Phylogenetic relationships within the Phyllidiidae (Opisthobranchia, Nudibranchia)

Bart E.M.W. Stoffels¹,², Sancia E.T. van der Meij²,³, Bert W. Hoeksema², Joris van Alphen², Theo van Alen¹, Maria Angelica Meyers-Muñoz¹, Nicole J. de Voogd², Yosephine Tutì⁴, Gerard van der Velde¹,²

¹ Radboud University Nijmegen, Institute for Water and Wetland Research, Heyendaalseweg 135, 6525 AJ Nijmegen, The Netherlands
² Naturalis Biodiversity Center, PO Box 9517, 2300 RA Leiden, The Netherlands
³ Oxford University Museum of Natural History, Parks Road, Oxford OX1 3PW, United Kingdom
⁴ Research Center for Oceanography (RCO), Indonesian Institute of Science (LIPI), Jl. Pasir Putih I, Ancol Timur, Jakarta 14430, Indonesia

Corresponding author: Bert W. Hoeksema (bert.hoeksema@naturalis.nl)

Abstract

The Phyllidiidae (Gastropoda, Heterobranchia, Nudibranchia) is a family of colourful nudibranchs found on Indo-Pacific coral reefs. Despite the abundant and widespread occurrence of many species, their phylogenetic relationships are not well known. The present study is the first contribution to fill the gap in our knowledge on their phylogeny by combining morphological and molecular data. For that purpose 99 specimens belonging to 16 species were collected at two localities in Indonesia. They were photographed and used to make a phylogeny reconstruction based on newly obtained cytochrome oxidase subunit (COI) sequences as well as sequence data from GenBank. All mitochondrial 16S sequence data available from GenBank were used in a separate phylogeny reconstruction to obtain information for species we did not collect. COI data allowed the distinction of the genera and species, whereas the 16S data gave a mixed result with respect to the genera Phyllidia and Phyllidiella. Specimens which could be ascribed to species level based on their external morphology and colour patterns showed low variation in COI sequences, but there were two exceptions: three specimens identified as Phyllidia cf. babai represent two to three different
species, while Phyllidiella pustulosa showed highly supported subclades. The barcoding marker COI also confirms that the species boundaries in morphologically highly variable species such as Phyllidia elegans, P. varicosa, and Phyllidiopsis krempfi, are correct as presently understood. In the COI as well as the 16S cladogram Phyllidiopsis cardinalis was located separately from all other Phyllidiidae, whereas Phyllidiopsis fissuratus was positioned alone from the Phyllidiella species by COI data only. Future studies on phyllidiid systematics should continue to combine morphological information with DNA sequences to obtain a clearer insight in their phylogeny.

**Keywords**
COI, Indonesia, mtDNA, nudibranch, phylogenetic relations, 16S

**Introduction**

Nudibranch gastropod molluscs have traditionally been classified with the Infraclass Opisthobranchia Milne Edwards, 1848, which consists of more than 6000 species (Yonow 2008). Although this taxon is not monophyletic and therefore is considered obsolete (Schrödl et al. 2011), taxonomic works still refer to “opisthobranchs” for practical reasons (e.g. Uribe et al. 2013) and Opisthobranchia is considered an “Informal Group” among the Heterobranchia (Wägele et al. 2014). These animals form, ecologically and morphologically, one of the most diverse groups of marine gastropods (Wägele et al. 2014). To avoid use of their misnomer, this well-known group of marine animals can also be referred to as sea slugs (Yonow 2015). Among these, the Nudibranchia Cuvier, 1817 form the largest order with an estimated number of more than 2000 species (Gosliner et al. 2008), although also estimates of nearly 3000 species are known (Vonnemann et al. 2005).

Much work has already been done to elucidate the phylogeny of the opisthobranchs by molecular analyses (e.g., Wollscheid and Wägele 1999, Grande et al. 2004a, 2004b, Vonnemann et al. 2005, Turner and Wilson 2008, Maeda et al. 2010, Pola and Gosliner 2010), but most of the phylogenetic relationships still remain unclear at family, genus, and species level, especially with regards to the nudibranchs. All nudibranch species and many other sea slugs are predators, which usually can be observed together with their prey (Behrens 2005, Pola and Gosliner 2010, van Alphen et al. 2011). Only rarely they are found together with potential predators such as sea anemones, mushroom corals, and pycnogonids (Piel 1991, Behrens 2005, van der Meij and Reijnen 2012, Mehrotra et al. 2015).

The present study aims to clarify the phylogenetic relationships within the Phyllidiidae Rafinesque, 1814, belonging to the Doridacea (Bouchet and Rocroi 2005). This family consists of more than 100 species divided over five genera: Ceratophyllidia Eliot, 1903, Phyllidia Cuvier, 1797, Phyllidiella Bergh, 1869, Phyllidiopsis Bergh, 1875, and Reticulidia Brunckhorst, 1990 (Bouchet 2015). The genera Fryeria JE Gray, 1853, and Reyfria Yonow, 1986, have been synonymised with Phyllidia (Valdés and Gosliner 1999).
Most nudibranchs of the family Phyllidiidae are commonly encountered on coral reefs, where they can easily be noticed because of their aposomatic colouration, which serves to deter possible predators from eating them (Ritson-Williams and Paul 2007). Nevertheless, only eight phyllidiid COI sequences can be found in GenBank, as well as two 18S sequences and 17 16S sequences. There are only a few published studies that incorporate even a single member of Phyllidiidae into a phylogenetic tree (e.g. Wollscheid-Lengeling et al. 2001) and even fewer deal with phylogenetic relationships among Phyllidiidae. Among the latter, most are using anatomical characters (Brunckhorst 1993, Valdés and Gosliner 1999, Valdés 2001, 2002) and only two are known to include a molecular and phylogenetic analysis (Valdés 2003, Cheney et al. 2014).

Phyllidiid slugs are characterized by their oval elongate and tough bodies, which generally possess hard notal tubercles on the dorsal side. Although their colouration is a main character used for their identification, many species cannot be identified based on colouration alone owing to their high intra-specific colour variation. Structure and pattern of the notal tubercles are important characters for identification. Other distinctive features of the Phyllidiidae are the retractile lamellate rhinophores, the compact digestive gland mass, and the triaulic reproductive system (Brunckhorst 1993). Another important character diagnosing the Phyllidiidae is the possession of numerous subdermal calcareous spicules of different microstructures (Chang et al. 2013). The Phyllidiidae have no jaws or radula and lack the dorsal, circumanal circlet of gills that is typical of other dorids (Brunckhorst 1993).

To study the phylogenetic relationships within the Phyllidiidae, a molecular analysis was performed based on DNA sequence data of the mitochondrial cytochrome oxidase I (COI) gene, combined with external morphological assessments of material collected in two areas in eastern Indonesia, the Raja Ampat islands (West Papua) and Ternate, off western Halmahera (Moluccas). Both locations are situated in the centre of maximum marine biodiversity, also known as the Coral Triangle (Hoeksema 2007). In earlier studies, high numbers of phyllidiid species were recorded from this area: 13 from the Bismarck Sea, Papua New Guinea (Domínguez et al. 2007), eleven from Ambon (Moluccas, Indonesia) (Yonow 2011), and eleven from the South China Sea (Sachidhanandam et al. 2000). Therefore, both of our areas were expected to show a high number of phyllidiid species that could be used for the present study.

**Materials and methods**

**Sampling**

Specimens were collected by SCUBA diving in West Papua by Gerard van der Velde in 2007, mostly in the coastal areas of Gam, Kri, Mansuar, and Batanta (Figures 1–2; see Hoeksema and van der Meij 2008). Additional specimens were mainly collected by Joris van Alphen and Nicole de Voogd, and also by Bert Hoeksema, Sancia van der Meij, and other expedition members (Hoeksema and van der Meij 2010) in
Figure 1. Location of field areas: Halmahera (including Ternate) and West Papua (including Raja Ampat).

Figure 2. Raja Ampat sites where Phyllidiidae were sampled in 2007.
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Figure 3. Halmahera and Ternate sites where Phyllidiidae were sampled in 2009.

2009 off Halmahera (northern Moluccas), especially around Ternate (Figures 1, 3). A locality list of the sampling stations is provided in Table 1. Collected slugs were first photographed and subsequently preserved in 96% ethanol (West Papua 2007). Halmahera specimens were transferred into fresh 96% ethanol and labelled in order to prepare them for DNA analysis. These have been deposited in the mollusc collection of Naturalis Biodiversity Center, Leiden (coded as RMNH.Mol.), with the exception of some specimens that dried out after sequencing (Table 1; Figures 5–15; Suppl. material 1: COI sequences).
Table 1. Information on analysed Phyllidiidae species: RMNH.MOL catalogue number or field code number in case voucher specimen became lost; Genbank number if available; collection site, station number (RAJ = Raja Ampat, TER=Ternate, Halmahera), coordinates.

| RMNH.MOL or Field nr. | Genbank accession number | Species | Locality | Station | Coordinates               |
|-----------------------|--------------------------|---------|----------|---------|---------------------------|
| 336464                | KX235918                 | Phyllidia babai | Tanjung Ebamadu | TER08   | N0°45’23.4", E127°24’26.5" |
| 336575                | KX235920                 | Phyllidia cf. babai | South Gam, shoal near mangroves | RAJ37   | S0°31’08.2", E130°38’28.0" |
| 336614                | KX235919                 | Phyllidia cf. babai | Tanjung Ratemu (South of river) | TER27   | N0°54’44.5", E127°29’09.9" |
| 336573                | KX235921                 | Phyllidia coelestis | Eastern entrance of passage | RAJ44   | S0°25’44.3", E130°33’56.8" |
| 336574                | KX235922                 | Phyllidia coelestis | Wallace Lake | RAJ13   | S0°26’31.1", E130°41’08.0" |
| 58                    |                          | Phyllidia elegans | Pulau Maka | TER13   | N0°54’42.7", E127°18’32.9" |
| 137                   |                          | Phyllidia elegans | Pulau Pilongga, North | TER34   | N0°42’49.8", E127°28’45.4" |
| 156                   |                          | Phyllidia elegans | Teluk Dodinga: Karang Ngeli West | TER40   | N0°46’25.3", E127°32’22.9" |
| 336475                | KX073972                 | Phyllidia elegans | Tanjung Tabam | TER12   | N0°50’05.1", E127°23’10.0" |
| 336478                | KX073973                 | Phyllidia elegans | Pulau Maka | TER13   | N0°54’42.7", E127°18’32.9" |
| 336488                | KX073974                 | Phyllidia elegans | Tanjung Pasir Putih | TER16   | N0°51’50.4", E127°20’36.7" |
| 336514                | KX073975                 | Phyllidia elegans | Dufadufa / Benteng Toloko | TER24   | N0°48’49.1", E127°23’21.6" |
| 336515                | KX073976                 | Phyllidia elegans | Idem | TER24   | N0°48’49.1", E127°23’21.6" |
| 336554                | KX073985                 | Phyllidia elegans | Passage | RAJ43   | S0°25’45.2", E130°33’37.3" |
| 336555                | KX073990                 | Phyllidia elegans | Akber Reef | RAJ14   | S0°34’15.2", E130°39’33.7" |
| 336556                | KX073988                 | Phyllidia elegans | Passage | RAJ43   | S0°25’45.2", E130°33’37.3" |
| 336557                | KX073987                 | Phyllidia elegans | Idem | RAJ43   | S0°25’45.2", E130°33’37.3" |
| 336558                | KX073984                 | Phyllidia elegans | Southwest Pulau Kri | RAJ40   | S0°33’58.1", E130°39’46.2" |
| 336559                | KX073991                 | Phyllidia elegans | South Gam, shoal near mangroves | RAJ37   | S0°31’08.2", E130°38’28.0" |
| 336560                | KX073983                 | Phyllidia elegans | Southwest Pulau Kri | RAJ40   | S0°33’58.1", E130°39’46.2" |
| 336561                | KX073986                 | Phyllidia elegans | Passage | RAJ43   | S0°25’45.2", E130°33’37.3" |
| 336562                | KX073989                 | Phyllidia elegans | Akber Reef | RAJ14   | S0°34’15.2", E130°39’33.7" |
| 336628                | KX073977                 | Phyllidia elegans | Pulau Gura Ici, East | TER32   | S0°01’17.3", E127°14’17.2" |
| 336629                | KX073978                 | Phyllidia elegans | Idem | TER32   | S0°01’17.3", E127°14’17.2" |
| 336631                | KX073979                 | Phyllidia elegans | Pulau Pilongga, North | TER34   | N0°42’49.8", E127°28’45.4" |
| 336632                | KX073980                 | Phyllidia elegans | Idem | TER34   | N0°42’49.8", E127°28’45.4" |
| Genbank accession number | Species                     | Locality                                | Station | Coordinates          |
|--------------------------|----------------------------|-----------------------------------------|---------|----------------------|
| KX073981                 | *Phyllidia elegans*        | Teluk Dodinga, Kening Ngebi West       | TER34   | N°4°22.2974, E127°27.53°37.5" |
| KX073982                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER40   | N°4°21.9874, E127°27.38°40.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| KX253526                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER19   | N°4°34.6674, E127°22.86°35.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| KX253526                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER19   | N°4°34.6674, E127°22.86°35.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| KX253526                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER19   | N°4°34.6674, E127°22.86°35.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| KX253526                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER19   | N°4°34.6674, E127°22.86°35.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| KX253526                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER19   | N°4°34.6674, E127°22.86°35.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| KX253526                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER19   | N°4°34.6674, E127°22.86°35.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| KX253526                 | *Phyllidia elegans*        | South of Tidore, Tanjung Ngebi         | TER19   | N°4°34.6674, E127°22.86°35.7" |
| KX253525                 | *Phyllidia exoleta*        | Northeast Gam, Frwen Wonda              | TER14   | N°4°34.6674, E127°22.86°35.7" |
| KX253524                 | *Phyllidia exoleta*        | Taufinat Ngebi, Northwest of Tobola    | TER19   | N°4°34.4074, E127°22.69°30.7" |
| Genbank accession number | Species | Locality | Coordinates |
|--------------------------|---------|----------|-------------|
| KX235938                 | Phyllidiella nigra | Idem | TER10 N°44°32.0' E127°31.5° |
| KX235949                 | Phyllidiella nigra | Idem | TER12 N°44°32.0' E127°31.5° |
| KX235950                 | Phyllidiella nigra | Idem | TER13 N°44°32.0' E127°31.5° |
| KX235951                 | Phyllidiella nigra | Idem | TER14 N°44°32.0' E127°31.5° |
| KX235952                 | Phyllidiella nigra | Idem | TER15 N°44°32.0' E127°31.5° |
| KX235953                 | Phyllidiella nigra | Idem | TER16 N°44°32.0' E127°31.5° |
| KX235954                 | Phyllidiella nigra | Idem | TER17 N°44°32.0' E127°31.5° |
| KX235955                 | Phyllidiella nigra | Idem | TER18 N°44°32.0' E127°31.5° |
| KX235956                 | Phyllidiella nigra | Idem | TER19 N°44°32.0' E127°31.5° |
| KX235957                 | Phyllidiella nigra | Idem | TER20 N°44°32.0' E127°31.5° |
| KX235958                 | Phyllidiella nigra | Idem | TER21 N°44°32.0' E127°31.5° |
| KX235959                 | Phyllidiella nigra | Idem | TER22 N°44°32.0' E127°31.5° |
| KX235960                 | Phyllidiella nigra | Idem | TER23 N°44°32.0' E127°31.5° |
| KX235961                 | Phyllidiella nigra | Idem | TER24 N°44°32.0' E127°31.5° |
| KX235962                 | Phyllidiella nigra | Idem | TER25 N°44°32.0' E127°31.5° |
| KX235963                 | Phyllidiella nigra | Idem | TER26 N°44°32.0' E127°31.5° |
| KX235964                 | Phyllidiella nigra | Idem | TER27 N°44°32.0' E127°31.5° |
| KX235965                 | Phyllidiella nigra | Idem | TER28 N°44°32.0' E127°31.5° |
| KX235966                 | Phyllidiella nigra | Idem | TER29 N°44°32.0' E127°31.5° |
| KX235967                 | Phyllidiella nigra | Idem | TER30 N°44°32.0' E127°31.5° |
| KX235968                 | Phyllidiella nigra | Idem | TER31 N°44°32.0' E127°31.5° |
| KX235969                 | Phyllidiella nigra | Idem | TER32 N°44°32.0' E127°31.5° |
| KX235970                 | Phyllidiella nigra | Idem | TER33 N°44°32.0' E127°31.5° |
| KX235971                 | Phyllidiella nigra | Idem | TER34 N°44°32.0' E127°31.5° |
| KX235972                 | Phyllidiella nigra | Idem | TER35 N°44°32.0' E127°31.5° |
| KX235973                 | Phyllidiella nigra | Idem | TER36 N°44°32.0' E127°31.5° |
| KX235974                 | Phyllidiella nigra | Idem | TER37 N°44°32.0' E127°31.5° |
| KX235975                 | Phyllidiella nigra | Idem | TER38 N°44°32.0' E127°31.5° |
| KX235976                 | Phyllidiella nigra | Idem | TER39 N°44°32.0' E127°31.5° |
| KX235977                 | Phyllidiella nigra | Idem | TER40 N°44°32.0' E127°31.5° |
| KX235978                 | Phyllidiella nigra | Idem | TER41 N°44°32.0' E127°31.5° |
| KX235979                 | Phyllidiella nigra | Idem | TER42 N°44°32.0' E127°31.5° |
| KX235980                 | Phyllidiella nigra | Idem | TER43 N°44°32.0' E127°31.5° |
| KX235981                 | Phyllidiella nigra | Idem | TER44 N°44°32.0' E127°31.5° |
| KX235982                 | Phyllidiella nigra | Idem | TER45 N°44°32.0' E127°31.5° |
| KX235983                 | Phyllidiella nigra | Idem | TER46 N°44°32.0' E127°31.5° |
| KX235984                 | Phyllidiella nigra | Idem | TER47 N°44°32.0' E127°31.5° |
| KX235985                 | Phyllidiella nigra | Idem | TER48 N°44°32.0' E127°31.5° |
| KX235986                 | Phyllidiella nigra | Idem | TER49 N°44°32.0' E127°31.5° |
| KX235987                 | Phyllidiella nigra | Idem | TER50 N°44°32.0' E127°31.5° |
| KX235988                 | Phyllidiella nigra | Idem | TER51 N°44°32.0' E127°31.5° |
| KX235989                 | Phyllidiella nigra | Idem | TER52 N°44°32.0' E127°31.5° |
| KX235990                 | Phyllidiella nigra | Idem | TER53 N°44°32.0' E127°31.5° |
| RMNH.MOL or Genbank accession number | Species | Locality | Station | Coordinates |
|--------------------------------------|---------|----------|---------|-------------|
| 336462  KX235973                    | Phyllidiopsis krempfi | Tanjung Ebamadu | TER08   | N0°45'23.4", E127°24'26.5" |
| 336466  KX235974                    | Phyllidiopsis krempfi | Idem | TER08   | N0°45'23.4", E127°24'26.5" |
| 336469  KX235975                    | Phyllidiopsis krempfi | West Maitara | TER09   | N0°43'47.6", E127°21'44.7" |
| 336512  KX235976                    | Phyllidiopsis krempfi | Dufadufa / Benteng Toloko | TER24   | N0°48'49.1", E127°23'21.6" |
| 336594  KX235979                    | Phyllidiopsis krempfi | Southwest Pulau Kre, Kuburan | RAJ15   | S0°33'42.8", E130°39'40.4" |
| 336595  KX235984                    | Phyllidiopsis krempfi | Southwest Pulau Kre | RAJ40   | S0°33'58.1", E130°39'46.2" |
| 336596  KX235983                    | Phyllidiopsis krempfi | Northwest Pulau Mansuar, Laios reef | RAJ49   | S0°32'53.5", E130°29'51.1" |
| 336597  KX235978                    | Phyllidiopsis krempfi | Southwest Pulau Kre, Kuburan | RAJ15   | S0°33'42.8", E130°39'40.4" |
| 336598  KX235980                    | Phyllidiopsis krempfi | North Batanta, North Pulau Yatiri | RAJ28   | S0°46'46.7", E130°42'42.7" |
| 336599  KX235982                    | Phyllidiopsis krempfi | East Kre, Sorido Wall | RAJ12   | S0°33'13.2", E130°41'16.9" |
| 336600  KX235981                    | Phyllidiopsis krempfi | Northeast Mansuar | RAJ38   | S0°34'05.0", E130°38'31.5" |
| 336650  KX235977                    | Phyllidiopsis shireenae | Teluk Dodinga: West Karang Ngeli | TER40   | N0°46'25.3", E127°32'22.0" |
| 336451  KX235985                    | Phyllidiopsis shireenae | Kampung Gina / Tapak 2 | TER06   | N0°47'15.0", E127°23'25.0" |
| 336652  KX235986                    | Phyllidiopsis shireenae | Teluk Dodinga: East Karang Luelue | TER41   | N0°46'32.8", E127°33'43.4" |
| 336591  KX235987                    | Phyllidiopsis xishaensis | Southeast Gam, Pulau Kerupir, Mike's Point | RAJ05   | S0°30'57.1", E130°40'22.1" |
| 336592  KX235988                    | Phyllidiopsis xishaensis | East Pulau Kre, Cape Kre | RAJ07   | S0°33'22.2", E130°41'28.7" |
| 336593  KX235989                    | Phyllidiopsis xishaensis | Eastern entrance of passage | RAJ44   | S0°25'44.3", E130°33'56.8" |
| 336640  KX235990                    | Reticulidia fungia | East Teluk Dodinga: North of Pulau Jere | TER36   | N0°50'47.8", E127°37'48.7" |
| 336455  KX235991                    | Reticulidia bulgerda | Kampung Gina / Tapak 2 | TER06   | N0°47'15.0", E127°23'25.0" |
Figure 4. Phylogeny reconstruction of the Phyllidiidae based on COI gene sequence data of 109 specimens (including outgroups). Topology derived from Bayesian inference 50% majority rule, significance values are posterior probabilities / bootstrap values. Numbers refer to GenBank accession numbers / RMNH.Moll catalogue numbers.

Morphological study

Collected specimens were identified according to their external morphology using Brunckhorst (1993), Yonow et al. (2002), and Yonow (2011). In addition, field guides showing in situ photographs were used (Gosliner et al. 2008). All individuals except
for three could be identified to species level. All specimens were photographed alive or in the preserved state (Figures 5–15); these photos can be linked to the phylogeny reconstruction of the Phyllidiidae based on COI gene sequence data (Figure 4).
Figure 6. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidia elegans*. Order of specimens (a–i) according to Figure 4 (d dorsal and ventral sides). Numbers refer to RMNH.Moll catalogue numbers and locality codes (137 and 156, dried-out).
Figure 7. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidia elegans* (a–f), *Phyllidia* sp. (g dorsal and ventral sides), *P. exquisita* (h), *P. coelastis* (i). Order of specimens (a–i) according to Figure 4. Numbers refer to RMNH.Moll catalogue numbers or locality code (058, dried-out).
Figure 8. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidia coelestis* (a), *P. varicosa* (b–i). Order of specimens (a–i) according to Figure 4 (d dorsal and ventral sides). Numbers refer to RMNH.Moll catalogue numbers.
Figure 9. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidia varicosa* (a–f), *P. ocellata* (g–i). Order of specimens (a–i) according to Figure 4 (c dorsal and ventral sides). Numbers refer to RMNH.Moll catalogue numbers or locality code (074, dried-out).
Figure 10. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidia picta* (a–c), *Phyllidia babai* (d), *Phyllidia cf. babai* (e–f), *Reticulidia fungia* (g), *Reticulidia halgerda* (h), *Phyllidiopsis fissuratus* (i). Order of specimens (a–i) according to Figure 4 (e dorsal and ventral sides). Numbers refer to RMNH.Moll catalogue numbers.
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Figure 11. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: Phyllidiella rudmani (a), Phyllidiella nigra (b–h), Phyllidiella pustulosa (i–j). Order of specimens (a–j) according to Figure 4. Numbers refer to RMNH.Moll catalogue numbers.
Figure 12. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: _Phyllidiella pustulosa_. Order of specimens (a–j) according to Figure 4. Numbers refer to RMNH.Moll catalogue numbers or locality code (75F, dried-out).
Figure 13. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidiella pustulosa* (a–h), *Phyllidiopsis xishaensis* (i–j). Order of specimens (a–j) according to Figure 4. Numbers refer to RMNH.Moll catalogue numbers.
Figure 14. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidiopsis xishaensis* (a), *Phyllidiopsis shireenae* (b–c), *Phyllidiopsis krempfi* (d–i). Order of specimens (a–i) according to Figure 4 (c dorsal and ventral sides). Numbers refer to RMNH.Moll catalogue numbers.
Figure 15. External morphology and colouration of Phyllidiidae specimens used for COI phylogeny reconstruction: *Phyllidiopsis krempfi*. Order of specimens (a–g) according to Figure 4 (f, g dorsal and ventral sides). Numbers refer to RMNH.Moll catalogue numbers.
DNA extraction

For each species encountered in the field surveys one or more individuals were chosen for DNA analysis as well as from the morphologically distinct unidentified specimens, resulting in a total of 99 samples (Table 1). DNA was extracted from tissue of small foot fragments with the DNeasy Blood & Tissue Kit (Qiagen, Germany) according to the manufacturer’s protocol. DNA was eluted in DEPC treated water. The quality of the extracted DNA was tested by agarose gel (0.7%) electrophoresis.

PCR amplification, purification, and sequencing

Extracted DNA was used for Polymerase Chain Reaction (PCR) to amplify fragments of the mitochondrial gene COI (cytochrome c oxidase subunit 1). The primers used for the amplification of the COI gene were: LCO1490 (5’GGT CAA CAA ATC ATA AAG ATA TTG G 3’) and HCO2198 (5’TAA ACT TCA GGG TGA CCA AAA AAT CA 3’) (Folmer et al. 1994). Thermal cycling conditions used for the amplification of the COI gene were: initial denaturing at 94 °C for 3 min followed by 38 amplification cycles of denaturation at 94 °C for 15 sec, primer annealing at 50 °C for 30 sec, and elongation at 72 °C for 1 min. A final elongation step at 72 °C for 5 min was performed. After checking by agarose (1%) electrophoresis if the PCR resulted the unique PCR fragments of the expected size (approximately 658 bp), the fragments were purified using the GeneJET PCR Purification Kit (Thermo Scientific, Landsmeer, NL). Purified PCR products were sequenced with corresponding primers.

Sequence alignment and phylogenetic analyses

The quality of the sequences was checked using Chromas Lite (Technelysium Pty Ltd.). Subsequently the sequences were edited in MEGA 6 (Tamura et al. 2013) and analysed by BLAST searches (http://www.ncbi.nlm.nih.gov). COI sequences of Dermatodes citrina (Cheeseman, 1881) and Doriopsilla areolata Bergh, 1880 were collected from GenBank and used as outgroups. Additional COI sequences of Phyllidia coelestis Bergh, 1905, Phyllidia elegans Bergh, 1869, Phyllidia ocellata Cuvier, 1804, Phyllidia picta Pruvot-Fol, 1957, Phyllidia varicosa Lamarck, 1801, Phyllidiella lizae Brunckhorst, 1993, Phyllidiella pustulosa (Cuvier, 1804), Phyllidiopsis cardinalis Bergh, 1875 were obtained from GenBank (Table 2).

The newly obtained COI sequences and the sequences from GenBank were aligned using the Guidance server (Clustal W; Penn et al. 2010), resulting in an alignment score of 1.000. There were no unreliable columns. Prior to the model-based phylogenetic analysis, the best-fit model of nucleotide substitution was identified by means of the Akaike Information Criterion (AIC) calculated with jModeltest (Posada 2008), resulting in TVM+I+G as the most suitable model. Phylogenetic reconstructions were
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Table 2. Mitochondrial COI sequences of Phyllidiidae (and outgroups) obtained from GenBank.

| Species                  | Accession number | Reference                           | Collection locality                  |
|--------------------------|------------------|-------------------------------------|--------------------------------------|
| Dendrodoris citrina      | GQ292043         | Shields et al. (2009 unpublished)   | Ross Sea, Antarctica?                |
| Doriopsilla areolata     | AJ223262         | Thollesson (2000)                   | Cadiz, Andalusia, Spain              |
| Phyllidia coelestis      | KJ001305         | Cheney et al. (2014)                | Lizard I., Queensland Australia      |
| Phyllidia elegans        | AJ223276         | Thollesson (2000)                   | Tab I., Papua New Guinea             |
| Phyllidia ocellata       | KJ001307         | Cheney et al. (2014)                | Mooloolaba, Queensland, Australia    |
| Phyllidia picta          | KJ001304         | Cheney et al. (2014)                | Lizard I., Queensland Australia      |
| Phyllidia varicosa       | KJ001306         | Cheney et al. (2014)                | Lizard I., Queensland Australia      |
| Phyllidiella lizae       | KJ001309         | Cheney et al. (2014)                | Lizard I., Queensland Australia      |
| Phyllidiella pustulosa   | KJ001310         | Cheney et al. (2014)                | Lizard I., Queensland Australia      |
| Phyllidiopsis cardinalis | KJ001308         | Cheney et al. (2014)                | Mooloolaba, Queensland, Australia    |

Initial phylogenetic analyses showed high intraspecific variation on the COI region between specimens identified as *Phyllidiella pustulosa*. Tests to estimate the average evolutionary divergence over sequence pairs between and within groups were carried out in MEGA 6.06. *Phyllidia elegans, P. varicosa, Phyllidiella nigra* (van Hasselt, 1824), *P. pustulosa*, and *Phyllidiopsis krempfi* Pruvot-Fol, 1957 were used as representatives for each of the species groups, because of the larger number of available sequences for these species. The *Phyllidiella pustulosa* sequence from GenBank (KJ001310) was excluded from this analysis: based on its position in the phylogeny reconstruction the identification of this specimen as *P. pustulosa* is doubtful. The web version of ABGD (Automatic Barcode Gap Discovery, Puillandre et al. 2012) was used to estimate the genetic distance corresponding to the difference between a speciation process versus intra-specific variation in *Phyllidiella pustulosa*. Runs were performed using the default range of priors (pmin = 0.001, pmax = 0.10) using the JC69 Jukes-Cantor measure of distance. The analysis involved 20 nucleotide sequences with a total of 588 positions in the final dataset.

All available mitochondrial 16S sequences of Phyllidiidae on GenBank (Thollesson 2000, Wolfscheid-Lengeling et al. 2001, Valdés 2003, Cheney et al. 2014, Shields et al. unpublished) were used for a phylogeny reconstruction based on this marker, which allowed us to study the phylogenetic position of 17 phyllidiid species including two species (*Phyllidia rueppelii* (Bergh, 1869) and *Phyllidiopsis sphingis* Brunckhorst, 1993) for which no COI data were available. *Doriopsilla albopunctata* (JG Cooper, 1863) was used as outgroup (Table 3). The sequences were aligned using the Guidance server.
Table 3. 16S sequences of Phyllidiidae obtained from GenBank.

| Species                      | Accession number | Reference                  | Collection locality            |
|------------------------------|------------------|----------------------------|--------------------------------|
| Doropsilla albopunctata      | AF430354         | Valdés (2003)              | Baja California, Mexico       |
| Phyllidia coelestis          | AF430361         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidia coelestis          | KJ018917         | Cheney et al. (2014)       | Lizard I., Queensland Australia|
| Phyllidia elegans            | AF430362         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidia elegans            | AJ225201         | Thollesson (2000)          | Tab I., Papua New Guinea      |
| Phyllidia ocellata           | AF430363         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidia picta              | KJ018916         | Cheney et al. (2014)       | Lizard I., Queensland Australia|
| Phyllidia rueppelli          | AF430358         | Valdés (2003)              | Hurghada, Egypt               |
| Phyllidiella lizae           | AF430365         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidiella lizae           | KJ018918         | Cheney et al. (2014)       | Lizard I., Queensland Australia|
| Phyllidiella pustulosa       | AF249232         | Wollscheid-Lengeling et al. (2001) | Great Barrier Reef, Australia |
| Phyllidiella pustulosa       | AF430366         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidia varicosa           | AF430364         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidiopsis cardinalis     | AF430367         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidiopsis sphingis       | AF430368         | Valdés (2003)              | Lifou I., New Caledonia       |
| Phyllidiopsis xishaensis*    | AF430369         | Valdés (2003)              | Lifou I., New Caledonia       |
| Reticulidia fungia          | AF430370         | Valdés (2003)              | Lifou I., New Caledonia       |
| Reticulidia halgerda         | AF430371         | Valdés (2003)              | Lifou I., New Caledonia       |

* Re-identification according to Yonow (pers. comm.)

(ClustalW; Penn et al. 2010), resulting in an alignment score of 0.996281. All unreliable columns (confidence score below 0.93) were removed. Prior to the model-based phylogenetic analysis, the best-fit model of nucleotide substitution was identified by means of the Akaike Information Criterion (AIC) calculated with jModeltest (Posada 2008), resulting in TVM+I+G. Because of the unavailability of TVM in MrBayes 3.1.2 (Ronquist and Huelsenbeck 2003), we used the most complex GTR+I+G model of nucleotide substitution. Bayesian inferences coupled with MCMC techniques (six chains) were run for 3,000,000 generations, with a sample tree saved every 1000 generations and the burnin set to 25%. Likelihood scores stabilized at a value of 0.005654. Consensus trees were visualized in FigTree v.1.3.1 (Rambaut 2009). A maximum likelihood analysis (GTR+I+G; 1000 bootstraps) was carried out with Phyml 3.1 (Guindon et al. 2010) using the Seaview platform (Gouy et al. 2010).

Results and discussion

Position of genera

The reconstruction based on COI (Figure 4) is derived from the Bayesian inference 50% majority rule consensus. This topology is congruent with the one resulting from
the maximum likelihood analysis. Three large groupings can be discerned (indicated as A, B, and C in Figure 4), albeit with low support for the higher taxonomic levels. The support values in the distal branches are high. The genera *Phyllidia*, *Phyllidiella*, *Phyllidiopsis*, and *Reticulidia* are retrieved in distinct clades, with *Reticulidia* as a sister clade to *Phyllidia*. *Phyllidiopsis fissuratus* Brunckhorst, 1993 formed a separate lineage basal to *Phyllidiella* species (albeit without support). *Phyllidiopsis cardinalis* does not cluster with its congeners, but instead forms a separate lineage in the *Phyllidiidae*.

The 16S phylogeny reconstruction is also derived from the Bayesian inference 50% majority rule consensus of the trees remaining after the burnin. There are low support values in the basal part of the tree and high support values in the distal phylogenetic branches (Figure 17). The Bayesian inference topology is congruent with the topology resulting from the maximum likelihood analysis. The outgroup *Doriopsilla albopunctata* is separated by a long branch. Within the overall clade four main groupings can be distinguished: *Phyllidiella*, *Phyllidiopsis*, and *Reticulidia*, and a mixed clade of *Phyllidiella* and *Phyllidia*. Based on this analysis only the genus *Reticulidia* is monophyletic. *Phyllidiopsis cardinalis* does not cluster with any of the other analysed taxa, and holds a separate position in the phylogeny reconstruction. The latter is in accordance with the COI reconstruction (Figure 4).

The arrangement of the four phyllidiid genera based on the molecular data (Figures 4, 16a) is similar to that of Brunckhorst (1993) that was based on morphological and anatomical data (Figure 16b). The only exception is the position of the genus *Fryeria*. Brunckhorst (1993) distinguished *Fryeria* from *Phyllidia* based on the position of the anus and other anatomical features. *Phyllidia picta* (with its synonyms *Fryeria picta* (Pruvot-Fol, 1957), *Fryeria menindie* Brunckhorst, 1993, *Phyllidia menindie* (Brunckhorst, 1993)) was included in our analyses which, according to Brunckhorst, should belong to the genus *Fryeria*. Valdés and Gosliner (1999) synonymized both genera, which was later followed by Valdés (2003) and Cheney et al. (2014). The present reconstruction based on COI (Figure 16a) reconfirms the inclusion of *Fryeria* in the genus *Phyllidia*.

The cladogram of the genera based on 16S mtDNA sequence data collected by Valdés (2003) (Figure 16c) is roughly similar to the cladogram based on COI, except for the different positions of *Phyllidiopsis* and *Phyllidiella*. The cladogram based on morphological and anatomical data as shown by Valdés (2002; Figure 16d) is different from the other proposed classifications (Figures 16a–c). Brunckhorst (1993) considered *Ceratophyllidia* a sister group to all the other genera (Figure 6b). Valdés (2002; Figure 16d) distinguished two larger groupings within the *Phyllidiidae*: *Ceratophyllidia* and *Phyllidiopsis* as one group and *Phyllidia*, *Phyllidiella*, and *Reticulidia* as the other group. *Phyllidia* and *Phyllidiella* in turn formed a sister group of *Reticulidia* (Figure 16d). The cladogram by Brunckhorst (1993) and our cladogram based on COI (Figure 4) both show that *Phyllidiella* is a sister clade of *Reticulidia* and *Phyllidia*. In contrast, *Phyllidiella* is not a sister group of *Phyllidia* but to all the other genera grouped together in the cladogram of Valdés (2003).
Figure 16. a Cladogram based on COI gene sequence data showing topology of four genera of Phyllidiidae. b Cladogram according to Brunckhorst (1993) based on morphological data showing topology of six genera of Phyllidiidae. c Cladogram based on 16S mtDNA sequence data showing topology of four genera of Phyllidiidae (Valdés 2003) d Cladogram based on morphological data (Valdés 2002) showing topology of five genera of Phyllidiidae.

Unfortunately no Ceratophyllidia specimens were available to complete our analysis at genus level. Up to this point the phylogenetic position of the genus Ceratophyllidia remains unclear, and additional molecular analyses are necessary to establish its position.
Figure 17. Phylogeny reconstruction of the Phyllidiidae based on 16S mtDNA of 17 specimens of 14 species (including outgroup). Topology derived from Bayesian inference 50% majority rule, significance values are posterior probabilities/bootstrap values. Numbers refer to GenBank accession numbers. *Re-identification according to Yonow (pers. comm.)

Species level analysis

Species level analysis was mainly based on COI (Figure 4). Four nominal species were sequenced in the genus Phyllidiella. Phyllidiella nigra formed a highly supported clade. In the clade containing *P. pustulosa* much variation is visible indicating larger genetic differences among individuals. The ABGD analysis shows that four Molecular Operational Taxonomic Units (MOTUs) are present in *Phyllidiella pustulosa*, suggesting the presence of cryptic species or, alternatively, high intraspecific variation. The *P. pustulosa* of Cheney et al. (2014) falls in between the group consisting of *P. nigra* and *P. pustulosa* on one side and *P. rudmani* Brunckhorst, 1993 on the other and probably represents another species. Our specimen of *P. rudmani* clustered with the specimen identified as *P. lizae* in Cheney et al. (2014). *Phyllidiella rudmani* and *P. lizae* resemble each other (Brunckhorst 1993) and hence it is possible that the species identified as *P. lizae* in Cheney et al. (2014) is in fact *P. rudmani*.
Specimens of seven nominal *Phyllidia* species were sequenced. Sequences of 25 individuals of *Phyllidia elegans* (including one from GenBank) formed a highly supported clade, just like the clades containing *P. ocellata*, *P. picta*, and *P. varicosa*. *Phyllidia coelestis* was also retrieved as a highly supported clade. An individual identified as *P. picta* by Cheney et al. (2014) was part of this group suggesting that it should probably be identified as *P. coelestis*. Brunckhorst (1993) already noticed the close similarity between the two species but still confused them (Yonow 1996), and hence identification errors are likely to occur. Individuals identified as *Phyllidia babai* Brunckhorst, 1993 and *P. cf. babai* were retrieved in two different clades. Specimens 336464 and 336614 differ in 75 base pairs, 336464 and 336575 by 68 base pairs and 336614 and 336575 by 32 base pairs. Differences based on COI suggest that they represent two, or possibly three, different species. The genus *Reticulidia* was retrieved as a sister group of *Phyllidia*.

Material of four nominal species in the genus *Phyllidiopsis* was sequenced, with additional data of one species from GenBank (*P. cardinalis*). *Phyllidiopsis fissuratus* clusters basal to *Phyllidiella*, without support. *Phyllidiopsis shireenae* Brunckhorst, 1990 and *P. xishaensis* (Lin, 1983) cluster as sister species, in highly supported clades. *Phyllidiopsis krempfi* also formed a clear group. *Phyllidiopsis cardinals* does not cluster with any of the phyllidiid genera based on either the 16S or the COI analysis. This result suggests that *P. cardinalis* should be separated from the other *Phyllidiopsis* species, but further morphological analyses are needed to confirm this outcome. Brunckhorst (1993) noted that *P. cardinalis* is the type species of the genus *Phyllidiopsis*, and that it has a unique and complex coloration totally different from that of any other known phyllidiid species, as well as a different anatomy, especially in the foregut. Valdés (2003) states “Additionally, the genus *Phyllidiopsis* is not monophyletic when molecular characters are used, because *Phyllidiopsis cardinals* is at the base of the Phyllidiidae clade, and not nested with the other members of *Phyllidiopsis*”. Surprisingly, in the analysis of Cheney et al. (2014), based on a concatenated dataset of 16S and COI mtDNA, *P. cardinalis* was retrieved in a highly supported clade with several species of *Phyllidiella* and *Phyllidia*.

**Variation within *Phyllidiella pustulosa***

*Phyllidiella pustulosa* is the only species in the COI cladogram (Figure 4) in which highly supported subclades can be discerned. To estimate the average evolutionary divergence within *Phyllidiella pustulosa* the base differences were compared per site for all grouped sequences of the species *Phyllidia elegans* (n = 24), *P. varicosa* (n = 15), *Phyllidiella nigra* (n = 7), *P. pustulosa* (n = 20), and *Phyllidiopsis krempfi* (n = 13) (Tables 4–5).

The genetic variation on the barcoding marker COI is much higher within *Phyllidiella pustulosa* (3.9%) than within the other four species, which showed genetic variations between 0.6 and 1.2% (Table 4). The interspecific genetic variation (involving three different genera) ranges between 10.5 and 18.9% (Table 5). The congeners *Phyl-
Phylogenetic relationships within the Phyllidiidae (Opisthobranchia, Nudibranchia)

Table 4. Estimates of average evolutionary divergence (p-distance) over sequence pairs within groups, in percentages.

| Species               | Distance (%) |
|-----------------------|--------------|
| Phyllidia elegans     | 0.7          |
| Phyllidia varicosa    | 0.7          |
| Phyllidiella nigra    | 0.6          |
| Phyllidiella pustulosa| 3.9          |
| Phyllidiopsis krempfi | 1.2          |

Table 5. Estimates of average evolutionary divergence (p-distance) over sequence pairs between groups, in percentages.

| Distance (%) | Phyllidia elegans | Phyllidia varicosa | Phyllidiella nigra | Phyllidiella pustulosa | Phyllidiopsis krempfi |
|--------------|-------------------|--------------------|--------------------|------------------------|------------------------|
| Phyllidia elegans |                   |                    |                   |                        |                        |
| Phyllidia varicosa | 12.1              |                    |                   |                        |                        |
| Phyllidiella nigra | 15.8              | 15.5               |                   |                        |                        |
| Phyllidiella pustulosa | 18.3              | 18.9               | 10.5               |                        |                        |
| Phyllidiopsis krempfi | 15.8              | 16.4               | 14.6               | 17.2                   |                        |

Phyllidiella nigra and P. pustulosa differ by 10.5%, and the congeners Phyllidia elegans and P. varicosa differ by 12.1%. The observed levels of genetic variation within Phyllidiella pustulosa (Table 4) and between the five species (Table 5) call for additional studies on possible cryptic speciation in P. pustulosa.

Conclusions

The barcoding marker COI works well to separate the different species in the Phyllidiidae, and confirms that the species boundaries in highly variable species, such as Phyllidia elegans, P. varicosa, and Phyllidiopsis krempfi, are correct as presently understood. However, a multi-locus approach, preferably including nuclear markers, is needed to improve the resolution for the higher taxonomic levels. With the exception of a few species that are difficult to place (Phyllidiopsis fissuratus, Phyllidiopsis cardinalis) the studied genera (Phyllidia, Phyllidiella, Phyllidiopsis, and Reticulidia) were retrieved as separate genera within the family. Additional representatives of Ceratophyllidia are needed to indicate the position of this genus within the Phyllidiidae. The observed groupings within Phyllidiella pustulosa suggest that multiple (cryptic) species could be present in this species, for which further analyses are needed including morphological data and multiple markers. Chang and Willan (2015) indicated that at least nine clades could be recognized in Phyllidiella pustulosa that could be separated slightly according
to morphological characters. We recommend that future studies combine DNA sequences with morphological characters, which can easily be done by adding pictures of the specimens to avoid increasing confusion in the identification of specimens.

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Supplementary material I

COI sequences of lost Phyllidiidae specimens
Authors: Bart E.M.W. Stoffels, Sancia E.T. van der Meij, Bert W. Hoeksema, Joris van Alphen, Theo van Alen, Maria Angelica Meyers-Muñoz, Nicole J. de Voogd, Yosephine Tuti, Gerard van der Velde
Data type: Adobe PDF file
Explanation note: COI sequences of Phyllidiidae specimens that dried out after sequencing (numbers and localities are indicated in Table 1.
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