# Molecular phylogeny of the marine snail genus Haminoea (Gastropoda, Cephalaspidea): A framework to study marine diversity and speciation 

Martina Turani ${ }^{1}$ | Ángel Valdés ${ }^{2} \odot \mid$ Manuel António E. Malaquias ${ }^{1}$ ©

${ }^{1}$ Department of Natural History, University Museum of Bergen, University of Bergen, Bergen, Norway
${ }^{2}$ Department of Biological Sciences, California State Polytechnic University, Pomona, California, USA

## Correspondence

Manuel António E. Malaquias, Department of Natural History, University Museum of Bergen, University of Bergen, PB 7800, 5020 Bergen, Norway.
Email: manuel.malaquias@uib.no

## Funding information

Fundação para a Ciência e Tecnologia, Grant/Award Number: SFRH/ BPD/26209/2006; Meltzers Foundation; Ministerio de Ciencia e Innovación, Grant/Award Number: CGL2010-17187 and CTM2008-05228-E/MAR; Spanish Ministry of Education and Science, Grant/Award Number: CGL2006-05182; SYNTHESYS programme, EU;
Universidad de Cádiz, Grant/Award Number: PR2018-039


#### Abstract

Haminoea are herbivorous, coastal snails occurring in temperate and tropical waters of the Atlantic and Eastern Pacific oceans, with one species present in temperate South Africa (Indian Ocean). The genus is taxonomically difficult as several available nominal species were introduced based on shell descriptions alone, or described based on subtle differences in morpho-anatomical features, without a phylogenetic molecular framework. Fifteen species are currently accepted as valid in recent scientific literature and field guides (eight Eastern Atlantic, one temperate Indian Ocean, four Western Atlantic and three Eastern Pacific). Here we generate the first complete phylogeny (Bayesian and Maximum Likelihood) of this genus based on multilocus molecular data (COI, 12S rRNA, 16 S rRNA, 28 S rRNA) using a taxon set accumulated over a period of 15 years, coupled with species delimitation analyses methods (ABGD, ASAP, bPTP) and morpho-anatomical studies. The goal of this study is to provide insights into the taxonomy, phylogenetic relationships and geographical distributions of species while generating a framework for future systematic reviews of the genus, as well as to study speciation and historical biogeography. Our results rendered four possible hypotheses of species diversity: with $14,15,19$ and 20 candidate species and point to the fact that several taxa presently regarded as valid might be conspecific (e.g. H. orteai-H. templadoi-H. exigua; and H. alfredensis-H. antillarum-H. orbignyana), while highlighting the existence of a complex of four or five species often identified as $H$. elegans. Pervasive nomenclatural problems in the genus, including with the type species $H$. hydatis, are highlighted and discussed.


## KEYWORDS

Atlantic, biodiversity, Eastern Pacific, Haminoeidae, Mollusca, systematics

[^0]
## 1 | INTRODUCTION

The marine gastropods of the genus Haminoea Turton \& Kingston in Carrington, 1830 have been long considered to have a worldwide distribution, inhabiting temperate and tropical shorelines (Burn \& Thompson, 1998; Malaquias \& Cervera, 2006; Rudman, 1971). Yet, Oskars et al. (2019) and Oskars and Malaquias (2019) found Haminoea to be a paraphyletic assemblage of five distinct evolutionary lineages; three mostly tropical and restricted to the Indo-West Pacific (Haloa Pilsbry, 1921, Lamprohaminoea Habe, 1952 and Bakawan Oskars \& Malaquias, 2020a), one confined to Australasian (Papawera Oskars \& Malaquias, 2020b) and Haminoea proper, geographically restricted to the Atlantic Ocean (including the Mediterranean Sea), the Eastern Pacific Ocean and with a single lineage represented in temperate stretches of the Indian Ocean coastline of South Africa.

The small, globose, semi-translucent and thin shells of these five genera are similar in shape, colour and size, which coupled with a lack of broad comparative morphological studies and phylogenetic frameworks, led the majority of authors to accept Haminoea as the only valid genus across the world (e.g. Burn \& Thompson, 1998; Cervera et al., 2004; Gosliner et al., 2008; Malaquias \& Cervera, 2006; Thompson, 1981; Valdés et al., 2006). The modern concept of Haminoea was proposed by Oskars et al. (2019) and Oskars and Malaquias (2019) who identified several diagnostic features to separate it from their Indo-West Pacific closely related genera, namely the higher number of lateral radular teeth, the presence of a muscular penis and a Hancock organ with a perfoliate structure.

Species of Haminoea are herbivorous and live predominantly in estuaries and coastal lagoons, where they are often found on seagrass, algae or sandy-muddy bottoms, but can also occur on rocky shores in tidepools or shallow depths always among algal mats (Boulch-Bleas, 1983; Malaquias et al., 2002, 2004, 2009; Rudman, 1971).

At present, eight species of Haminoea are recognized as valid in the Eastern Atlantic (EA) between the southern shores of the British Isles and Angola in West Africa, namely H. hydatis (Linnaeus, 1758; type locality Mediterranean Sea), H. navicula (Da Costa, 1778; type locality Weymouth, Dorset, England), H. orbignyana (Férussac, 1822; type locality near La Rochelle, Bay of Biscay, France), H. elegans (Gray, 1825; type locality south of British Isles and Mediterranean Sea), H. orteai Talavera, Murillo \& Templado, 1987 (type locality Salinas del Rasall, Murcia, Spain), H. templadoi García, Pérez-Hurtado \& García-Gómez, 1991 (type locality Huelva, Spain), H. exigua (Schaefer, 1992; type locality Adriatic Sea, Italy/ Croatia) and H. fusari (Álvarez, García \& Villani, 1993; type
locality Lake Fusaro, Italy) (Malaquias \& Cervera, 2006; Martínez \& Ortea, 1997; Rolán \& Ryall, 1999). In the Western Atlantic (WA) four species are often accepted as valid in current literature occurring between Florida, USA and Rio Grande do Sul in Brazil, namely H. elegans, H. antillarum (d'Orbigny, 1841; type locality Saint Thomas, U.S. Virgin Islands), H. petitii (d'Orbigny, 1841; type locality Cuba) and H. succinea (Conrad, 1846; type locality Tampa Bay, Florida, USA) (Caballer et al., 2015; García et al., 2008; Rios, 2009; Valdés et al., 2006). In the Eastern Pacific (EP) three species are commonly recognized as valid between Alaska and Panama, namely $H$. ovalis Pease, 1868 (type locality Tahiti, French Polynesia), H. virescens (Sowerby, 1833; type locality Pitcarin Island or California, USA; see Valdés, 2019) and H. vesicula (Gould, 1855; type locality San Diego, California, USA) (Behrens \& Hermosillo, 2005; Hermosillo et al., 2006; Valdés \& Camacho-Garcia, 2004). In addition, one species of Haminoea occurs in temperate stretches of the Indian Ocean coastline of South Africa (tWIO), namely H. alfredensis Bartsch, 1915 (type locality Port Alfred, South Africa), distributed on both sides of the Cape Peninsula eastwards up to East London (Gosliner, 1987).

In total, 15 species of Haminoea are currently accepted as valid in current scientific literature and field guides. However, in a literature search we were able to identify 48 nominal species, most of them of uncertain taxonomic status because of short and ambiguous species descriptions based only on shells, which are similar in shape, colour and dimensions (e.g. Leach, 1852 for H. dilatata; A. Adams, 1850 for H. glabra; Baker \& Hanna, 1927 for H. angelensis; Petuch, 1987 for $H$. taylorae).

Furthermore, even among the 'well-established' species, there are questions about the taxonomic status of several of them. For example, the definition of the type species of the genus-H. hydatis-is problematic. This species was described by Linnaeus (1758) based on shells (unclear if only one or several) from the Mediterranean Sea but later assumed by various authors to be conspecific with specimens occurring between the British Isles and the Adriatic Sea, and characterized by having a smooth shell, a bilobed prostate separated by a constricted region and a radula with the first lateral tooth denticulated (Pruvot-Fol, 1954; Tchang, 1931; Thompson, 1981; Thompson \& Brown, 1976; Vayssière, 1885). Another case is the species name $H$. elegans introduced by Gray (1825) based on shells from the British Isles and the Mediterranean Sea, yet, the name is commonly attributed to one of the tropical western Atlantic species (e.g. Caballer et al., 2015; Malaquias, 2014; Marcus, 1976; Marcus \& Marcus, 1967; Redfern, 2001; Valdés et al., 2006) and also to spiralled shells occurring in tropical West Africa (Gabon, Republic of the Congo, São Tomé and

Príncipe, Angola; Bernard, 1984; Martínez \& Ortea, 1997; Rolán \& Ryall, 1999). Likewise, the name H. ovalis is commonly employed to designate animals with tiny orange or yellow dots on the body occurring in the Eastern Pacific, between Mexico and Peru (Behrens \& Hermosillo, 2005; Hermosillo et al., 2006; Oskars \& Malaquias, 2019; Valdés \& Camacho-Garcia, 2004); nonetheless, H. ovalis was described by Pease (1868) from Tahiti and was recently reassigned to the genus Lamprohaminoea by Oskars and Malaquias (2020c), who confirmed the species to be widespread in the Indo-West Pacific and absent from the Eastern Pacific. In fact, in a previous study, Oskars \& Malaquias (2019, as Haminoea sp.1475) showed that dotted orange haminoeids from Peru were phylogenetically related to all other Atlantic and Eastern Pacific Haminoea species.

In the present study, we generate the first complete phylogeny of the genus Haminoea based on multilocus molecular characters using a taxon set accumulated over a period of 15 years, which we believe to likely cover the entire diversity of the genus and include a comprehensive geographical coverage of the distribution of species. The main goals of this paper are to define the number of species in Haminoea and provide insights on their taxonomy, phylogenetic relationships and geographical distributions while establishing a framework for future detailed systematic reviews and studies on speciation and historical biogeography of this genus.

## 2 | MATERIALS AND METHODS

## 2.1 | Sampling of taxa

Specimens of Haminoea were obtained during fieldwork in Bermuda (2009), Venezuela (2010), Brazil (2012), Bahamas (2013), Portugal (2014) and Florida Keys, USA (2015), from donations from colleagues, and loans of museum collections; e. g., University Museum of Bergen, Norway (ZMBN), The Natural History Museum, London, UK (NHMUK), The Natural History Museum of Florida, USA (UF), Museu Municipal do Funchal (História Natural) (MMF(HN)), California Academy of Sciences (CAS), Museo de Ciencias Naturales de Madrid (MCNM), Bavarian State Collections of Zoology (ZSM) and Cal Poly Pomona Invertebrate Collection, USA (CPIC).

For the majority of the 15 recognized valid species (see Introduction) our dataset includes specimens from the type localities or nearby places $(50-100 \mathrm{~km})$. The exceptions are $H$. orbignyana, $H$. orteai, $H$. antillarum and $H$. succinea, but in these last four cases, specimens were still assembled from the same biogeographical areas of the type localities (see Introduction and Table 1).

Outgroup taxa consisted of species from two additional genera, namely Haloa (represented by four species) and Lamprohaminoea (represented by one species). The trees were rooted with Smaragdinella, a genus closely related to Haminoea (Oskars et al., 2019). In total, this study includes 206 specimens (94 EA Haminoea, 69 WA Haminoea, 18 EP Haminoea, 5 tWIO Haminoea and 16 outgroup taxa) and a total of 608 sequences, of which 426 were newly generated for this study (Table 1).

### 2.2 DNA extraction, amplification and sequencing

DNA was extracted from tissue obtained from the foot or parapodial lobes using the Qiagen DNeasy Blood and Tissue Kit (catalogue no. 69504) following the protocol recommended by the manufacturer. For small specimens with shell height between 2 and 3 mm , the whole specimen was digested and hard parts such as the shell, radula and gizzard plates were collected for morphological examination.

Partial sequences of the mitochondrial genes cytochrome c oxidase subunit I (COI; primers: LCO1490 (F) GGTCA ACAAATCATAAAGATATTGG and HCO2198 (R) TAAAC TTCAGGGTGACCAAAAATCA by Folmer et al., 1994; C_ GasF1_t1 (F) TGTAAAACGACGGCCAGTTTTCAACAAA CCATAARGATATTGG and GasR1_t1 (R) CAGGAAACAG CTATGACACTTCWGGRTGHCCRAARAATCARAA by Steinke et al., 2016), 16S rRNA (16S; primers: 16Sar-L (F) CGCCTGTTTATCAAAAACAT and $16 \operatorname{Sbr}-\mathrm{H}$ (R) CCGGT CTGAACTCAGATCACGT by Palumbi et al., 1991), and 12S rRNA (12S; primers: 12SA-L (F) AAACTGGGATTAGA TACCCCACTAT and 12SB-H (R) GAGGGTGACGGGCG GTGTGT by Palumbi, 1996), as well as the nuclear gene 28S rRNA (28S; LSU5-F TAGGTCGACCCGCTGAAYTTAAGCA by Littlewood et al., 2000; 900-F CCGTCTTGAAACACGGA CCAAG by Olson et al., 2003; LSU1600-R AGCGCCATCCA TTTTCAGG by Williams et al., 2003; ECD2S-R CTTGGTCC GTGTTTCAAGACGG modified from Littlewood et al., 2000 by Williams et al., 2003) were amplified and sequenced. Polymerase chain reactions (PCR) were performed in $25 \mu \mathrm{~L}$ volume and for the COI and 28S genes followed the protocols described by Malaquias et al. (2009), whereas for the 16 S the protocol described by Oskars et al. (2015) was used, and for the 12 S gene we applied the protocol described by Oskars and Malaquias (2019). Annealing temperatures were $45^{\circ} \mathrm{C}$ for the COI gene, $51.5^{\circ} \mathrm{C}$ for $16 \mathrm{~S}, 49.4^{\circ} \mathrm{C}$ for 12 S and $52^{\circ} \mathrm{C}$ for the 28S gene.

For samples that did not amplify with Qiagen Taq, additional $25 \mu \mathrm{~L}$ reactions were set with TaKaRa Ex Taq Polymerase HS (250 U) (Cat. number: RR006A), following the protocol described by Oskars et al. (2015). For some
TABLE 1 List of specimens of Haminoea and outgroups used for phylogenetic analyses, with sampling localities, voucher numbers and GenBank accession numbers (numbers with prefix OR' are novel sequences generated for this study).

| Taxon | DNA extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. navicula | C51 | 1 | Hampshire, South England, UK | BMNH 20060324 | DQ974676 | OR343604 | OR351715 | DQ927233 |
| H. navicula | 129 | 1 | Aveiro, Portugal | BMNH 20070018 | KF615838 | MK474364 | MK474221 | KF615804 |
| H. navicula | 130 | 1 | Aveiro, Portugal | BMNH 20070020 | KF615837 | OR343606 | OR351711 | KF615803 |
| H. navicula | 131 | 1 | Aveiro, Portugal | BMNH 20070020 | KF615839 | MK474365 | MH933144 | KF615805 |
| H. navicula | 147 | 1 | Hampshire, South England, UK | BMNH 20070021 | KF615836 | MK474366 | MH933145 | KF615806 |
| H. navicula |  | 1 | Aveiro, Portugal | BMNH 20070018.2 | EU314804 | - | - | - |
| H. navicula | 398 | 1 | France, Mediterranean | ZMBN 81646 | OR352608 | OR343608 | OR351713 | OR346082 |
| H. navicula | 294 | 1 | France, Mediterranean | ZMBN 81647.1 | OR352609 | OR343607 | OR351712 | OR346083 |
| H. navicula | 295 | 1 | France, Mediterranean | ZMBN 81647.2 | OR352610 | OR343605 | OR351714 | OR346084 |
| Haminoea sp. | 526 | 2 | Žirje I., Croatia | ZMBN 121357 | OR352611 | - | - | - |
| Haminoea sp. | 543 | 3 | Roses, Catalonia, Spain, Mediterranean | ZMBN 112904 | OR352612 | OR343603 | OR351650 | OR346094 |
| H. 'fusari' | 535 | 4 | Taranto, Italy | ZMBN 106885 | OR352613 | OR343609 | OR351726 | - |
| H. 'fusari' | 167 | 4 | Naples, Italy | BMNH 20070177 | KF615840 | MK474368 | MH933152 | KF615801 |
| H. 'fusari' | 511 | 4 | Rovinj, Croatia | ZMBN 119729.1 | OR352614 | OR343610 | OR351729 | OR346085 |
| H. 'fusari' | 512 | 4 | Rovinj, Croatia | ZMBN 119729.2 | OR352621 | OR343611 | OR351728 | OR346087 |
| H. 'fusari' | 522 | 4 | Bene, Croatia | ZMBN 119735 | OR352618 | OR343612 | OR351730 | OR346086 |
| H. 'fusari' | 515 | 4 | Žut I., Croatia | ZMBN 119731.1 | OR352619 | OR343613 | OR351716 | OR346097 |
| H. 'fusari' | 516 | 4 | Žut I., Croatia | ZMBN 119731.2 | OR352620 | OR343614 | OR351727 | OR346098 |
| H. 'fusari' | 525 | 4 | Žirje I., Croacia | ZMBN 121357 | OR352615 | - | - | - |
| H. 'fusari' | 527 | 4 | Žirje I., Croacia | ZMBN 121358.1 | OR352616 | - | - | - |
| H. 'fusari' | 528 | 4 | Žirje I., Croacia | ZMBN 121358.2 | OR352617 | - | - | - |
| Haminoea sp. | C50 | 5 | Selvagem Grande I., Madeira Archipelago, Portugal | BMNH 20070024 | OR352622 | OR343615 | OR351717 | OR346081 |
| Haminoea sp. | 173 | 5 | Selvagem Grande I., Madeira Archipelago, Portugal | BMNH 20070025 | - | OR343616 | - | - |
| H. 'hydatis' | 524 | 6 | Žut I., Croatia | ZMBN 121356 | OR352623 | - | - | - |
| H. 'hydatis' | 530 | 6 | Žut I., Croatia | ZMBN 121355 | OR352625 | - | - | - |
| H. 'hydatis' | 513 | 6 | Iž I., Croacia | ZMBN 119730.1 | OR352626 | - | - | - |
| H. 'hydatis' | 514 | 6 | Iž I., Croacia | ZMBN 119730.2 | OR352624 | - | - | - |

TABLE 1 (Continued)

| Taxon | DNA extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. 'hydatis' | 529 | 6 | Iž I., Croacia | ZMBN 121355 | OR352627 | - | - | - |
| H. 'hydatis' | 540 | 6 | Roses, Catalonia, Spain, Mediterranean | ZMBN 112900 | OR352635 | OR343617 | OR351719 | OR346092 |
| H. 'hydatis' | 541 | 6 | Roses, Catalonia, Spain, Mediterranean | ZMBN 112901 | OR352632 | OR343624 | OR351725 | OR346093 |
| H. 'hydatis' | 542 | 6 | Port - Leucate, France, Mediterranean | ZMBN 112902 | OR352629 | OR343622 | OR351723 | OR346089 |
| H. 'hydatis' | 545 | 6 | Port - Leucate, France, Mediterranean | ZMBN 112906 | OR352630 | OR343623 | OR351722 | OR346090 |
| H. 'hydatis' | 546 | 6 | Cerbére, France, Mediterranean | ZMBN 112907 | OR352636 | OR343619 | OR351721 | OR346095 |
| H. 'hydatis' | 166 | 6 | Port Barcarès, France, Mediterranean | BMNH 20060326 | KF615841 | MK474367 | KJ022796 | KF615802 |
| H. 'hydatis' | 533 | 6 | Mataró, Barcelona, Spain | ZMBN 106861 | OR352634 | OR343618 | OR351710 | - |
| H. 'hydatis' | 561 | 6 | Cadaqués, Caials, Catalonia, Spain | ZMBN 131054 | OR352633 | OR343625 | OR351720 | OR346096 |
| H. 'hydatis' | C53 | 6 | France, Mediterranean | BMNH 20060326 | DQ974674 | MK474369 | MH933208 | DQ927231 |
| H. 'hydatis' | 559 | 6 | France Mediterranean | ZMBN 130226 | OR352631 | OR343621 | OR351724 | OR346091 |
| H. 'hydatis' | 259 | 6 | Lazaro, Calabria, Italy | ZMBN 81714 | OR352628 | OR343620 | OR351718 | OR346088 |
| H. 'elegans 4' | 311 | 7 | Guanacabibes, Cuba | MNCN Madrid | OR352637 | OR343591 | OR351697 | OR346044 |
| H. 'elegans 4' | 461 | 7 | Morrocoy, Venezuela | ZMBN 84942 | - | - | OR351693 | OR346045 |
| H. 'elegans 4' | 462 | 7 | Morrocoy, Venezuela | ZMBN 84943 | - | OR343592 | OR351696 | OR346046 |
| H. 'elegans 4' | 463 | 7 | Morrocoy, Venezuela | ZMBN 84903.1 | - | OR343593 | OR351695 | - |
| H. 'elegans 4' | 464 | 7 | Morrocoy, Venezuela | ZMBN 84903.2 | - | OR343590 | OR351694 | - |
| H. 'elegans 4' | 465 | 7 | Morrocoy, Venezuela | ZMBN 84939.1 | - | - | OR351692 | OR346047 |
| H. 'elegans 4' | 466 | 7 | Morrocoy, Venezuela | ZMBN 84939.2 | - | OR343594 | OR351698 | - |
| H. vesicula | 202 | 8 | Bodega Harbor, Sonoma Co., California, USA | CAS97502 | KF615843 | MK474362 | MH933161 | KF615789 |
| H. vesicula | 574 | 8 | Cat Harbor, Catalina I., California, USA | CPIC 01237 | OR352641 | OR343601 | OR351690 | OR346058 |
| H. vesicula | 575 | 8 | Long Beach, California | CPIC 00606 | OR352639 | - | OR351691 | OR346057 |
| H. vesicula | 577 | 8 | San Pedro, California | CPIC 01030 | OR352640 | OR343599 | OR351689 | - |
| H. vesicula | 571 | 8 | San Diego, California | CPIC 00222 | OR352638 | OR343602 | OR351688 | - |
| H. vesicula |  | 8 | British Columbia, Indian Arm, Canada | BIOUG12670-G07 | MG423188 | - | - | - |

TABLE 1 (Continued)

| Taxon | DNA extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. vesicula |  | 8 | British Columbia, Indian Arm, Canada | 10BCMOL-00307 | KF643501 | - | - | - |
| H. vesicula |  | 8 | Washington, USA |  | JQ693571 | - | - | - |
| H. vesicula |  | 8 | False Bay, San Juan Island, Washington, USA | BMBM-0081 | MH242779 | - | - | - |
| H. vesicula |  | 8 | Haida Gwaii, British Columbia, Canada | 10BCMOL-00152 | KF643269 | - | - | - |
| H. vesicula |  | 8 | Haida Gwaii, British Columbia, Canada | 10BCMOL-00154 | KF643444 | - | - | - |
| H. vesicula |  | 8 | Haida Gwaii, British Columbia, Canada | 10BCMOL-00348 | KF643861 | - | - | - |
| H. vesicula |  | 8 | Haida Gwaii, British Columbia, Canada | 10BCMOL-00345 | KF643877 | - | - | - |
| H. vesicula |  | 8 | Haida Gwaii, British Columbia, Canada | 10BCMOL-00346 | KF643968 | - | - | - |
| H. vesicula |  | 8 | Haida Gwaii, British Columbia, Canada | 10BCMOL-00347 | KF644011 | - | - | - |
| H. vesicula | 481 | 8 | Long Beach, California | ZMBN 88214 | - | - | - | MK474262 |
| H. orteai |  | 9 | Greece | CPIC 01879 | KX683878 | - | - | - |
| H. orteai |  | 9 | Greece | CPIC 01880 | KX683879 | - | - | - |
| H. orteai |  | 9 | Greece | CPIC 01878 | KX683877 | - | - | - |
| H. orteai | 197 | 9 | Boca das Caldeirinhas, Faial I., Azores, Portugal | BMNH 20070458.1 | KF615844 | MK474363 | MH933160 | KF615791 |
| H. orteai | 198 | 9 | Boca das Caldeirinhas, Faial I., Azores, Portugal | BMNH 20070458.2 | KF615845 | KF615790 | MK474238 | KF615790 |
| H. orteai | 508 | 9 | Rovinj, Croatia | ZMBN 119726.1 | OR352652 | - | - | - |
| H. orteai | 510 | 9 | Rovinj, Croatia | ZMBN 119726.2 | OR352653 | - | - | - |
| H. orteai | 517 | 9 | Zadar, Croatia | ZMBN 119728.1 | OR352654 | - | - | - |
| H. orteai | 518 | 9 | Zadar, Croacia | ZMBN 119728.2 | OR352655 | - | - | - |
| H. orteai | 519 | 9 | Zadar, Croatia | ZMBN 119728.3 | OR352656 | - | - | - |
| H. orteai | 520 | 9 | Brac I., Maslinova Bay, Croatia | ZMBN 119734.1 | OR352648 | - | - | - |
| H. orteai | 521 | 9 | Brac I., Maslinova Bay, Croatia | ZMBN 119734.2 | OR352658 | - | - | - |
| H. orteai | 523 | 9 | Lubinski, Croatia | ZMBN 119736 | OR352647 | - | - | - |

TABLE 1 (Continued)

| Taxon | DNA extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. orteai | 507 | 9 | Rovinj, Croatia | ZMBN 119726 | OR352643 | - | - | - |
| H. orteai | 281 | 9 | Tenerife, Canary Is, Spain | BMNH 20030802 | OR352659 | - | - | - |
| H. orteai | 258 | 9 | Naples, Italy | ZMBN 81761 | OR352657 | - | OR351699 | OR346048 |
| H. orteai | 282 | 9 | Tenerife, Canary Is, Spain | BMNH 20030802 | OR352642 | OR343596 | OR351706 | - |
| H. orteai | 369 | 9 | Naples, Italy | ZMBN 83030 | OR352644 | - | OR351702 | OR346054 |
| H. orteai | 292 | 9 | Zoomarine, Algarve, Portugal | ZMBN 81696 | OR352650 | - | OR351701 | OR346050 |
| H. orteai | 293 | 9 | Zoomarine, Algarve, Portugal | ZMBN 81696 | OR352645 | - | OR351709 | OR346051 |
| H. orteai | C54 | 9 | Cape Ferret, France, Atlantic | BMNH 20070030 | OR352651 | - | - | - |
| H. orteai | 370 | 9 | Naples, Italy | ZMBN 83030 | OR352646 | - | OR351703 | OR346053 |
| H. orteai | 254 | 9 | Naples, Italy | ZMBN 81701.2 | KX383914 | - | OR351708 | OR346056 |
| H. orteai | 255 | 9 | Naples, Italy | ZMBN 81701.3 | KX383913 | - | OR351704 | OR346049 |
| H. orteai | 48 | 9 | Tenerife, Canary Is, Spain | BMNH 20030836 | KF615846 | - | MK474239 | KF615792 |
| H. orteai | 151 | 9 | Sal I., Cape Verde Archipelago | BMNH 20070023 | KX383915 | - | OR351707 | OR346052 |
| H. orteai | 283 | 9 | Porto Velho do Varadouro, Faial I., Azores, Portugal | BMNH 20070459 | KC404963 | OR343598 | KC404960 | KC404962 |
| H. orteai | 536 | 9 | Cap Ferret, France, Atlantic | ZMBN 112891 | OR352649 | OR343597 | OR351700 | OR346055 |
| H. orteai | 253 | 9 | Naples, Italy | ZMBN 81701.1 | KX383912 | - | MH933172 | MH933367 |
| H. orteai | 144 | 9 | Porto Moniz, Madeira I., Portugal | MMF(HN) 36,229 | - | - | OR351705 | - |
| H. 'ovalis' | 578 | 10 | Baya de Banderas, Mexico | CPIC 00177 | OR352660 | - | OR351605 | - |
| H. 'ovalis' | 475 | 10 | Máncora, Piura, Peru | $\begin{aligned} & \text { ZSM } \\ & \quad \text { Mol-20,100,737 } \end{aligned}$ | - | OR343519 | OR351606 | - |
| H. virescens | 489 | 11 | Long Beach, California USA | ZMBN 88213 | - | - | OR351609 | MK474255 |
| H. virescens | MT 572 | 11 | Palos Verdes, California, USA | CPIC 01055 | OR352662 | OR343540 | OR351607 | OR346018 |
| H. virescens | MT 573 | 11 | Long Beach, California, USA | CPIC 00186 | OR352661 | OR343539 | OR351610 | OR346017 |
| H. virescens | MT 580 | 11 | Catalina I., California, USA | CPIC 01238 | OR352663 | OR343541 | OR351608 | OR346016 |
| H. virescens | HVCal | 11 | Venice, California, USA |  | AF156142 | AF156110 | AF156126 | - |
| Haminoea sp. | 307 | 12 | Rio de Janeiro, Brazil | ZMBN 81796.1 | OR352665 | OR343536 | OR351601 | OR346019 |
| Haminoea sp. | 308 | 12 | Rio de Janeiro, Brazil | ZMBN 81796.2 | OR352666 | OR343538 | OR351604 | OR346020 |
| Haminoea sp. | 479 | 12 | São Sebastião, São Paulo State, Brazil | ZMBN 88219.1 | OR352664 | OR343537 | OR351602 | - |
| Haminoea sp. | 480 | 12 | São Sebastião, São Paulo State, Brazil | ZMBN 88219.2 | - | - | OR351603 | - |

TABLE 1 (Continued)

| Taxon | DNA <br> extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. alfredensis | 174 | 13 | Kariega river estuary, Kenton, South Africa | BMNH 20070314 | KF615816 | MK474355 | MH933154 | KF615774 |
| H. alfredensis | 182 | 13 | Kenton on Sea, Kariega river estuary, South Africa | BMNH 20070315.1 | KF615815 | MK474353 | OR351612 | KF615775 |
| H. alfredensis | 183 | 13 | Kenton on Sea, Kariega river estuary, South Africa | BMNH 20070315.2 | KF615814 | MK474354 | MH933155 | KF615773 |
| H. alfredensis | 459 | 13 | Knysna lagoon, South Africa | ZMBN 86406.1 | MK473513 | MK474343 | MK474188 | MK474257 |
| H. alfredensis | 460 | 13 | Knysna lagoon, South Africa | ZMBN 86406.2 | MK473514 | - | MK474184 | MK474256 |
| H. orbignyana | 278 | 14 | Niger Delta, Nigeria | ZMBN 81652.1 | OR352681 | OR343569 | OR351629 | OR346036 |
| H. orbignyana | 279 | 14 | Niger Delta, Nigeria | ZMBN 81652.2 | OR352682 | OR343571 | OR351624 | OR346031 |
| H. orbignyana | 531 | 14 | Praia Grande do Sul, Principe Is., São Tomé and Principe | ZMBN 106875 | OR352680 | OR343570 | OR351625 | OR346035 |
| H. orbignyana | 548 | 14 | Ayamonte, Huelva, Spain | ZMBN 130159 | OR352677 | OR343556 | OR351611 | OR346021 |
| H. orbignyana | 391 | 14 | Gran Canaria I., Canary Is., Spain | ZMBN 86408 | OR352669 | - | OR351640 | - |
| H. orbignyana | 396 | 14 | Rabat, Morocco | ZMBN 81793 | OR352671 | OR343555 | OR351619 | - |
| H. orbignyana | 397 | 14 | Aghroud, Morocco | ZMBN 81797 | - | OR343547 | - | - |
| H. orbignyana | 399 | 14 | Djerba, Tunisia | ZMBN 86407 | OR352678 | OR343559 | OR351614 | OR346038 |
| H. orbignyana | 275 | 14 | Rabat, Morocco | ZMBN 81791 | MH933103 | MK474352 | MH933174 | MH933369 |
| H. orbignyana | 277 | 14 | Agadir, Morocco | ZMBN 81799 | OR352672 | OR343552 | OR351623 | OR346037 |
| H. orbignyana | 296 | 14 | Aveiro, Portugal | ZMBN 81674.1 | OR352674 | OR343551 | OR351622 | OR346022 |
| H. orbignyana | 297 | 14 | Aveiro, Portugal | ZMBN 81674.2 | OR352668 | OR343545 | OR351639 | OR346039 |
| H. orbignyana | 256 | 14 | Naples, Italy | ZMBN 81714.1 | KC404964.1 | OR343548 | OR351616 | KC404961.1 |
| H. orbignyana | 257 | 14 | Naples, Italy | ZMBN 81714.2 | OR352670 | OR343550 | OR351616 | OR346033 |
| H. orbignyana | 498 | 14 | Lake Qarun, Egypt | ZMBN 99936.2 | KT339766 | - | - | - |
| H. orbignyana | 499 | 14 | Lake Qarun, Egypt | ZMBN 99936.1 | KT339765 | - | - | - |
| H. orbignyana | 549 | 14 | Isla Cristina, Huelva, Spain | ZMBN 130161 | OR352679 | OR343544 | OR351618 | OR346034 |
| H. orbignyana | 001 | 14 | Faro, Ria Formosa, Portugal | BMNH 20030296 | KF615813 | MK474360 | KJ022794 | KF615776 |
| H. orbignyana | 557 | 14 | Cartaya, Huelva, Spain | ZMBN 130169 | OR352667 | OR343557 | OR351641 | OR346027 |
| H. orbignyana | 558 | 14 | Huelva, Spain | ZMBN 130197 | OR352673 | OR343558 | OR351615 | OR346028 |
| H. orbignyana | 249 | 14 | Djerba, Golfo de Gabes, Tunisia | ZMBN 81710 | OR352675 | OR343546 | OR351620 | OR346040 |
| H. orbignyana | 250 | 14 | Djerba, Golfo de Gabes, Tunisia | ZMBN 81710 | OR352676 | OR343549 | OR351621 | OR346041 |
| H. orbignyana | 148 | 14 | Ria Formosa, Portugal | BMNH 20030296 | KF615812 | MK474359 | OR351613 | KF615777 |

TABLE 1 (Continued)

| Taxon | DNA extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. orbignyana | 367 | 14 | Agadir, Morocco | ZMBN 83029.1 | - | OR343553 | - | - |
| H. orbignyana | 368 | 14 | Agadir, Morocco | ZMBN 83029.2 | - | OR343554 | - | - |
| H. orbignyana | 390 | 14 | Gran Canaria I., Canary Is., Spain | ZMBN 86408 | - | - | OR351638 | - |
| H. 'antillarum 2' | 157 | 15 | Caburná, Yucatan, Mexico | BMNH 20070091 | KF615819 | OR343542 | OR351626 | KF615782 |
| H. 'antillarum 2' | 158 | 15 | Bocana, Sisal, Yucatan, Mexico | BMNH 20070094 | KF615811 | MK474356 | MK474186 | KF615779 |
| H. 'antillarum 2' | 159 | 15 | Caburná, Yucatan, Mexico | BMNH 20070092 | KF615818 | OR343543 | OR351627 | KF615780 |
| H. 'antillarum 2' | 160 | 15 | Bocana, Sisal, Yucatan, Mexico | BMNH 20070093 | KF615820 | MK474357 | MH933150 | KF615781 |
| H. 'antillarum 1' |  | 16 | Indian River Lagoon, Florida, USA | FTP_0224 | KP255198 | - | - | - |
| H. 'antillarum 1' | 176 | 16 | Jupiter inlet, Palm Beach, Florida, USA | BMNH 20070316 | KF615817 | MK474358 | MH933151 | KF615778 |
| H. 'antillarum 1' | 210 | 16 | Fort Pierce, Florida, USA | UF 369433 | OR352683 | OR343564 | OR351628 | OR346026 |
| H. 'antillarum 1' | 553 | 16 | Key Largo, Florida Keys, USA | ZMBN 99926 | OR352689 | - | - | - |
| H. 'antillarum 1' | 555 | 16 | Key Largo, Florida Keys, USA | ZMBN 99908 | OR352690 | OR343568 | OR351630 | - |
| H. 'antillarum 1' | 310 | 16 | Florida Keys USA | ZMBN 81766.1 | OR352685 | OR343560 | OR351632 | OR346024 |
| H. 'antillarum 1' | 271 | 16 | Long Key, Florida, USA | ZMBN 81769 | OR352687 | OR343565 | OR351636 | OR346032 |
| H. 'antillarum 1' | 309 | 16 | Florida Keys USA | ZMBN 81766.2 | OR352686 | OR343561 | OR351631 | - |
| H. 'antillarum 1' | 269 | 16 | Long Key, Florida, USA | ZMBN 81767 | OR352688 | OR343567 | OR351634 | OR346025 |
| H. 'antillarum 1' | 364 | 16 | Tobacco Bay, Bermuda | ZMBN 82989 | OR352684 | OR343562 | OR351637 | OR346029 |
| H. 'antillarum 1' | 270 | 16 | Long Key, Florida, USA | ZMBN 81768 | - | OR343566 | OR351635 | OR346030 |
| H. 'antillarum 1' | 303 | 16 | Florida Keys USA | ZMBN 81751 | - | OR343563 | OR351633 | OR346023 |
| H. 'elegans 3' | 47 | 17 | Bimini Is, Bahamas | BMNH 20060100 | - | OR343528 | OR351658 | OR346059 |
| H. 'elegans 3' | 265 | 17 | Abaco Is, Bahamas | ZMBN 81771 | - | OR343533 | OR351652 | OR346061 |
| H. 'elegans 3' | 266 | 17 | Abaco Is, Bahamas | ZMBN 81765 | OR352692 | OR343535 | OR351657 | OR346066 |
| H. 'elegans 3' | 267 | 17 | Abaco Is, Bahamas | ZMBN 81770 | - | OR343532 | OR351653 | OR346062 |
| H. 'elegans 3' | 268 | 17 | Abaco Is, Bahamas | ZMBN 81773 | - | OR343531 | OR351651 | - |
| H. 'elegans 3' | 304 | 17 | Abaco Is, Bahamas | ZMBN 81713 | - | OR343534 | OR351655 | OR346063 |
| H. 'elegans 3' | 305 | 17 | Abaco Is, Bahamas | ZMBN 81764 | - | OR343530 | OR351654 | OR346064 |
| H. 'elegans 3' | 306 | 17 | Abaco Is, Bahamas | ZMBN 81763 | - | - | OR351656 | OR346065 |
| H. 'elegans 3' | 568 | 17 | Great Oyster Pond, Eleuthera I., Bahamas | ZMBN 91089 | OR352691 | OR343529 | OR351659 | OR346060 |
| H. 'elegans 1' | 168 | 18 | Sal-Rei, Boavista I., Cape Verde | BMNH 20020716 | OR352693 | - | - | - |

TABLE 1 (Continued)

| Taxon | DNA <br> extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. 'elegans 1' | 175 | 18 | Kiwanis Park, Florida, USA | BMNH 20070318 | KF615827 | OR343573 | OR351663 | KF615800 |
| H. 'elegans 1' | 188 | 18 | Banana River, Florida, USA | BMNH 20070448.1 | KF615828 | MK474349 | MK474213 | KF615798 |
| H. 'elegans 1' | 189 | 18 | Banana River, Florida, USA | BMNH 20070448.2 | KF615831 | OR343574 | OR351661 | KF615799 |
| H. 'elegans 1' | 190 | 18 | Pine Channel, Florida, USA | BMNH 20070603.1 | KF615832 | MK474350 | MK474214 | KF615795 |
| H. 'elegans 1' | 152 | 18 | Banana River, Florida, USA | BMNH 20070180 | KF615829 | MK474348 | MK474212 | KF615797 |
| H. 'elegans 1' | 552 | 18 | Summerland Key, Florida, USA | ZMBN 99935 | OR352695 | - | OR351665 | OR346067 |
| H. 'elegans 1' | 554 | 18 | Key Largo, Florida, USA | ZMBN 99925 | OR352694 | OR343572 | OR351666 | OR346068 |
| H. 'elegans 1' | 191 | 18 | Pine Channel, Florida, USA | BMNH 20070603.2 | KF615830 | OR343575 | OR351660 | KF615796 |
| H. 'elegans 1' | 482 | 18 | Banana River Lagoon, Florida, USA | ZMBN 88212 | OR352696 | - | OR351664 | - |
| H. 'elegans 1' | 162 | 18 | Yucatan, Mexico | BMNH 20070089 | - | - | OR351662 | - |
| H. 'elegans 2' | 154 | 19 | Veracruz, Mexico | BMNH 20070175 | KF615834 | OR343584 | OR351682 | KF615794 |
| H. 'elegans 2' | 161 | 19 | Bocana, Sisal, Yucatan, Mexico | BMNH 20070090 | KF615833 | MK474347 | - | KF615793 |
| H. 'elegans 2' | 312 | 19 | Guanacabibes, Cuba | MNCN Madrid | OR352700 | OR343588 | OR351669 | OR346073 |
| H. 'elegans 2' | 313 | 19 | Guanacabibes, Cuba | MNCN Madrid | OR352701 | OR343582 | OR351667 | OR346074 |
| H. 'elegans 2' | 314 | 19 | Guanacabibes, Cuba | MNCN Madrid | OR352702 | OR343583 | OR351668 | OR346075 |
| H. 'elegans 2' | 550 | 19 | Great Oyster Pond, Eleuthera I. Bahamas | ZMBN 91086 | OR352698 | OR343576 | OR351672 | OR346069 |
| H. 'elegans 2' | 566 | 19 | El Ocho's lagoon, Morrocoy National Park, Venezuela | ZMBN 84926 | OR352710 | OR343589 | OR351670 | OR346071 |
| H. 'elegans 2' | 569 | 19 | Sweeting Pond, Eleuthera I., Bahamas | ZMBN 91093 | OR352699 | OR343577 | OR351683 | - |
| H. 'elegans 2' | 570 | 19 | Turtle Pond, Eleuthera I., Bahamas | ZMBN 91103 | OR352711 | - | OR351673 | - |
| H. 'elegans 2' | 298 | 19 | Abaco Is, Bahamas | ZMBN 81759 | OR352712 | OR343587 | OR351674 | OR346070 |
| H. 'elegans 2' | 365 | 19 | Tom Moore's pond, Bermuda | ZMBN 82999.1 | OR352707 | OR343585 | OR351684 | OR346079 |
| H. 'elegans 2' | 366 | 19 | Tom Moore's pond, Bermuda | ZMBN 82999.2 | OR352706 | OR343586 | OR351685 | OR346080 |
| H. 'elegans 2' | 371 | 19 | Tom Moore's pond, Bermuda | ZMBN 83024.1 | OR352705 | OR343578 | OR351675 | OR346077 |
| H. 'elegans 2' | 372 | 19 | Tom Moore's pond, Bermuda | ZMBN 83024.2 | OR352709 | OR343579 | OR351676 | OR346076 |
| H. 'elegans 2' | 373 | 19 | Tom Moore's pond, Bermuda | ZMBN 82983.1 | OR352703 | OR343580 | OR351671 | OR346078 |
| H. 'elegans 2' | 374 | 19 | Tom Moore's pond, Bermuda | ZMBN 82983.2 | OR352704 | OR343581 | OR351677 | OR346072 |
| H. 'elegans 2' | 484 | 19 | São Sebastião, São Paulo, Brazil | ZMBN 88217.1 | OR352697 | OR343595 | OR351680 | - |
| H. 'elegans 2' | 485 | 19 | São Sebastião, São Paulo, Brazil | ZMBN 88217.2 | - | - | OR351681 | - |
| H. 'elegans 2' | 486 | 19 | Veracruz, Mexico | ZMBN 88204 | OR352713 | - | OR351679 | - |

TABLE 1 (Continued)

| Taxon | DNA extraction code | Group No | Locality | Voucher No | COI | 12S rRNA | 16S rRNA | 28S rRNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| H. 'elegans 2' | 492 | 19 | Ilha de Santo Amaro, Guarujá, São Paulo, Brazil | ZMBN 88218.1 | - | - | OR351687 | - |
| H. 'elegans 2' | 493 | 19 | Ilha de Santo Amaro, Guarujá, São Paulo, Brazil | ZMBN 88218.2 | OR352708 | - | OR351686 | - |
| H. 'elegans 2' | 477 | 19 | Quintana Roo, Mexico | ZMBN 88203 | - | - | OR351678 | - |
| Haloa japonica | c52 |  | Barcarès, France, Mediterranean | BMNH 20070029 | KF615824 | OR343525 | OR351645 | KF615786 |
| Haloa japonica | 149 |  | Barcarès, France, Mediterranean | BMNH 20070065 | KF615823 | OR343523 | OR351646 | KF615787 |
| Haloa japonica | 164 |  | Barcarès, France, Mediterranean | BMNH 20070028 | KF615822 | OR343526 | OR351647 | KF615785 |
| Haloa japonica | 251 |  | Bahia de Ogrove, Galicia, Spain | ZMBN 81700.1 | OR352606 | OR343524 | OR351648 | OR346042 |
| Haloa japonica | 252 |  | Bahia de Ogrove, Galicia, Spain | ZMBN 81700.2 | OR352607 | OR343527 | OR351649 | OR346043 |
| Haloa japonica |  |  | San Juan Islands, WA |  | JN830723 | - | - | - |
| Haloa japonica |  |  | Hokinawa, Japon | ZMBN 91233 | MK473509 | - | MK474248 | MK474264 |
| Haloa crocata |  |  | Beach House, Kauai, Hawaii | ZMBN 97228 | MK473470 | MK474393 | MK474222 | MK474278 |
| Haloa pemphis |  |  | Hamamatsu, Japan | NHMUK 20070446 | MH933106 | MK474379 | MH933156 | MH933343 |
| Haloa musetta |  |  | Heshikiya, Okinawa | ZMBN 112928 | MK473467 | MK474400 | MK474232 | MK474290 |
| Lamprohaminoea ovalis |  |  | Manza Beach, Okinawa | ZMBN 88230 | MK473488 | MK474428 | MK474194 | MK474303 |
| Lamprohaminoea ovalis |  |  | Airport Beach, Maui, Hawaii | NHMUK 20070031 | MK473489 | MK474427 | MK474173 | MK474250 |
| Lamprohaminoea ovalis | 262 |  | Calabria, Italy | ZMBN 81762.1 | - | OR343520 | OR351644 | OR346101 |
| Lamprohaminoea ovalis | 263 |  | Calabria, Italy | ZMBN 81762.2 | OR352604 | OR343521 | OR351642 | OR346099 |
| Lamprohaminoea ovalis | 276 |  | Malta | ZMBN 81803 | OR352605 | OR343522 | OR351643 | OR346100 |
| Smaragdinella sp. |  |  | Mozambique | ZMBN 125447 | MK473459 | MK474345 | MK474249 | MK474253 |

samples, the amount of $\mathrm{MgCl}_{2}$ and DNA was increased, and the volume of water adjusted accordingly in the PCR cocktail. In addition, 10x dilutions of DNA extractions were attempted for samples that did not yield results with all previous approaches.

The quality and quantity of PCR products were assessed by gel electrophoresis following standard methods (see Eilertsen \& Malaquias, 2013). Successful PCR products were purified according to the EXO-SAP method described by Eilertsen and Malaquias (2013). Sequence reactions were run on an ABI 3730XL DNA Analyser (Applied Biosystems).

## 2.3 | Phylogenetic analyses

Geneious (v. R11, Kearse et al., 2012) was used to inspect, edit, and assemble the chromatograms of the forward and reverse DNA strands. All sequences were blasted in GenBank to check for contamination. Single gene sequences were aligned with Muscle (Edgar, 2004) implemented in Geneious. Alignments were trimmed to a position where at least $50 \%$ of the sequences had nucleotides and missing positions at the ends were coded as missing data (?). All sequences were deposited in GenBank (Table 1).

Blocks of ambiguous data in the single gene alignments of the ribosomal genes were identified and excluded using Gblocks with stringent and relaxed settings (Talavera \& Castresana, 2007) (Appendix S1). The JModeltest software (Darriba et al., 2012) was used to find the best-fit model of evolution for each single gene dataset under the Akaike information criterion (Akaike, 1974). Seven individual gene analyses were initially performed: COI (Appendix S2; 681 bp ; GTR + G), 12S Gblocks-relaxed (AppendixS3;355bp;TVM + G), 12 S Gblocks-stringent (Appendix S4; $267 \mathrm{bp} ; \operatorname{TrN}+\mathrm{I}+\mathrm{G}$ ), 16S Gblocks-relaxed (Appendix S5; $420 \mathrm{bp} ;$ GTR + G + I), 16S Gblocks-stringent (Appendix S6; 393bp; TVM+I+G), 28S Gblocks-relaxed (Appendix S7; $1047 \mathrm{bp} ; \operatorname{TrN}+\mathrm{I}+\mathrm{G}$ ) and 28S Gblocks-stringent (Appendix S8; 1036 bp; GTR + I + G).

Bayesian inference analyses (BI) using MrBayes (Huelsenbeck \& Ronquist, 2001) were run through the portal CIPRES Science gateway V.3.3 (https://www.phylo.org) on the initial single gene datasets (Appendix S2-S8) and all-genes concatenated dataset (Figure 1, Appendices S9 and S10; 2492 bp ). For the ribosomal genes, the datasets selected for concatenation were those that yielded the best-resolved trees with higher node support. All samples with sequences available for two or more genes were used in the concatenation analysis. In addition, samples with a single gene from unique geographical localities or with a unique phylogenetic position in the single gene trees were also included in the concatenated dataset. The analyses
used three parallel runs of 5 million generations for the single gene analyses and 15 million generations for the concatenated dataset, with sampling every 100 generations. The concatenated dataset was partitioned by gene and each partition was run under the best-fit model of evolution. Convergence of runs was inspected in Tracer v1.7 (Rambaut et al., 2018) with a burn-in set to $25 \%$ by comparing the likelihood of trees drawn by the independent runs. Posterior probabilities (PP) higher than 0.95 were considered statistically significant (Alfaro et al., 2003; Huelsenbeck et al., 2001). A Maximum Likelihood analysis (ML) of the concatenated dataset was run with the RAxML (v.8.2; Stamatakis, 2014) plug-in implemented in Geneious. The analysis was partitioned by gene and run under the 'rapid bootstrapping and search for best scoring ML tree' algorithm, using a random starting tree and the model GTR $+\mathrm{G}+\mathrm{I}$ with 1000 bootstrap (BS) replicates. Bootstrap values higher than $75 \%$ were considered significantly supported (Felsenstein, 1985). Consensus phylograms were converted to graphics in FigTree v1.3.1 (Rambaut \& Drummond, 2009).

COI uncorrected $p$-distances were calculated in MEGA (ver. 7, Kumar et al., 2016) (Table 2) within and between candidate species, by plotting pairwise uncorrected $p$ distances against total distances (transversions + transitions). This is a common approach to calculate genetic distances in Heterobranchia taxonomic research (e.g. Austin et al., 2018; Carmona et al., 2011; Jörger et al., 2012; Kienberger et al., 2016).

## 2.4 | Molecular species delimitation analyses

We used the DNA sequences of the COI gene to evaluate candidate species by using the Automatic Barcode Gap Discovery delimitation method (ABGD) (Puillandre et al., 2012) and the Assembling Species by Automatic Partitioning (ASAP) (Puillandre et al., 2021) under default settings and three different models of molecular evolution (Jukes-Cantor (JC69), Kimura TS/TV $=2.0$ (K80), Simple Distance). In addition, we used the bPTP method (Poison tree processes) on the same COI dataset. This method is intended to delimiting species that are consistent with the phylogenetic species concept and model speciation in terms of the number of substitutions (Zhang et al., 2013).

## 2.5 | Haplotype network analyses

Haplotype networks were generated based on the COI DNA sequences for the groups recognized by the phylogenetic analyses as putative candidate species, but which were

FIGURE 1 Cartoon based on the Bayesian phylogeny of Haminoea species depicted in Appendix S9 and resulting from the combined analysis of the mitochondrial COI, 12 S rRNA and 16 S rRNA and nuclear 28 S rRNA gene markers. Figures above branches are Bayesian posterior probabilities and those below branches are bootstrap values derived by maximum likelihood. The trees were rooted with Smaragdinella sp . and representatives of the genera Haloa and Lamprohaminoea were included as outgroups. Both rooting and outgroups were removed for clarity (see Appendices S9 and S10 for complete trees). Images groups $1,7,14,19$ by Manuel Malaquias; images groups 8,10 , 11 by Ángel Valdés; images groups 2, 4, 6, 9 courtesy of Jakov Prkic; image group 3 courtesy of Marina Poddubetskaia; image group 12 courtesy of Joana Bahia; image group 13 courtesy of George Branch; image group 15 courtesy of Jazmin Ortigosa; images groups 16,17 courtesy of Colin Redfern; image group 18 courtesy of Marlo Krisberg. Image not available for group 15.

clustered together by all or some of the ABGD and ASAP species delimitation analyses. This was the case for groups $4+5+6$; groups $13+14+15+16$; and groups $18+19$. The COI alignment derived for phylogenetic inference was edited to remove all sequences from non-target groups using the text editor programme Notepad++ v.8.3.3. Empty positions at both ends of the alignments were treated as missing data, yielding final alignments of 677 bp . The programme DnaSP v. 5 (Librado \& Rozas, 2009) was used to identify the number and sequences of the different haplotypes. Notepad++ was additionally used to generate trait files with geographic area codes based on a binary coding where 0 stands for sample absent and 1 for sample present and to edit the file into nexus format (nex). Alignments and trait files were finally run in PopArt v. 1.7 (Population Analysis with Reticulate Trees; Leigh \& Bryant, 2015) to create a standard tight compact spanning (TCS; Clement et al., 2002) network analysis to visualize the relationships and distances between the individual haplotypes from different groups and geographical areas. The TCS haplotype networks were edited in PopArt for more satisfying visualization.

## 2.6 | Morpho-anatomical analysis

The shell and anatomical features of selected specimens were studied to aid in interpreting the taxonomic status of some problematic lineages or to address complex taxonomic cases resulting from the molecular phylogenetic analyses (see Section 4 for details).

The shells were gently separated from the animals with the aid of forceps. The male reproductive system, gizzard and buccal bulb were dissected by a dorsal incision through the cephalic shield. Shells were photographed with a digital DSLR camera equipped with a macro lens and strobe lights. Shell height ( $H$ ) was measured with a digital Vernier calliper. The reproductive system was drawn using a stereo microscope fitted with a drawing tube and the penial sheath was removed to expose the penial papilla. The gizzard and buccal bulb were placed in a solution containing $180 \mu \mathrm{~L}$ buffer ATL with $20 \mu \mathrm{~L}$ of proteinase K solution (both from the Qiagen DNeasy ${ }^{\circledR}$ Blood and Tissue Kit) and incubated at $56^{\circ} \mathrm{C}$ at night in order to clean the gizzard plates, jaws and radulae. The penial papillae, gizzard plates, jaws and radulae were mounted on metallic stubs using carbon sticky tabs and then sputter-coated with gold-palladium for scanning electron microscopy (SEM). Prior to sputter coating and SEM, the gizzard plates and penial papillae were dehydrated with Hexamethyldisilazane (HMDS) by covering each sample inside small square watch glasses and left to dry
between 30 min and 1 h inside a fume hood. All samples were scanned and imaged with a Fei Quanta 450 scanning electron microscope.

## 3 | RESULTS

## 3.1 | Phylogenetic analyses

The ribosomal gene datasets selected for concatenation were the 12 S -relaxed, 16 S -relaxed and the 28 S -stringent. Though, it must be stressed that differences between relaxed and stringent datasets were minor in all cases (Appendix S5-S8). Thus, the all-genes concatenated dataset was based on the COI (178 sequences), 12S-relaxed (136 sequences), 16S-relaxed (162 sequences) and 28Sstringent (134 sequences).

The COI gene analyses rendered 19 groups putatively compatible with candidate species of Haminoea (Appendix S2). All groups but one (H. orbignyana; $\mathrm{PP}=0.88$ ), received maximum or nearly maximum support. The 12 S rendered 18 groups, but clade support was comparatively lower and often below statistical thresholds; the group missing is a singleton only represented in the COI dataset (Haminoea sp. 256 Croatia) (Appendix S2). The 16 S tree rendered 15 groups and clustered together with no support $(\mathrm{PP}=0.72)$ four groups recognized in the COI analyses (groups $13+14+15+16$ ). None of these four groups formed supported sub-clades. Only group 13 (H. alfredensis) was nearly supported ( $\mathrm{PP}=0.91$ ) but one representative branched apart (Appendix S5). The 28S gene tree was the less resolved with several of the groups recognized by the mitochondrial gene analyses, rendered non-monophyletic (Appendix S8). On the contrary, the concatenated analyses rendered the same 19 groups as the COI analysis, with two groups represented by singletons (Haminoea sp. 256 [Croatia; group 2] and Haminoea sp. 543 [Spanish Mediterranean; group 3]), one group (group 14) with moderate support $(\mathrm{PP}=0.95)$ and all remaining 16 groups with maximum support $(\mathrm{PP}=1)$ (Figure 1, Appendices S9 and S10).

### 3.2 Molecular species delimitation analyses and genetic distances

The ABGD analyses using the simple distance (SD) and Jukes-Cantor (JC69) models suggested the presence of 13 candidate species of Haminoea, clustering together groups $4+5+6$; groups $13+14+15+16$; and groups $18+19$. With the K2P model, the ABGD hypothesized the presence of 14 species by considering groups 15 and 16 distinct lineages (Appendix S11).

The ASAP results based on the three best ASAP-scores and for the same three evolutionary models used in ABGD rendered 11 candidate species of Haminoea (SD ASAPscore $=4$; JC69 ASAP-score $=3.5$; K2P ASAP-score $=3$; SD genetic threshold $=9.3 \%$ ), 14 species (SD ASAP-score $=5$; JC69 ASAP-score $=5$; K2P ASAP-score $=1.5$; SD genetic threshold $=5.9 \%$ ) and 17 species (SD ASAP-score $=4.5$; JC69 ASAP-score $=4.5$; K2P ASAP-score $=6$; SD genetic threshold $=3.1 \%$ ) (Appendix S12). The best ASAP score was 1.5 retrieved with the K2P model, suggesting 14 candidate species. This hypothesizes the same scenario as with the ABGD method under the same model, rendering 14 species and clustering groups $4+5+6$ and groups $13+14+15+16$, but separating groups 18 and 19 as distinct lineages.

The bPTP analysis suggested 20 candidate species, corresponding to the same 19 groups rendered by the COI and concatenated analyses, yet group 17 (H. 'elegans 3') represented by two samples was inferred to correspond to two putative species (sample 266 from Abaco, Bahamas and sample 568 from Eleuthera, Bahamas; Table 1, Appendices S2 and S13).

The estimated uncorrected $p$-distances between the 19 groups depicted in Figure 1 varied between a maximum of $19.8 \%$ (Haminoea 'elegans 4’; Western Atlantic [group 7] and Haminoea sp. Mediterranean Sea [group 2]) and a minimum of $2.4 \%$ (Haminoea alfredensis [group 13] and Haminoea orbignyana [group 14]). Genetic distances between several sister groups, namely those considered by the species delimitation methods to be conspecific were
comparatively low; for example, between groups 13,14 , $15,16(2.4 \%-4.7 \%)$, between groups $4,5,6(5.1 \%-7.4 \%)$ and between groups 18, 19 (7.5\%). All other sister groups have genetic distances equal to or higher than $10 \%$ (Table 2). The genetic distance between the two samples of $H$. 'elegans 3' recognized as distinct lineages by the bPTP analyses was $2.4 \%$. This corresponds to the intraspecific genetic distance for this species depicted in Table 2.

## 3.3 | Haplotype network analyses

The haplotype network of groups $4+5+6$ formed by samples from the eastern and central Mediterranean Sea was well structured with 14 haplotypes and three recognizable haplogroups separated by 28 substitutions(between groups 4 and 6) and 21 substitutions (between groups 5 and 6). Only one case of shared haplotypes was detected in group 6 between samples from Spain and France (Figure 2). The haplotype network of groups $13+14+15+16$ with samples from Europe, West Africa, temperate South Africa and Caribbean Sea, includes 19 haplotypes and four recognizable haplogroups connected through hypothetical haplotypes (black circles; Figure 3). The highest number of substitutions among haplogroups was 23 between group 13 (H. alfredensis) and 15 (H. 'antillarum 2'). The haplotype network of groups $18+19$ formed by samples from Brazil, Caribbean Sea and Cape Verde Islands includes 21 distinct haplotypes and two recognizable haplogroups separated by 36 substitutions (Figure 4).


FIGURE 2 COI haplotype network produced with the TCS method in PopART for groups 4, 5 and 6 . Colours of circles refer to the geographic origin of each haplotype. The relative size of circles is proportional to the number of sequences of that same haplotype. Black circles refer to hypothetical haplotypes and black bars to mutations.


Group 16-Haminoea "antillarum 1"



FIGURE 4 COI haplotype network produced with the TCS method in PopART for groups 18 and 19. Colours of circles refer to the geographic origin of each haplotype. The relative size of circles is proportional to the number of sequences of that same haplotype. Black circles refer to hypothetical haplotypes and black bars to mutations.
monophyletic, the genetic distances among them are comparatively lower varying between a minimum of 2.4\% (H. alfredensis and H.orbignyana) and a maximum of $4.5 \%$ ( $H$. antillarum group 16 and $H$. orbignyana), with the genetic distance between the two lineages of H. antillarum estimated at $3.9 \%$. These four lineages are only supported by the COI gene, whereas the 12 S and 28 S genes rendered support for the clade containing the specimens of $H$. antillarum from Yucatan (group 15), and the latter gene also provided support for the clade with specimens of $H$. alfredensis (Figure 1, Appendices S2, S3, S8, S10, S11 and S11). None of the four lineages was supported by the 16 S gene analysis, which nevertheless clustered all representatives together but with low node support ( $\mathrm{PP}=0.72$ ).

The species $H$. orbignyana, $H$. antillarum and $H$. alfredensis are well established in the literature, yet they were never studied in a comparative framework. A closer look at the literature together with our own preliminary data on the morphology of specimens reveals that all these nominal species share a pear-shaped smooth shell and a body colouration characterized by dense black pigmentation along the edges of the cephalic shield and parapodial lobes. In contrast, H. antillarum has mildly denticulated inner lateral radular teeth, whereas in H. alfredensis and H. orbignyana these teeth are smooth. Likewise, whereas in the latter two species, the proximal lobe of the prostate is wider, conferring the prostate an acorn-like shape, in $H$. antillarum seems to be the opposite with the distal lobe wider compared to the proximal one (Gosliner, 1987; Macnae, 1962; Malaquias \& Cervera, 2006; Marcus \& Marcus, 1967; Thompson, 1981; Valdés et al., 2006;
personal observations), but this requires further anatomical investigations in order to be confirmed.

Even if our molecular results based on the species delimitation analyses and genetic distances suggest the occurrence of a single ubiquitous species with amphiAtlantic distribution encompassing the Iberian Peninsula, the Mediterranean Sea, West Africa including the Canary Islands, the temperate shores of South Africa in the Indian Ocean and the western Atlantic along the Yucatan Peninsula, Florida and Bermuda, this warrants caution and further corroboration by conchological and morphoanatomical data. As highlighted above, H. antillarum seems to be characterized by relevant anatomical differences from the digestive and reproductive systems, and even if genetic distances are comparatively low, this could be due to different evolutionary rates between species of Haminoea.

On the contrary, and even in the absence of sound data on the duration of the pelagic larval stage of Haminoea (Schaefer, 1996), the confirmed occurrence of specimens attributed to H. elegans (group 18) on both sides of the Atlantic (Figure 1; only $0.3 \%$ different in the COI gene; Table 2) supports a high dispersal capability, at least in some species of the genus (Martínez \& Ortea, 1997; current study as $H$. 'elegans 1 ' [group 18]; Figure 1). Thus, we cannot discard that representatives of the orbignyana-alfredensis-antillarum complex may have larvae with high dispersal capability favouring gene flow between distant populations. However, we must admit that the genetic distance between the two putative lineages of $H$. antillarum from nearby locations, namely the Yucatan side of the Gulf of Mexico (group 15) and the Florida Keys/Florida

Peninsula-Bermuda (group 16), estimated at $3.9 \%$ and 14 substitutions between these two haplogroups (Figure 3) challenges this view. Even if the prevalent ocean current system in the area suggests putative connectivity between Yucatan and the Florida Peninsula through the Loop Current (Gyory et al., 2011), faunal breaks between tropical Florida and the more temperate/sub-tropical Gulf of Mexico have been documented for several groups of molluscs and fish (Briggs, 1974; Lee \& Ó Foighil, 2004; Mikkelsen \& Bieler, 2000; Reeb \& Avise, 1990), likely reflecting seasonal changes in the current systems and water temperatures oscillations. These factors may hinder gene flow between the Yucatan and the Florida-Bermuda populations, creating periods of temporary isolation that could explain the observed genetic discontinuity.

Another interesting aspect is the sister relationship between the lineages $H$. orbignyana (Eastern Atlantic) and $H$. alfredensis (temperate South Africa). The cold water of the Benguela current established at the end of the Miocene (Siesser, 1980) is regarded as a strong barrier for temperate and tropical marine coastal species isolating the faunas of the Atlantic and Indian Oceans, while at the same time, paleontological and morphological evidence suggests that this barrier was sporadically bridged by several coastal invertebrate organisms (Briggs, 1995; Vermeij \& Rosenberg, 1993). Few molecular evidence for dispersal from the Atlantic into the Indian Ocean is still available. This is the case for reef fish (Floeter et al., 2008; Rocha et al., 2005) and sea slugs (Churchill et al., 2014; Golestani et al., 2019), which seem to have taken advantage of the disruption of the Benguela and Agulhas currents during warmer interglacial periods of the Pleistocene. Because Haminoea is a genus of Atlantic and Eastern Pacific affinity, the sister relationship between $H$. orbignyana and H. alfredensis is more parsimoniously explained as being the result of dispersal of larvae or H. orbignyana into the Indian Ocean during these warmer periods, with the establishment of viable populations followed by isolation after the reestablishment of the current system.

## 4.2 | The Haminoea elegans complex

Haminoea elegans is characterized by having whitish translucent spiralled shells and it has been regarded as widely distributed in the Western Atlantic throughout the Gulf of Mexico and the Caribbean Sea southwards to Brazil (Valdés et al., 2006) with records in West Africa between the Gulf of Guinea and Angola (Bernard, 1984; Martínez \& Ortea, 1997; Rolán \& Ryall, 1999). However, the attribution of the name 'elegans' to this tropical amphi-Atlantic species stems certainly from a misidentification perpetuated in the literature over time. The name H. elegans was
introduced by Gray (1825) based on spiralled shells from the British Isles and the Mediterranean Sea, and the name is most certainly a junior synonym of Haminoea navicula, the only European species with a deeply spiralled shell (Malaquias \& Cervera, 2006).

Our results showed the existence of cryptic diversity in this 'species' with specimens provisionally ascribed by us to $H$. elegans splitting in four (or five) clades of possible species status (groups 7, 17-19; Figure 1, Appendices S2, S9, S10 and S13). Representatives of groups 17, 18 and 19 clustered together with maximum support, whereas group 7 branched off elsewhere in the tree (Figure 1, Appendix S2, S9 and S10).

If, in contrast, our results unequivocally support group 7 as a good species, they are not conclusive about the eventual status of group 17, with one of the species delimitation analysis (bPTP), suggesting the possible occurrence of two lineages in this group. However, none of the single gene and combined analyses retrieved reciprocal monophyly between sub-clades within group 17. When present, the sub-clades are not statistically supported (Appendices S3-S10).

The results are also not entirely conclusive about the conspecificity of groups 18 and 19 (see Section 3 - theme 3.2). Groups 18 and 19 are the only two in the complex with a genetic distance between themselves below $10 \%$, but still moderately high ( $=7.5 \%$ ). Moreover, in the haplotype network analysis, they were separated by 36 substitutions, the largest number of substitutions between putative conspecific groups among all our haplotype network analyses (Figure 4).

There are several names available in the literature that could be regarded as previous attempts to describe the shells variability in the H. elegans complex (e.g. H. guildingii (Swainson, 1840) [shells globose with visible spiral striae], H. petitii (d'Orbigny, 1841) [shells lacking or with inconspicuous spiral striae], H. succinea (Conrad, 1846) [shells cylindrical with tightly arranged spiral striae], $H$. taylorae (Petuch, 1987) [shells globose with numerous fine spiral striae]). These names have been in part considered synonyms of $H$. elegans (MolluscaBase, 2022; Valdés et al., 2006) or hardly used in scientific literature, but our results show the need to carefully re-evaluate the status of these names since some of them may apply to lineages revealed by our analyses.

The only study that provided a comparative analysis of the various types of shells of 'H. elegans' in the Western Atlantic was by Redfern (2013: 266-268). This author recognized five different types of whitish shells that could be associated with $H$. elegans; four with spiral striae and one apparently smooth. One of these forms was named by Redfern (2013) Haminoea elegans proper and the other four Haminoea A, B, C and D. According to the author,

Haminoea elegans and Haminoea sp. A are characterized by globose-quadrate opaque shells with wavy-spiral striae and a partially concealed involute spire; Haminoea sp. B by a more oval translucent shell, with numerous tightly arranged spiral striae and a spire concealed by a callus; Haminoea sp. C by a globose opaque smooth shell and a spire concealed by a callus; Haminoea sp. D by cylindrical translucent shells, with lightly impressed spiral striae and a spire concealed by a callus.

Here we provide for the first time a phylogenetic framework to properly explore the diversity of the Haminoea 'elegans' species-complex. Yet confirming whether our four (or five) candidate species correspond to the shell types identified by Redfern (2013) and are compatible with the names available in the literature, requires additional taxonomic work based on detailed analyses of conchological and morpho-anatomical characters.

## 4.3 | The Haminoea hydatisfusari complex

Another difficult case consists of groups 4, 5 and 6 in our phylogeny (Figure 1), which were rendered a single candidate species by the ABGD and ASAP molecular species delimitation methods. These three clades received maximum or nearly maximum support in both BI and ML analyses, but interestingly if considered together as a single clade the support lowers to 0.84 (PP) and $59 \%$ (BS), although this seems to be mostly influenced by the 16 S gene data (Appendix S5). On the contrary, the haplotype network analysis (Figure 2) recovered the three groups as distinct, separated by 28 substitutions (between groups 4 and 6) and 21 substitutions (between groups 5 and 6) and showed a lack of shared haplotypes. Genetic distances were moderately high, ranging between $5.1 \%$ (between groups 5 and 6), 6.2\% (between groups 4 and 6) and 7.4\% (between groups 4 and 5).

This larger clade, including the three groups $(4+5+6)$, contains only specimens from the Mediterranean Sea and one from the Eastern Atlantic island of Selvagem Grande (Madeira Archipelago). They are all characterized by a distinct anatomical feature among Haminoea, namely a prostate with a constricted zone between the proximal and distal lobes (see Thompson, 1988). This feature has been described for the type species of the genus $H$. hydatis (Talavera et al., 1987; Tchang, 1931; Thompson, 1976, 1981, 1988) and H. fusari (Álvarez et al., 1993). According to the literature these two species are basically distinguished by the presence of denticulated inner lateral teeth in H. hydatis (Talavera et al., 1987; Tchang, 1931; Vayssière, 1885) while they are smooth in $H$. fusari (Álvarez et al., 1993).

Haminoea hydatis is the type species of the genus described by Linnaeus (1758) based on shells from the Mediterranean Sea. The type specimen illustrated in the webpage of the Linnean Collections, London (https:// linnean-online.org/16897/\#?s=0\&cv=0\&z=0.0365\%2C$0.0109 \% 2 \mathrm{C} 1.232 \% 2 \mathrm{C} 1.503$ ), is a shell about 9 mm in height with a smooth surface. Vayssière (1885) studied specimens from the Gulf of Marseille on the Mediterranean French coast with smooth shells and a radula with inner lateral teeth denticulated, which he identified as $H$. hydatis. Later, Tchang (1931) described the male reproductive system of specimens from the same region as having a prostate with the proximal and distal lobes separated by a narrow tubular region, and Talavera et al. (1987) mentioned a smooth, cylindrical pointed penis. Thus, progressively it became established in the scientific literature the idea that $H$. hydays (originally only known from shells) was characterized by having smooth shells, radulae with denticulated inner lateral teeth and a prostate with a narrow region separating the two lobes. This view was reinforced by the fact that up until the end of the first half of the 20th century, the European fauna of Haminoea was basically restricted to two accepted species; either $H$. hydatis with its small smooth shells or H. navicula with larger and deeply spiralled shells.

However, several species with smooth shells accepted as valid (see Introduction) were described during the second half of the last century, one of them (H. fusari) also with a prostate with two lobes separated by a narrow region, but with smooth radular inner lateral teeth (Álvarez et al., 1993). But the study of the holotype of $H$. fusari (MNCN 15.05/5356) revealed in fact the presence of mostly smooth inner lateral teeth, but interestingly some of them had the lower half denticulated. Intraspecific radular variability was described by Malaquias and Cervera (2006) for H. navicula and might occur also in specimens identified as $H$. fusari. This would basically make the two species anatomically undistinguishable and thus likely conspecific, rendering the name $H$. fusari a junior synonym of $H$. hydatis. In addition, the colour patterns of specimens in groups 4 and 5 are alike (data not available for group 3 ), with large unpigmented peri-ocular areas, dark upper sides of the parapodial lobes and fine bright-white dots along the edge of the cephalic shield, which further supports their conspecificity (see Figure 1).

### 4.4 The eastern Pacific species

In the Eastern Pacific coastlines of North and Central America, there are three species of Haminoea commonly recognized as valid between Alaska and Panama, namely H. ovalis, H. virescens and H. vesicula (Behrens
\& Hermosillo, 2005; Hermosillo et al., 2006; Valdés \& Camacho-Garcia, 2004). For this region, our analyses recognized lineages compatible with these three species. The species $H$. virescens (group 11) with pear-shaped shells and monolobated prostates (Gibson \& Chia, 1989; Valdés, 2019; personal observations) split off as sister to the eastern Atlantic/Indian Ocean complex H. alfreden-sis-H. orbignyana-H. antillarum, although with no support ( $\mathrm{PP}=0.83, \mathrm{BS}=59 \%$; Figure 1). A similar pattern was found for the species $H$. vesicula (group 8) characterized by a globose-quadrate shell and bilobed prostate (Gibson \& Chia, 1989; Valdés, 2019; personal observations), which was rendered sister to the eastern European H. orteai, but again with no support ( $\mathrm{PP}=0.51, \mathrm{BS}=45 \%$; Figure 1). This phylogenetic pattern with putative speciation across these two oceanic realms could be explained by processes related to the uplift of the Isthmus of Panama, which separated the Atlantic from the eastern Pacific around 3 Mya (Coates \& Obando, 1996), however, the low support values hamper any sound explanation.

Finally, we retrieved a lineage morphologically compatible with what has been named in recent literature as H. ovalis (Behrens \& Hermosillo, 2005; Hermosillo et al., 2006; Valdés \& Camacho-Garcia, 2004). However, as explained in the Introduction this name applies to an Indo-West Pacific species in the genus Lamprohaminoea (Oskars \& Malaquias, 2020c; Pease, 1868). This Eastern Pacific species has globose smooth shells and a body doted by abundant tiny orange dots. We could not find any available name fitting the features known for this lineage, which thus may represent an undescribed species.

### 4.5 Notes on other Atlantic species

An additional five species were recovered by the phylogenetic and species delimitation analyses ( $H$. navicula [group 1], Haminoea sp. [group 2], Haminoea sp. [group 3], H. orteai [group 9], Haminoea sp. [group 12]). The species $H$. navicula is well established, characterized by large shells, with conspicuous spiral striae and an armed penis (for details see Lobo-da-Cunha et al., 2018; Malaquias \& Cervera, 2006). The species $H$. orteai was the first described European species characterized by a penis with an apical crest with 10 lamellae and the peculiarity of lacking an unpigmented periocular area in the cephalic shield (Talavera et al., 1987). However, our observations revealed that the description of this latter feature is not entirely accurate; in fact, like all other species of the genus, $H$. orteai has an unpigmented periocular area yet rounded and of a much smaller diameter. This can even be seen in the original description of the species (Talavera et al., 1987: 66, figure 15). Later, two additional species also with
penises with apical crests were described for European waters, namely $H$. templadoi (García et al., 1991) and $H$. exigua (Schaefer, 1992). These two species are morphoanatomically very similar to $H$. orteai when it comes to the radula, the male reproductive system, the shell and the reduced diameter of the periocular area. Haminoea templadoi was described as having a shell with transverse folds interconnecting longitudinal growth lines and a radula with the first two inner lateral teeth denticulated (García et al., 1991: 396, figures 2, 3), but the study of the holotype (MNCN 15.05/854) showed a radula with only the inner lateral teeth denticulated. Moreover, according to our interpretation, the distinct shell structure likely relates to the comparatively larger size of the shell $(H=21.6 \mathrm{~mm})$. A thorough systematic review of species combining detailed morphological work and the current phylogenetic framework is necessary to address the putative conspecificity of the species $H$. orteai, H. templadoi and $H$. exigua.

Three additional and unidentified species were rendered by our analyses; Haminoea sp. [group 12] from Brazil, and two species represented by single individuals, namely Haminoea sp. [group 2] from Croatia and Haminoea sp. [group 3] from Roses, Girona, Spain (Mediterranean Sea). The specimens from Brazil externally resemble $H$. 'ovalis' from the eastern Pacific and are interestingly part of a fully supported clade which includes the later species (Figure 1). The remaining two Mediterranean species might represent undescribed taxa, but this requires additional work to be confirmed. The colourations of these species are quite unique among European species with more or less uniform orangish and brownish background colour patterns (Figure 1).

## 4.6 | Concluding remarks

The genus Haminoea is a difficult taxonomic group with many available names introduced based on shell descriptions alone or described based on their morpho-anatomical features outside a phylogenetic molecular framework, several of them grounded on subtle differences.

The current literature consensus accepts eight species as valid in the Eastern Atlantic (including the amphiAtlantic H. 'elegans'), four in the Western Atlantic (including H. 'elegans'), three in the Eastern Pacific and one in temperate South Africa. As explained in the Discussion, we open the possibility that several of the Eastern Atlantic taxa, presently regarded as valid species, might be conspecific (e.g. H. orteai, H. templadoi, H. exigua). A re-evaluation of the literature and type material, preliminary anatomical work, combined with our molecular phylogenetic framework seem to indicate that several of the subtle differences used in the past to
introduce new species might not be sound enough. On the contrary, there were several cases where our results were not sufficient to reach definitive conclusions about the taxonomic status of certain species (e.g. H. alfredensis, H. antillarum, H. elegans, H. orbignyana) and further morphological work is necessary to understand their diversity and draw a robust taxonomic hypothesis. A preliminary combination of conchological, morphological and phylogenetic data demonstrated that putative cases of cryptic diversity may, in fact, reflect previously detected differences in shell characters that led to the description of species currently considered invalid (e.g. the cryptic species complex H. 'elegans').

There are several paradigmatic examples of pervasive nomenclature confusion that need to be evaluated, such as the status of the type species $H$. hydatis, H. elegans-a name introduced for a European species but largely in use for western Atlantic animals, and $H$. ovalis-an IWP species in the genus Lamprohaminoea, but a name commonly used to also refer to an Eastern Pacific lineage.

With this work, progress was made to underhand the diversity of Haminoea snails and the relationships between species, and for the first time a phylogeny of the genus is presented. This new framework combined with a detailed study of shells, morpho-anatomy of wet specimens, revision of original descriptions and type material, can help solve the many issues remaining with the taxonomy of Haminoea snails in the Atlantic and Eastern Pacific Oceans.

## ACKNOWLEDGEMENTS

This work would have not been possible without the generosity of many colleagues, friends and institutions that kindly provided access to specimens, museum collections and type material. We are especially indebted to Alan Hodgson (South Africa), George Branch (South Africa), Charles Griffiths (South Africa), Nelson Miranda (South Africa), Alberto Zecalupo (Tunisia), Andrea Spinelli (Italy), Guido Villani (Italy), Angelo Vazzana (Italy), Marta Pola (Italy), Andrea Zamora (Mexico), Jazmin Ortigosa (Mexico), Andreia Salvador and Kathy Way (Natural History Museum, London), António D. Abreu (Museum Municipal do Funchal (História Natural)), Ariane Dimitris (Florida, USA), Craig Hoover (California, USA), Jeff Goddard (California, USA). Marlo Krisberg (Florida, USA), Luiz Simone (Museu de Zoologia, University of São Paulo, Brazil), Carlo Cunha (Brazil), Vinicius Padula (Brazil), Juliana Bahia (Brazil), Colin Redfern (Bahamas), Edwin Cruz-Rivera (Egypt), Emilio Rólan (Spain), Irene Figueroa and Guillem Mas (Spain), José Martín (Spain), Manuel Ballesteros (Spain), Karla Araújo (Spain), Gonçalo Calado (Portugal and Mexico), Jakov Prkić (Croatia), Jeroen Goud (Naturalis Biodiversity Center, Leiden), José
P. Borges (Canary Islands), José Templado and Rafael Araujo (Museo de Ciencias Naturales de Madrid), Julio Magaña (Mexico and Costa Rica), Marina Pudibouskaya (France and Bahamas), Justine Siegwald (France), Pirjo Peller (France), Juan Lucas Cervera (Bermuda, Brazil and Spain), Samuel Narciso (Venezuela), Manuel Caballer (Venezuela), Matthias Glaubrecht (Museum für Naturkunde, Berlin), Nenibarini Zabbey (Nigeria), Noufal Tamsouri (Morocco), Patrick Krug (Jamaica, Baja California, Mexico), Peter Wirtz (Cape Verde), Philippe Bouchet and Virginie Héros (Muséum national d'Histoire naturelle, Paris), Ricardo Serrão Santos (Department of Oceanography and Fisheries, University of the Azores), Rita Coelho (Portugal), Thierry Backeljau (Royal Belgium Institute of Natural Sciences).

We are thankful to Chari Martín-Hervás (University of Cádiz) for help with the bPTP analysis, and to David Rees and Louise Lindblom (DNA Lab, Department of Biological Sciences/University Museum of Bergen, University of Bergen) for help with molecular work, and to Irene Heggstad (Department of Geological Sciences, University of Bergen) for help with electron microscopy work.

MAE Malaquias are grateful to David Reid (Natural History Museum, London) for hosting him as a postdoctoral fellow between 2006 and 2007 to work with Haminoea snails.

We are indebted to the Department of Marine Resources, Government of The Bahamas for granting a collecting permit (MAF/FIS/12) to MAE Malaquias to sample in Eleuthera Island.

We are very thankful to JL Cervera (University of Cádiz, Spain) for inviting MAE Malaquias to field trips to the Azores, Bermuda and Brazil as part of four research projects (CGL2010-17187, Spanish Ministry of Science, Innovation and Universities; PR2018-039, University of Cadiz; CGL2006-05182, Spanish Ministry of Education and Science; CTM2008-05228-E/MAR Spanish Ministry of Science and Innovation). To the Meltzers Foundation, University of Bergen (Norway) for awarding funding to MAE Malaquias for a field trip to the Bahamas. To the Fundação para a Ciência e Tecnologia (FCT, Portuguese Government, SFRH/BPD/26209/2006) for a post-doctoral grant to MAE Malaquias.

Visits to several European natural history museums received support from the SYNTHESYS Project http:// www.synthesys.info/ which is financed by European Community Research Infrastructure Action under the FP6 'Structuring the European Research Area’ Programme.

## ORCID

Ángel Valdés © https://orcid.org/0000-0002-2347-4896 Manuel António E. Malaquias © https://orcid. org/0000-0002-9668-945X

## REFERENCES

Adams, A. (1850). Monograph of the family Bullidae. In G. B. Sowerby (Ed.), Thesaurus Conchyliorum, or monographs of genera of shells (Vol. II, pp. 553-608). Sowerby.
Akaike, H. (1974). A new look at the statistical model identification. IEEE Transactions on Automatic Control, 19, 716-723. https:// doi.org/10.1109/TAC.1974.1100705
Alfaro, M. E., Zoller, S., \& Lutzoni, F. (2003). Bayes or bootstraps? A simulation study comparing the performance of Bayesian Markov chain Monte Carlo sampling and bootstrapping in assessing phylogenetic confidence. Molecular Biology and Evolution, 20, 255-266. https://doi.org/10.1093/molbev/msg028
Álvarez, L. A., Garcia, F. J., \& Villani, G. (1993). A new Mediterranean species of Haminaea Leach, 1820 (Gastropoda: Opisthobranchia: Cephalaspidea). Journal of Molluscan Studies, 59, 339-345. https://doi.org/10.1093/mollus/59.3.339
Austin, J., Gosliner, T., \& Malaquias, M. A. E. (2018). Systematic revision, diversity patterns, and trophic ecology of the tropical Indo-West Pacific sea slug genus Phanerophthalmus A. Adams, 1850 (Cephalaspidea, Haminoeidae). Invertebrate Systematics, 32, 1336-1387. https://doi.org/10.1071/IS17086
Baker, F., \& Hanna, G. D. (1927). Expedition of the California Academy of Sciences to the Gulf of California in 1921. Proceedings of the California Academy of Sciences, 16, 123-135.
Bartsch, P. (1915). Report on the Turton collection of South African marine mollusks, with additional notes on other South African shells contained in the United States National Museum. Bulletin of the United States National Museum, 91, 1-305.
Behrens, D. W., \& Hermosillo, A. (2005). Eastern Pacific nudibranchs. A guide to the opisthobranchs from Alaska to Central America. Sea Challengers 137 p .
Bernard, P. A. (1984). Coquilages du Gabon.
Boulch-Bleas, D. (1983). A propos du regime alimentaire d' Haminea hydatis (Linne, 1758), (Mollusque, Opisthobranche). Haliotis, 13, 45-52.
Briggs, J. C. (1974). Marine zoogeography. McGraw-Hill.
Briggs, J. C. (1995). Global biogeography. Elsevier.
Burn, R., \& Thompson, T. E. (1998). Order Cephalaspidea. In P. L. Beesley, G. J. B. Ross, \& A. A. Wells (Eds.), Fauna of Australia, Mollusca: The southern synthesis (Part B, VIII, Vol. 5', pp. 943959). CSIRO Publishing.

Caballer, M., Ortea, J., Rivero, N., Carias, G., Malaquias, M. A. E., \& Narciso, S. (2015). The opisthobranch gastropods (Mollusca: Heterobranchia) from Venezuela: An annotated and illustrated inventory of species. Zootaxa, 4034, 201-256. https://doi. org/10.11646/zootaxa.4034.2.1
Carmona, L., Gosliner, T. M., Pola, M., \& Cervera, J. L. (2011). A molecular approach to the phylogenetic status of the aeolid genus Babakina Roller, 1973 (Nudibranchia). Journal of Molluscan Studies, 77(4), 417-422. https://doi.org/10.1093/mollus/eyr029
Cervera, J. L., Calado, G., Gavaia, C., Malaquias, M. A., Templado, J., Ballesteros, M. B. V., Garcia-Gomez, J. C., \& Megina, C. (2004). An annotated and updated checklist of the ophisthobranchs (Mollusca: Gastropoda) from Spain and Portugal (including islands and archipelagos). Boletín del Instituto Español de Oceanografía, 20(1-4), 1-122.
Churchill, C. K., Valdés, A., \& Ó Foighil, D. (2014). Afro-Eurasia and the Americas present barriers to gene flow for the cosmopolitan neustonic nudibranch Glaucus atlanticus.

Marine Biology, 161, 899-910. https://doi.org/10.1007/s0022 7-014-2389-7
Clement, M., Snell, Q., Walker, P., Posada, D., \& Crandall, K. (2002). TCS: estimating gene genealogies. In Parallel and distributed processing symposium, international (Vol. 3, p. 184). IEEE Computer Society.
Coates, A. G., \& Obando, J. A. (1996). The geological evolution of the Central American isthmus. In J. B. C. Jackson, A. F. Budd, \& A. G. Coates (Eds.), Evolution and environment in tropical America (pp. 21-56). University of Chicago Press.
Conrad, T. A. (1846). Descriptions of new species of fossil and recent shells and corals. Proceedings of the Academy of Natural Sciences of Philadelphia, 3, 19-27.
Da Costa, E. M. (1778). The British conchology. Elmsley and Robson Booksellers.
Darriba, D., Taboada, G. L., Doallo, R., \& Posada, D. (2012). jModelTest 2: More models, new heuristics and parallel computing. Nature Methods, 9, 772. https://doi.org/10.1038/nmeth. 2109
d'Orbigny, A. (1841). In R. De la Sagra (Ed.), Mollusques. In 'Histoire Physique, Politique et Naturelle de l'Ile de Cuba' (Vol. 1, pp. 1240). Arthus Bertrand.

Edgar, R. C. (2004). MUSCLE: Multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Research, 32(5), 1792-1797. https://doi.org/10.1093/nar/gkh340
Eilertsen, M. H., \& Malaquias, M. A. E. (2013). Systematic revision of the genus Scaphander (Gastropoda, Cephalaspidea) in the Atlantic Ocean, with a molecular phylogenetic hypothesis. Zoological Journal of the Linnean Society, 167(3), 389-429. https://doi.org/10.1111/zoj. 12013
Felsenstein, J. (1985). Confidence limits on phylogenies: An approach using the bootstrap. Evolution, 39, 783-791. https://doi. org/10.2307/2408678
Férussac, D. D. (1822). Dictionnaire Classique d'Histoire Naturelle (Vol. 2). Rey et Gravier, Libraires-Editeurs.
Floeter, S. R., Rocha, L. A., Robertson, D. R., Joyeux, J. C., SmithVaniz, W. F., Wirtz, P., Edwards, A. J., Barreiros, J. P., Ferreira, C. E. L., Gasparini, J. L., Brito, A., Falcón, J. M., Bowen, B. W., \& Bernardi, G. (2008). Atlantic reef fish biogeography and evolution. Journal of Biogeography, 35, 22-47.
Folmer, O., Black, M., Hoeh, W., Lutz, R., \& Vrijenhoek, R. (1994). DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. Molecular Marine Biology and Biotechnology, 3, 294-299.
García, F. J., Domínguez, M., \& Troncoso, J. S. (2008). Opistobranquios de Brasil. Descripción y distribución de opistobranquios del litoral de Brasil y del Archipiélago Fernando de Noronha. 215 p. Feito S. L.
García, F. J., Pérez-Hurtado, A., \& García-Gómez, J. C. (1991). Haminaea templadoi, a new species of cephalaspidean opisthobranch from the Atlantic Iberian coast. Journal of Molluscan Studies, 57, 395-399. https://doi.org/10.1093/mollus/57.4.395
Golestani, H., Crocetta, F., Padula, V., Camacho-Garcia, Y., Langeneck, J., Poursanidis, D., Pola, M., Yokeş, M. B., Cervera, J. L., Jung, D. W., \& Gosliner, T. M. (2019). The little Aplysia coming of age: From one species to a complex of species complexes in Aplysia parvula (Mollusca: Gastropoda: Heterobranchia). Zoological Journal of the Linnean Society, 187(2), 279-330. https://doi.org/10.1093/zoolinnean/zlz028
Gosliner, T. M. (1987). Nudibranchs of Southern Africa. A guide to Opisthobranch Molluscs of Southern Africa. Sea Challengers: Monterey, CA, USA; Jeff Hartman: El Cajon, CA, USA; EJ.

Brill: Leiden, Netherlands; in association with the California Academy of Sciences.
Gosliner, T. M., Behrens, D. W., \& Valdés, Á. (2008). Indo-Pacific nudibranchs and sea slugs: A field guide to the World's most diverse Fauna. Sea Challengers Natural History Books and California Academy of Sciences.
Gould, A. A. (1855). Catalogue of shells collected by W. P. Blake, with descriptions of the new species. In W. P. Blake (Ed.), Preliminary geological report of a reconnaissance and survey in California in connection with explorations for a predictable railway route from the Mississippi River to the Pacific Ocean in 1853 by Lt. R. S. Williamson'. Doc. 129, 33rd Congress, 1st session. Appendix, part 2 (pp. 22-28). House of Representatives.
Gray, J. E. (1825). A list and description of some species of shells not taken notice of by Lamarck. Annals of Philosophy, 9, 134-415.
Gyory, J., Mariano, A. J., \& Ryan, E. H. (2011). The loop current, Ocean Surface Currents. https://oceancurrents.rsmas.miami. edu/caribbean/loop-current.html
Habe, T. (1952). Atydae in Japan. Illustrated Catalogue of Japanese Shells, 20, 137-152.
Hermosillo, A., Behrens, D. W., \& Ríos, E. (2006). Opistobranquios de México. CONABIO 143 p.
Huelsenbeck, J. P., \& Ronquist, F. (2001). MRBAYES: Bayesian inference of phylogenetic trees. Bioinformatics, 17(8), 754-755. https://doi.org/10.1093/bioinformatics/17.8.754
Huelsenbeck, J. P., Ronquist, F., Nielsen, R., \& Bollback, J. P. (2001). Bayesian inference of phylogeny and its impact on evolutionary biology. Science, 294, 2310-2314. https://doi.org/10.1126/scien ce. 1065889
Jörger, K. M., Norenburg, J. L., Wilson, N. G., \& Schrödl, M. (2012). Barcoding against a paradox? Combined molecular species delineations reveal multiple cryptic lineages in elusive meiofaunal sea slugs. BMC Evolutionary Biology, 12, 1-18.
Kearse, M., Moir, R., Wilson, A., Stones-Havas, S., Cheung, M., Sturrock, S., Buxton, S., Cooper, A., Markowitz, S., \& Duran, C. (2012). Geneious basic: An integrated and extendable desktop software platform for the organization and analysis of sequence data. Bioinformatics, 28(12), 1647-1649. https://doi. org/10.1093/bioinformatics/bts199
Kienberger, K., Carmona, L., Pola, M., Padula, V., Gosliner, T. M., \& Cervera, J. L. (2016). Aeolidia papillosa (Linnaeus, 1761) (Mollusca: Heterobranchia: Nudibranchia), single species or a cryptic species complex? A morphological and molecular study. Zoological Journal of the Linnean Society, 177(3), 481506. https://doi.org/10.1111/zoj. 12379

Kumar, S., Stecher, G., \& Tamura, K. (2016). MEGA7: Molecular evolutionary genetics analysis version 7.0 for bigger datasets. Molecular Biology and Evolution, 33, 1870-1874. https://doi. org/10.1093/molbev/msw054
Leach, W. E. (1852). A synopsis of the Mollusca of Great Britain. John van Voorst.
Lee, T., \& Ó Foighil, D. (2004). Hidden Floridian biodiversity: Mitochondrial and nuclear gene trees reveal four cryptic species within the scorched mussel, Brachidontes exustus, species complex. Molecular Ecology, 13, 3527-3542. https://doi. org/10.1111/j.1365-294X.2004.02337.x
Leigh, J.W., \& Bryant, D. (2015). POPART:Full-feature software for haplotype network construction. Methods in Ecology and Evolution, 6(9), 1110-1116. https://doi.org/10.1111/2041-210X. 12410

Librado, P., \& Rozas, J. (2009). DnaSP v5: A software for comprehensive analysis of DNA polymorphism data. Bioinformatics, 25, 1451-1452. https://doi.org/10.1093/bioinformatics/ btp187
Linnaeus, C. (1758). Systema Naturae per regna Tria Naturae, Secundum classes, ordines, genera, species, cum Characteribus, Differentiis, Synonymis, Locis. Editio decima (edn 10), reformat'a. Laurentius Salvius.
Littlewood, D. T. J., Curini-Galletti, M., \& Herniou, E. A. (2000). The interrelationships of Proseriata (Platyhelminthes: Seriata) tested with molecules and morphology. Molecular Phylogenetics and Evolution, 16(3), 449-466. https://doi.org/10.1006/ mpev.2000.0802
Lobo-da-Cunha, A., Alves, Â., Oliveira, E., \& Malaquias, M. (2018). Histological, histochemical and ultrastructural investigation of the male copulatory apparatus of Haminoea navicula (Gastropoda, Cephalaspidea). Journal of Morphology, 279(4), 554-565. https://doi.org/10.1002/jmor. 20790
Macnae, W. (1962). Tectibranch molluscs from southern Africa. Annals of the Natal Museum, 15, 183-199.
Malaquias, M. A. E. (2014). New data on the heterobranch gastropods ('opisthobranchs') for The Bahamas (tropical western Atlantic Ocean). Marine Biodiversity Records, 7, e27. https://doi. org/10.1017/S175526721400030X
Malaquias, M. A. E., Mackenzie-Dodds, J., Bouchet, P., Gosliner, T., \& Reid, D. G. (2009). A molecular phylogeny of the Cephalaspidea sensu lato (Gastropoda: Euthyneura): Architectibranchia redefined and Runcinacea reinstated. Zoologica Scripta, 38, 23-41. https://doi.org/10.1111/j.1463-6409.2008.00354.x
Malaquias, M. A. E., \& Cervera, J. L. (2006). The genus Haminoea (Gastropoda: Cephalaspidea) in Portugal, with a review of the European species. Journal of Molluscan Studies, 72, 89-103. https://doi.org/10.1093/mollus/eyi052
Malaquias, M. A. E., Condinho, S., Cervera, J. L., \& Sprung, M. (2004). Diet and feeding biology of Haminoea orbygniana (Mollusca: Gastropoda: Cephalaspidea). Journal of the Marine Biological Association of the United Kingdom, 84, 767-772. https://doi. org/10.1017/S0025315404009890h
Malaquias, M. A. E., Martínez, E., \& Abreu, A. D. (2002). Cephalaspidea (Mollusca: Opisthobranchia) of the Madeira archipelago and Selvagens Islands. American Malacological Bulletin, 17, 65-83.
Marcus, E., \& Marcus, E. (1967). American opisthobranch mollusks. Studies in Tropical Oceanography, 6, 1-256.
Marcus, E. V. (1976). Marine euthyneuran gastropods from Brazil. Studies on Neotropical Fauna and Environment, 11, 5-23.
Martínez, E., \& Ortea, J. (1997). Haminoea elegans (Gray, 1825) (Opisthobranchia: Cephalaspidea), a truly amphi-Atlantic species. The Veliger, 40(4), 281-291.
Mikkelsen, P. M., \& Bieler, R. (2000). Marine bivalves of the Florida keys: Discovered biodiversity. Geological Society, London, Special Publications, 177(1), 367-387.
MolluscaBase. (2022). MolluscaBase. Haminoea elegans (Gray, 1825). Accessed through: World Register of Marine Species at: https:// www.marinespecies.org/aphia.php?p=taxdetails\&id=140071
Olson, P. D., Cribb, T. H., Tkach, V. V., Bray, R. A., \& Littlewood, D. T. J. (2003). Phylogeny and classification of the Digenea (Platyhelminthes: Trematoda). International Journal of Parasitology, 33, 733-755.

Oskars, T. R., Bouchet, P., \& Malaquias, M. A. E. (2015). A new phylogeny of the Cephalaspidea (Gastropoda: Heterobranchia) based on expanded taxon sampling and gene markers. Molecular Phylogenetics and Evolution, 89, 130-150. https://doi. org/10.1016/j.ympev.2015.04.011
Oskars, T. R., \& Malaquias, M. A. E. (2019). A molecular phylogeny of the Indo-West Pacific species of Haloa sensu lato gastropods (Cephalaspidea: Haminoeidae): Tethyan vicariance, generic diversity, and ecological specialization. Molecular Phylogenetics and Evolution, 139, 106557. https://doi.org/10.1016/j. ympev.2019.106557
Oskars, T. R., \& Malaquias, M. A. E. (2020a). Systematic revision of the Indo-West Pacific mangrove-associated snails of the genus Bakawan (Cephalaspidea: Haminoeidae). Journal of Molluscan Studies, 86, 323-341.
Oskars, T. R., \& Malaquias, M. A. E. (2020b). The temperate Australasian genus Papawera Oskars and Malaquias, 2019 (Gastropoda: Cephalaspidea: Haminoeidae), with a redescription of P. zelandiae and P. maugeansis. Journal of Natural History, 54, 1343-1362.
Oskars, T. R., \& Malaquias, M. A. E. (2020c). Systematic revision of the Indo-West Pacific colourful bubble-snails of the genus Lamprohaminoea Habe, 1952 (Cephalaspidea: Haminoeidae). Invertebrate Systematics, 34, 727-756.
Oskars, T. R., Too, C. C., Rees, D., Mikkelsen, P. M., Willassen, E., \& Malaquias, M. A. E. (2019). A molecular phylogeny of the gastropod family Haminoeidae sensu lato (Heterobranchia: Cephalaspidea): A generic revision. Invertebrate Systematics, 33, 426-472.
Palumbi, S. R. (1996). Nucleic acids II: The polymerase chain reaction. In D. M. Hillis, C. Moritz, \& B. K. Mable (Eds.), Molecular systematics (pp. 205-247). Sinauer \& Associates Inc.
Palumbi, S. R., Martin, A., Roman, S., McMillan, W., Stice, L., \& Grabowski, G. (1991). The simple fool's guide to PCR. Special Publications of the Department of Zoology, University of Hawaii.
Pease, W. H. (1868). Descriptions of marine Gasteropodae, inhabiting Polynesia. American Journal of Conchology, 4, 71-80.
Petuch, E. J. (1987). New Caribbean molluscan faunas (154 pp., 29 pls; addendum 2 pp., 1 pl). The Coastal Education and Research Foundation.
Pilsbry, H. A. (1921). Marine molluscs of Hawaii, XIV, XV. Proceedings of the Academy of Natural Sciences of Philadelphia, 72, 360-382.
Pruvot-Fol, A. (1954). Mollusques Opisthobranches. Faune de France 58. Lechevalier.

Puillandre, N., Brouillet, S., \& Achaz, G. (2021). ASAP: Assemble species by automatic partitioning. Molecular Ecology Resources, 21(2), 609-620. https://doi.org/10.1111/1755-0998.13281
Puillandre, N., Lambert, A., Brouillet, S., \& Achaz, G. (2012). ABGD, automatic barcode gap discovery for primary species delimitation. Molecular Ecology, 21, 1864-1877. https://doi. org/10.1111/j.1365-294X.2011.05239.x
Rambaut, A., \& Drummond, A. (2009). FigTree v1.3.1. Computer program and documentation distributed by the author.
Rambaut, A., Drummond, A. J., Xie, D., Baele, G., \& Suchard, M. A. (2018). Posterior summarization in Bayesian phylogenetics using tracer 1.7. Systematic Biology, 67(15), 901-904.
Redfern, C. (2001). Bahamian seashells: A thousand species from Abaco, Bahamas. Bahamianseashells.com, Inc 280 pp.

Redfern, C. (2013). Bahamian seashells: 1161 species from Abaco, Bahamas. Bahamian seashells.com, Inc 501 pp.
Reeb, C. A., \& Avise, J. C. (1990). A genetic discontinuity in a continuously distributed species: Mitochondrial DNA in the American oyster, Crassostrea virginica. Genetics, 124(2), 397406. https://doi.org/10.1093/genetics/124.2.397

Rios, E. (2009). Compedium of Brazilian sea shells (p. 668p). Editora Evangraf Lda.
Rocha, L. A., Robertson, D. R., Roman, J., \& Bowen, B. W. (2005). Ecological speciation in tropical reef fishes. Proceedings of the Royal Society B: Biological Sciences, 272, 573-579. https://doi. org/10.1098/2004.3005
Rolán, E., \& Ryall, P. (1999). Checklist of the Angolan marine molluscs. Reseñas Malacologicas (Sociedad Española de Malacologia), 10, 1-132.
Rudman, W. B. (1971). On the opisthobrach genus Haminoea Turton \& Kingston. Pacific Science, 25, 545-559.
Schaefer, K.(1992). Haminaea exigua (Gastropoda, Opisthobranchia), a new cephalaspid species from the Mediterranean Sea. Journal of Molluscan Studies, 58, 329-336. https://doi.org/10.1093/ mollus/58.3.329
Schaefer, K. (1996). Review of data on cephalaspid reproduction, with special reference to the genus Haminaea (Gastropoda, Opisthobranchia). Ophelia, 45, 17-37. https://doi. org/10.1080/00785326.1996.10432460
Siesser, W. G. (1980). Late Miocene origin of the Benguela upwelling system of northern Namibia. Science, 208, 283-285. https://doi. org/10.1126/science.208.4441.283
Sowerby, G. B. I. (1833). The genera of recent and fossil shells, for the use of students, in conchology and geology. Published in 42 numbers. Vol. 1, pls 1-126 [1821-1825]; vol. 2, pls 127-262 + text (unpaginated) [1825-1834].
Stamatakis, A. (2014). RAxML version 8: A tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics, 30, 1312-1313. https://doi.org/10.1093/bioinformatics/btu033
Steinke, D., Prosser, S. W., \& Hebert, P. D. (2016). DNA barcoding of marine metazoans. In S. J. Bourlat (Ed.), Marine genomics: Methods and protocols (pp. 155-168). Humana Press.
Swainson, W. (1840). A treatise on malacology; or the natural classification of shells and shell fish. Longman, Orme, Brown, Green and Longman.
Talavera, G., \& Castresana, J. (2007). Improvement of phylogenies after removing divergent and ambiguously aligned blocks from protein sequence alignments. Systematic Biology, 56, 564-577. https://doi.org/10.1080/10635150701472164
Talavera, P., Murillo, L., \& Templado, J. (1987). The genus Haminoea Turton and Kingston, 1830 (Opisthobranchia, Bullomorpha) in the south east of Spain with the description of a new species. Bollettino Malacologico, 23, 53-68.
Tchang, S. (1931). Contribution à l'étude des mollusques opisthobranches de la côte Provençale. PhD thesis, Imprimerie de Trevoux, Lyon, France.
Thompson, T. E. (1976). Biology of opisthobranch molluscs (Vol. 1). The Ray Society.
Thompson, T. E. (1981). Taxonomy of three misunderstood opisthobranchs from the northern Adriatic Sea. Journal of Molluscan Studies, 47, 73-79. https://doi.org/10.1093/oxfordjournals.mollus.a065559
Thompson, T. E. (1988). Molluscs: Benthic opisthobranchs (Mollusca: Gastropoda). E. J. Brill/Dr. W. Backhuys.

Thompson, T. E., \& Brown, G. H. (1976). British Opisthobranch Molluscs. The Linnean Society of London \& Academic Press 203 p.
Valdés, A. (2019). Northeast Pacific benthic shelled sea slugs. Zoosymposia, 13, 242-304. https://doi.org/10.11646/zoosy mposia.13.1.21
Valdés, A., \& Camacho-Garcia, Y. (2004). "Cephalaspidean" Heterobranchs (Gastropoda) from the Pacific coast of Costa Rica. Proceedings of the California Academy of Sciences, 55(ser. 4), 459-497.

Valdés, Á., Hamann, J., Behrens, D., \& DuPont, A. (2006). Caribbean Sea slugs. A field guide to the opisthobranch mollusks from the tropical northwestern Atlantic. Sea Challengers Natural History Books 309 p.
Vayssière, M. A. (1885). Recherches zoologiques et anatomiques sur les mollusques Opisthobranches du Golfe de Marseille. Première partie tectibranches. Annales du Musée d'histoire Naturelle de Marseille, 2(3), 1-181.
Vermeij, G. J., \& Rosenberg, G. (1993). Giving and receiving: The tropical Atlantic as a donor and recipient region for invading species. American Malacological Bulletin, 10, 181-194.
Williams, S. T., Reid, D. G., \& Littlewood, D. T. J. (2003). A molecular phylogeny of the Littorininae (Gastropoda: Littorinidae): Unequal evolutionary rates, morphological parallelism, and
biogeography of the Southern Ocean. Molecular Phylogenetics and Evolution, 28(1), 60-86. https://doi.org/10.1016/S1055 -7903(03)00038-1
Zhang, J., Kapli, P., Pavlidis, P., \& Stamatakis, A. (2013). A general species delimitation method with applications to phylogenetic placements. Bioinformatics, 29(22), 2869-2876. https://doi. org/10.1093/bioinformatics/btt499

## SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Turani, M., Valdés, Á., \& Malaquias, M. A. E. (2023). Molecular phylogeny of the marine snail genus Haminoea (Gastropoda, Cephalaspidea): A framework to study marine diversity and speciation. Zoologica Scripta, 00, 1-26. https://doi.org/10.1111/zsc. 12627


[^0]:    This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.
    © 2023 The Authors. Zoologica Scripta published by John Wiley \& Sons Ltd on behalf of Royal Swedish Academy of Sciences.

