sequences of *P. andrica*, *P. arctica*, and *P. grimaldii* displayed intraspecific and intragenomic polymorphism, the CO1 data from these species were consistent, i.e., the sequences from the individuals of the same species were identical. Based on these data, *P. andrica* and *P. arctica* were sisters, although the support for this relationship was weak (Fig. 2).

Among other species, intraspecific polymorphism was revealed in *Polymastia boletiformis* and *P. invaginata*. All seven *P. boletiformis* studied had identical 28S rDNA and grouped with *Quasillina brevis* (Fig. 1). CO1 was obtained only from two individuals of *P. boletiformis*, of which one, ZMBN 98047, differed from *Q. brevis* just by one nucleotide in this gene, while the other, ZMBN 98089, differed from *Q. brevis* by six nucleotides, that resulted in a grouping of ZMBN 98047 with *Q. brevis* instead of the conspecific ZMBN 98089 (Fig. 2).

Table 3 Results of the testing of two conflicting topological hypotheses about the relationships of *Polymastia boletiformis* + *Quasillina brevis* with other taxa

Phylogenetic marker	Topological hypothesis	Mean marginal likelihood, natural log units	Compared hypotheses	$2 \times \log$ Bayes factor
CO1	H0: <i>P. boletiformis</i> + <i>Q. brevis</i> is sister to Clade I + <i>Polymastia</i> sp. 1 + <i>Polymastia</i> sp. 2	-4082.3	H1 vs. H0	-32.66
	H1: <i>P. boletiformis</i> + <i>Q. brevis</i> is sister to <i>P. invaginata</i>	-4065.97		
28S rDNA	H0: <i>P. boletiformis</i> + <i>Q. brevis</i> is sister to Clade I + <i>Polymastia</i> sp. 1 + <i>Polymastia</i> sp. 2	-7609.05	H1 vs. H0	34.96
	H1: <i>P. boletiformis</i> + <i>Q. brevis</i> is sister to <i>P. invaginata</i>	-7626.53		

Cloning of the PCR products confirmed these relationships. Two individuals of *P. invaginata*, ZMBN 98093 and ZMBN 98094, had identical 28S rDNA, whereas individual ZMBN 98046 differed from them by two nucleotides (Fig. 1). Conversely, CO1 of ZMBN 98046 and ZMBN 98093 were identical, while ZMBN 98094 differed from them by 19 nucleotides (Fig. 2).

Homoplasy of the morphological characters

Only two characters were fully consistent—character 5, “Exhalant papillae”, and character 9, “Oscula on the body surface” (Online Resources 1–3). All polymastiid taxa studied possess exhalant papillae and lack oscula on the surface, i.e.,

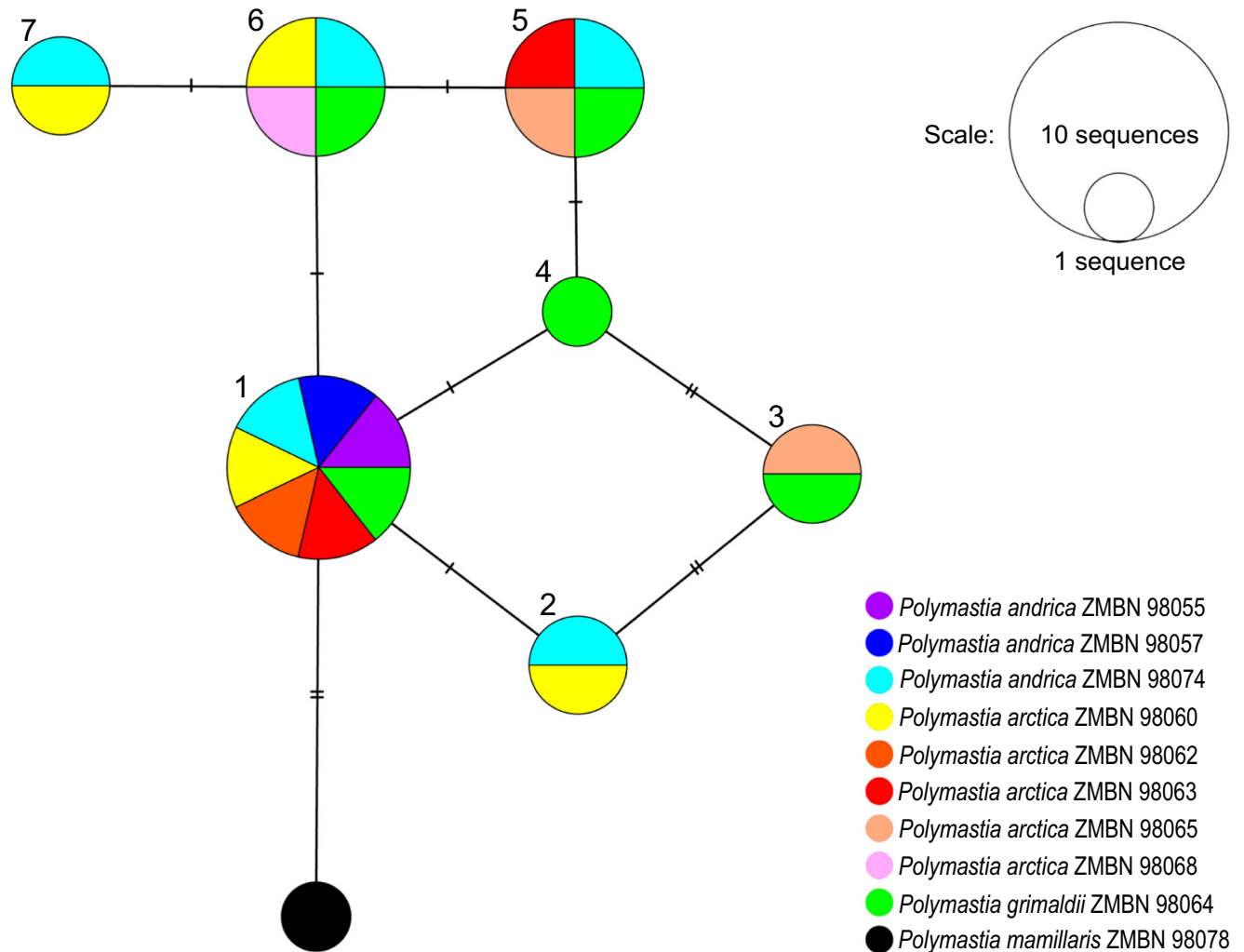


Fig. 5 Minimum-spanning network reconstructed from the dataset of the clones and direct sequences of the D1–D19 region, 28S rDNA from *Polymastia andrica*, *P. arctica*, and *P. grimaldii* with *P. mamillaris* as

outgroup. Numbers denote gene versions described in the text. Hatch marks on the branches indicate mutations

these features are the synapomorphies of the polymastiid clade compared with the given outgroups. All other characters were more or less homoplastic (Online Resources 1–3).

Most polymastiids possess one or several well-developed exhalant papillae (characters 6 and 7 in Online Resources 1–3, Fig. 6a–c). Weakly developed papillae (Fig. 6d–e) are present in the taxa from three remote groupings (Fig. 6b)—in all species from Clade III + *Tentorium semisuberites*, in *Sphaerotylus capitatus* from Clade II and in *Quasillina brevis* from the pair *Polymastia boletiformis* + *Q. brevis*. Among these taxa, only *Q. brevis* and three species of *Radiella*, *R. sarsi*, *R. cf. sarsi*, and *Radiella* sp., always possess just single papillae (Fig. 6d), whereas *Spinularia spinularia*, *Ten. semisuberites*, and *Sph. capitatus* may have several exhalant papillae as all other polymastiids studied (Fig. 6e).

Exotyles, i.e., extraordinary spicules protruding above the sponge surface (character 14 in Online Resources 1–3), are present in three remote groupings (Fig. 6f). Within the *Sphaerotylus borealis* + *Polymastia* cf. *conigera* + *Weberella bursa* group, the first two species possess exotyles, whereas *W. bursa* does not. All three species of Clade II have exotyles (Fig. 6h–i). Of the two species grouping with Clade II in the 28S rDNA tree, one species, *Sphaerotylus antarcticus*, possesses exotyles, while the other, *Polymastia corticata*, has no exotyles. Among the species of Clade I, *P. andrica* possesses exotyles.

The presence of a marginal fringe of extra-long spicules (character 16 in Online Resources 1–3, Fig. 6g) is shared by all species of Clade III, but also recorded in two species falling into two different subclades of Clade I, *Radiella hemisphaerica* (Fig. 6j) and *Polymastia grimaldii*.

Most polymastiids share the presence of a well-developed and regular choanosomal skeleton with one of the outgroup species, *Tethya citrina* (character 17 in Online Resources 1–3, Fig. 6k). The only exception is *Quasillina brevis* (Fig. 6m), which shares the presence of an irregular and reduced choanosomal skeleton with the other outgroup taxon, *Suberites ficus*.

The main choanosomal skeleton in most polymastiids as well as in *Tet. citrina* is radial (character 18 in Online Resources 1–3, Fig. 6l, n). Reticulate choanosomal skeleton (Fig. 6o) is recorded in taxa from four remote groupings—in *Weberella bursa* (*Sphaerotylus borealis* + *Polymastia* cf. *conigera* + *W. bursa*), in *Polymastia corticata* (*P. corticata* + Clade II), in *Polymastia boletiformis* (*P. boletiformis* + *Quasillina brevis*), and in *Polymastia thielei* (Clade I).

Discussion

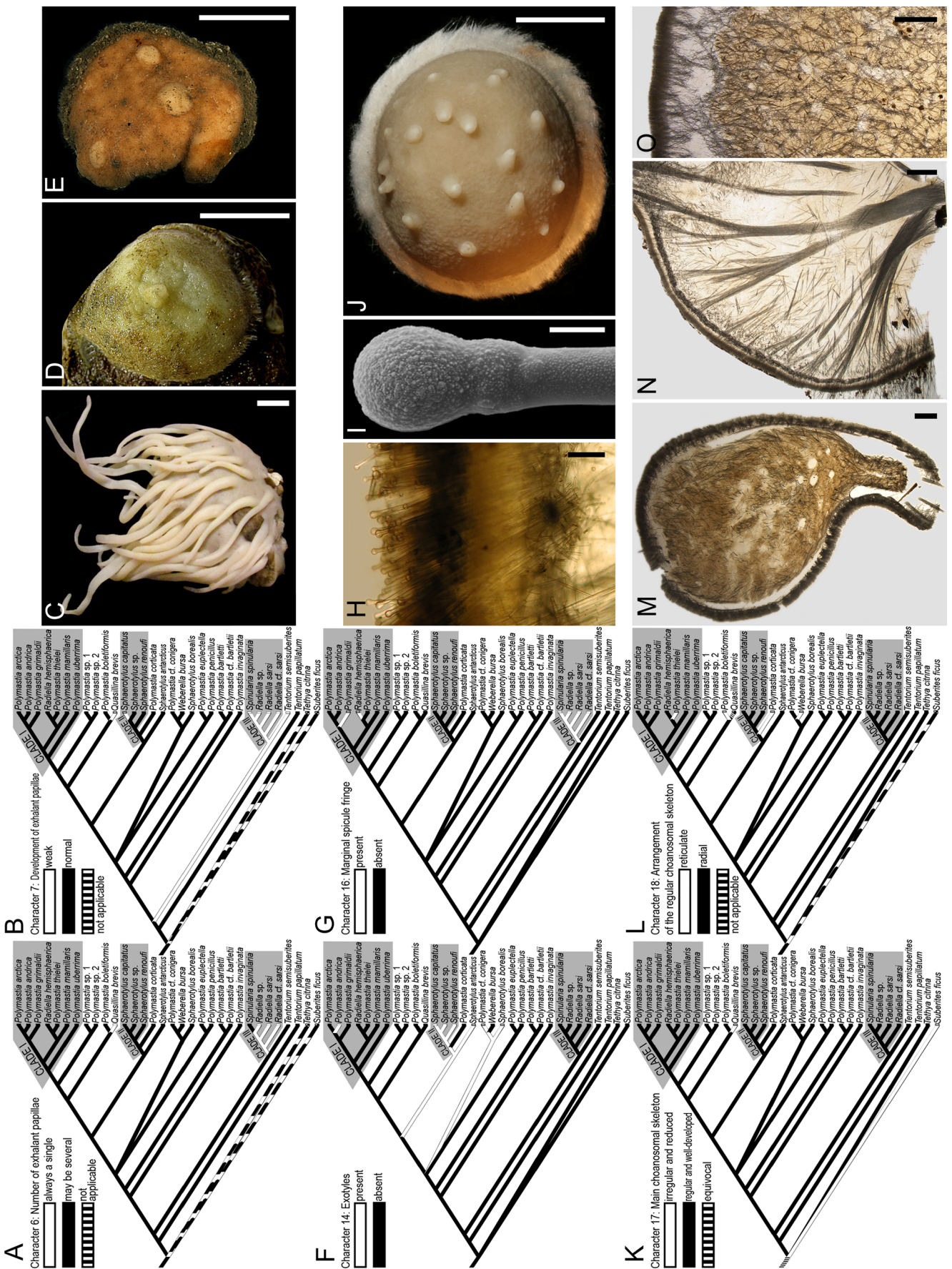
Monophyly of Polymastiidae and taxa of uncertain family affiliation

Our analyses demonstrated the monophyly of the clade formed by all polymastiid taxa studied when using a suberitid

and a tethyid as outgroups, and these results are congruent with most previous studies (Morrow et al. 2012, 2013; Redmond et al. 2013; Vargas et al. 2015). However, it should be noted that due to the lack of reliable molecular data, our analyses did not consider the two taxa with uncertain family affiliation, the genus *Pseudotrachya* Hallmann, 1914 and the species *Aaptos papillata*, which seem to be important for understanding of the polymastiid early evolution. *A. papillata* is usually regarded as a suberitid (Van Soest et al. 2015), despite that it displays strong morphological similarities with Polymastiidae, and in the trees reconstructed from a morphological dataset by Plotkin et al. (2012), it formed a polytomy with *Tentorium semisuberites* and the clade of other polymastiids. *Pseudotrachya* was commonly regarded as a polymastiid genus (Boury-Esnault 2002). But the type species of this genus, *P. hystrix*, did not group with other polymastiids in the phylogenies based on morphological data (Plotkin et al. 2012), and similarly *Pseudotrachya* sp. fell outside the main polymastiid clade in the tree reconstructed from a 28S rDNA dataset (Nichols 2005). Meanwhile, the taxonomic identification in the latter study raised some doubts because in the same 28S rDNA tree, another polymastiid, *Polymastia* sp. 1, also did not group with the main polymastiid clade, while in the CO1 trees, *Pseudotrachya* sp., *Polymastia* sp. 1, and *Polymastia* sp. 2 appeared in different clades. Additionally, the 28S rDNA sequences of these three species recovered by Nichols (2005) were much shorter than those used in our study, and therefore, we did not include Nichols' sequences in the analyses.

Molecular phylogenies contradict the morphology-based classification of polymastiids

Our most important outcome is the inapplicability of morphological characters, the majority of which has appeared to be highly homoplastic, for the phylogenetic reconstruction and hence for the natural classification of Polymastiidae. Homoplasy is a general problem in morphological taxonomy of the demosponges (e.g., Cárdenas et al. 2011; Morrow et al. 2013). Meanwhile, inside Polymastiidae, our study has recovered three clades strongly supported by the data from both genes studied. Each clade includes the type species of the certain genus, Clade I—the type species of *Polymastia*, *P. mamillaris*; Clade II—the type species of *Sphaerotylus*, *Sph. capitatus*; and Clade III—the type species of *Spinularia*, *Spi. spinularia*, and hence, these clades may be used as reference points in future classification of the family. However, no morphological synapomorphies can at present be defined for the clades revealed. Moreover, about 58 % of the species studied do not fall into any of these clades. Thus, for the time being, we cannot propose a satisfactory classification of Polymastiidae.



◀ **Fig. 6** Key morphological characters of Polymastiidae: depiction of states and tracing of evolution along the Bayesian consensus tree reconstructed from the concatenated dataset CO1 + reduced 28S rDNA (the same as in Fig. 3, but with the branches corresponding to different individuals of the same species collapsed). **a** Evolution of the character “Number of exhalant papillae” (N 6 in Online Resources 1–3); **b** evolution of the character “Development of exhalant papillae” (N 7 in Online Resources 1–3); **c** numerous normally developed papillae in *Polymastia bartletti* ZMBN 98111 (University Museum of Bergen); **d** single weakly developed papilla of *Radiella* sp. ZMBN 98040 (sampled from the Norwegian Sea); **e** three weakly developed papillae in *Spinularia spinularia* (not deposited); **f** evolution of the character “Exotyles” (N 14 in Online Resources 1–3); **g** evolution of the character “Marginal spicule fringe” (N 16 in Online Resources 1–3); **h** exotyles projecting above the cortex, histological section through the body of *Sphaerotylus capitatus* BMNH 10.1.1.1199 (paralectotype, Natural History Museum, London); **i** distal ornamentation of an exotyle, SEM image, preparation from *Sphaerotylus capitatus* ZMB 10855 (Museum für Naturkunde, Berlin); **j** prominent marginal spicule fringe bordering the body of *Radiella hemisphaerica* NHMO-B862 (holotype, Natural History Museum, University of Oslo); **k** evolution of the character “Main choanosomal skeleton” (N 17 in Online Resources 1–3); **l** evolution of the character “Arrangement of the regular choanosomal skeleton” (N 18 in Online Resources 1–3); **m** irregular and reduced choanosomal skeleton, histological section through the body of *Quasillina brevis* (not deposited); **n** regular radial choanosomal skeleton, histological section through the body of *Polymastia arctica* ZMBN 98068 (University Museum of Bergen); **o** regular reticulate choanosomal skeleton, histological section through the body of *Weberella bursa* (not deposited). Scale bars **c–e**: 1 cm, **h**: 0.1 mm; **i**: 0.01 mm; **j**: 1 cm; **m–o**: 1 mm

Abandonment of *Radiella*

We can, however, propose to transfer *Radiella hemisphaerica* to *Polymastia* since this species groups with the type species of *Polymastia* and five other *Polymastia* spp. forming Clade I. Likewise, *Radiella sarsi*, *Radiella* cf. *sarsi*, and *Radiella* sp. can be allocated to *Spinularia* since these three species and the type species of *Spinularia* form a monophyletic group, Clade II. The status of the genus *Radiella* is controversial (Boury-Esnault 2002; Plotkin and Janussen 2008; Plotkin et al. 2012), and the affinities of its type species, *R. sol* Schmidt, 1870, still remain ambiguous. Type material is lost, fresh material is not available, and the age-old nontype specimen identified as *R. sol* by Schmidt (1880) and redescribed by Boury-Esnault (2002) displays similarity to *R. hemisphaerica*, that does not match the drawing in the original description (Schmidt, 1870), which rather shows similarity between *R. sol* and *R. sarsi* (Plotkin and Janussen 2008; Plotkin et al. 2012). Nevertheless, no matter whether *R. sol* is related to *R. hemisphaerica* or *R. sarsi*, we propose to abandon *Radiella* since *R. hemisphaerica* is placed in *Polymastia*, *R. sarsi* is placed in *Spinularia*, and both *Polymastia* Bowerbank, 1862 and *Spinularia* Gray, 1867 are older names than *Radiella* Schmidt, 1870.

Disjunct distributions of polymastiid species are questioned

Our study questions the bipolar distribution of two polymastiid species, *Tentorium semisuberites* and *Radiella sarsi*. In the 28S rDNA tree, morphologically very similar *Ten. semisuberites* from the North Atlantic and *Ten.* cf. *semisuberites* from the Antarctic (see Table 1 for details on geography) did not group together. The type locality of *Ten. semisuberites* is Greenland (Schmidt 1870), and hence, we assume that the Antarctic sponges may be a separate species. Likewise, the pair of morphologically similar *R. sarsi* from the Norwegian Sea and *Radiella* cf. *sarsi* from Mozambiquean Coast was nonmonophyletic in both CO1 and 28S rDNA trees that questioned the allocation of these two to the same species.

Another example calling for reflection on disjunct distributions of sponge species is the pair *Polymastia euplectella* and *Polymastia bartletti* de Laubenfels, 1942, which display strong morphological similarities. The former species was recorded from the Barents Sea and adjacent areas (Rezvoj 1927; Plotkin 2004), whereas the latter was, before our study, known only from the type locality in the Baffin Sea (de Laubenfels 1942). We got genetic data from four Norwegian individuals identified as *P. euplectella*, one Canadian individual identified as *P. bartletti*, and a juvenile sponge from Sweden considered as *P.* cf. *bartletti*. These sponges were very similar in morphology, but in all phylogenetic trees, the Canadian individual and the Swedish sponge fell quite distantly from the Norwegian *P. euplectella*. All *P. euplectella* had identical CO1 and 28S rDNA except for one individual, for which no CO1 was obtained. *P. bartletti* and *P.* cf. *bartletti* slightly differed in both genes, but still were sisters. These results argue for that *P. euplectella* and *P. bartletti* are valid species. However, without studying more material, we cannot judge whether the small genetic differences between the Canadian *P. bartletti* and the Swedish *P.* cf. *bartletti* is just an intraspecific polymorphism, or these two are different, recently diverged species. Consequently, we cannot conclude whether *P. bartletti* is geographically isolated from *P. euplectella* or not.

Possible reasons for inconsistency between the single-gene phylogenies

Our analyses and tests supported dissimilar relationships of *Polymastia boletiformis* + *Quasillina brevis* with other clades and taxa in the single-gene trees. In the CO1 tree, this pair was the sister to *Polymastia invaginata*, whereas in the 28S rDNA tree, it was the sister to Clade I + *Polymastia* sp. 1 + *Polymastia* sp. 2. This may be due to real differences in genealogical histories of the mitochondrial and nuclear genes. However, on the other hand, the position of *P. boletiformis* + *Q. brevis* in the CO1 tree may be affected by very low resolution leading to Clade I. Likewise, unresolved

relationships of *Polymastia corticata* along with weakly supported grouping of *Sphaerotylus antarcticus* with Clade II in the CO1 tree are obviously due to low resolution and hence to weak phylogenetic signal provided by our CO1 data.

Intraspecific polymorphism indicates incomplete lineage sorting?

Several inconsistencies between the CO1 and 28S rDNA phylogenies within three small terminate subclades were caused by intraspecific polymorphism. Our study could not resolve the relationships between three sibling species of *Polymastia*, *P. andrica*, *P. arctica*, and *P. grimaldii*. The CO1 data on these species were congruent with the morphological differences between them—individuals of the same species possessed identical CO1 and morphologically similar *P. andrica* and *P. arctica* grouped together against morphologically more distinct *P. grimaldii*, although the Bayesian and ML bootstrap support for this relationship was rather weak. At the same time, the analysis of 28S rDNA failed to resolve the relationships between these three species because of the intraspecific and intragenomic polymorphism. Intraspecific polymorphism was also revealed in *P. boletiformis* and *P. invaginata*. Two individuals of *P. boletiformis* possessed identical 28S rDNA, but fairly different CO1. In three individuals of *P. invaginata*, the identity by the mitochondrial gene mismatched that by the nuclear gene.

The ascertained cases of intraspecific polymorphism may indicate incomplete lineage sorting in the closely related polymastiid species and their populations. For example, each lineage may carry one unique version of CO1, but several versions of 28S rDNA, if its ancestor was polymorphic by this gene, and vice versa. When further divergence of the lineages takes place, some gene versions inherited from the polymorphic ancestor may be lost owing to genetic drift or selection (Rogers and Gibbs 2014). Another explanation of the revealed polymorphism may be a gene flow through hybridization between different species, but this assumption requires more thorough studies.

Insufficient variability in the 5'-end barcoding region of CO1

To reconstruct the CO1 phylogeny of Polymastiidae we used “Folmer’s” barcoding region, which was successfully applied to recover phylogenies of many invertebrate taxa (Folmer et al. 1994), in particular two large sponge families, Geodiidae (Cárdenas et al. 2010) and Halichondriidae (Erpenbeck et al. 2012). Our results revealed the insufficiency of variation of this region in Polymastiidae that might cause some inconsistencies between the CO1 and 28S rDNA

phylogenies (see above) and also hindered the separation of the species in Clades II and III based on CO1 alone, while these species were otherwise successfully separated by the 28S rDNA data. The similar problem with the “Folmer’s” region was reported for some other sponge families, e.g., Lubomirskiidae (Schröder et al. 2002), Clionidae (Ferrario et al. 2010), and Irciniidae (Pöppe et al. 2010). To solve this problem, sequencing of an additional downstream region of the CO1 gene providing more variability was recommended (Erpenbeck et al. 2006, Sponge Barcoding Project at <http://www.palaeontologie.geo.uni-muenchen.de/SBP/>). The analyses of the extended CO1 barcoding region may probably reduce the inconsistency between the CO1 and 28S rDNA phylogenies of Polymastiidae and resolve the relationships between sibling polymastiid species.

Conclusions

Our study presents the first comprehensive phylogenetic reconstruction of the family Polymastiidae based on molecular data. Our results show that its classification based on morphology is in a strong conflict with molecular phylogenies. Accordingly, the majority of previously assumed morphological synapomorphies appear to be highly homoplastic, and a natural classification of Polymastiidae will require a thorough and comprehensive taxonomic revision. Here we have set up a sound molecular framework for this task and recovered several strongly supported clades. In order to determine the morphological synapomorphies of these clades, a reinterpretation of the currently used characters and a selection of additional characters are needed. Furthermore, we have reported evidence for that sorting of lineages of different genes may follow different ways under the evolutionary divergence of sponge species and that the gene flow between populations of recently diverged species may also take place. Finally, we have demonstrated that the standard 5'-end barcoding region of CO1 provides insufficient data that may result in some inconsistency between the CO1 and 28S rDNA phylogenies and failure to reconstruct the relationships between some polymastiid species, which are otherwise recovered with 28S rDNA data. Hence, we argue once again for the advantages of multigene datasets and extended barcoding regions for reconstructing of phylogenies at the family and generic level.

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Authors' contributions Conceiving the study: AP and HTR. Sampling the material: AP and HTR. Primary treatment and taxonomic identification of the samples: AP. DNA extraction, amplification and sequencing: AP. Contributing reagents/consumables/tools: HTR. Reconstruction of the secondary structure, 28S rDNA: OV. Alignments and phylogenetic analyses: OV, AP and EW. Tracing morphological characters along the molecular tree: EW and AP. Writing the manuscript: AP. Revising the manuscript: OV, EW and HTR.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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