# Phylogenetic revision of the shrimp genera Ephyrina, Meningodora and Notostomus (Acanthephyridae: Caridea) 

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#### Abstract

The shrimp genera Ephyrina, Meningodora and Notostomus have an unusual carapace strengthened with carinae and a half-serrated mandible, which may suggest a possible monophyly of this group. Here we test this hypothesis and present the first phylogenetic study of these genera based on 95 morphological characters (all valid species coded) and six molecular markers ( $71 \%$ of valid species sequenced). Representatives of all genera of Oplophoridae (sister to Acanthephyridae) were outgroups, 32 species belonging to all genera and potentially different clades of Acanthephyridae were ingroups. Both morphological and molecular analyses retrieve trees with similar topology. Our results reject the hypothesis of a clade formed by Ephyrina + Meningodora + Notostomus. We show that Ephyrina and Notostomus are monophyletic, both on morphological and on molecular trees, Meningodora gains support only on morphological trees. Evolutionary traits in the Ephyrina and Meningodora + Notostomus clades are different. Synapomorphies are mostly linked to adaptations to forward motion in Ephyrina (oar-like meri and ischia of pereopods, stempost-like rostrum) and to progressive strengthening of the carapace and pleon in Meningodora and Notostomus (net of sharp carinae). Unusual mandibles evolved in the clades independently and represent convergent adaptations to feeding on gelatinous organisms.


ADDITIONAL KEYWORDS: Crustacea - evolution - phylogeny - plankton biology - shrimp.

## INTRODUCTION

Among pelagic decapods, Oplophoroidea is one of the most diverse superfamilies occurring in the widest geographic and depth ranges. Indeed, nearly one hundred species have been recorded from polar to equatorial regions (WoRMS, 2020), from the upper mixed layer to bathyal depths. Their role in pelagic trophic chains is important: Oplophoroidea are a dominant group explaining nearly half of the total zooplankton stock in the Atlantic tropical and equatorial waters (Vereshchaka et al., 2019b).
Historically, Oplophoroidea was considered on morphological grounds as a single family Oplophoridae (e.g. Chace, 1986; De Grave et al., 2009), but molecular data suggest that it consists of two families, Oplophoridae and Acanthephyridae, within the superfamily Oplophoroidea (e.g. Bracken et al., 2009; Chan et al., 2010; Lunina et al., 2019b; WoRMS, 2020).

[^0]Following this two-family concept, we have revised the global fauna of the family Oplophoridae (Lunina et al., 2019b) and now take the next step to start a phylogenetic revision of the family Acanthephyridae, which is more species-rich than Oplophoridae ( 55 vs . 16 currently accepted species, seven vs. three genera; WoRMS, 2020).

Acanthephyridae is morphologically heterogeneous in many aspects (Chace, 1986). For example, the rostrum may have more teeth on the ventral than on the dorsal margin (Heterogenys Chace, 1986) or vice versa (other genera), the hepatic spine and three lateral carinae on the carapace may be present (Kemphyra Chace, 1986) or absent (other genera), the carapace is ventrally dentate (Notostomus A. Milne-Edwards, 1881) or smooth (other genera), the dorsal pleonic carina is absent (Ephyrina Smith, 1885 and Hymenodora Sars, 1877) or developed (other genera), eggs are large and few $(<50)$ (Ephyrina and Hymenodora) or small and numerous ( $>80$ ) (other genera). One of the basic phylogenetic characters in Crustacea, the morphology of the mandible, also varies. In Acanthephyridae, this
character, which is conservative in decapods at family and even superfamily level, may have two different states (Chace, 1986): (1) subtriangular and armed along the entire margin (Fig. 1A) and (2) subtruncate and unarmed in the distal half beyond the apex, except the terminal tooth (Fig. 1B). The first state is common for most pelagic carnivorous shrimps (Burukovsky, 2009), while the second state is restricted only to the genera Ephyrina, Meningodora Smith, 1882 and Notostomus; its adaptive value has never been assessed. In addition to the remarkable mandible, Ephyrina, Meningodora
and Notostomus are characterized by strong ridges and carinae along the lateral sides of the carapace (Fig. 1C-E). These structures are also unusual for most pelagic shrimps, having a more or less smooth and streamlined carapace. Both unusual characters (a half-serrated mandible and a strengthened carapace) may suggest possible monophyly of this group. Here we test this hypothesis and thus start a revision of the global Acanthephyridae fauna. Using morphological and molecular data, we examine whether the unusual mandible and the strengthened carapace represent


Figure 1. Morphological characters of Acanthephyridae.A, typical mandible of Oplophoroidea (exemplified by Acanthephyra brevicarinata). B, mandible of the Ephyrina-Meningodora-Notostomus group (exemplified by Ephyrina bifida). C, schematic carapace view of Notostomus (exemplified by N. elegans). D, schematic carapace view of Meningodora (exemplified by M. compsa). E, schematic carapace view of Ephyrina (exemplified by E. bifida). Dotted lines indicate additional carinae, which are present alongside the obligatorily carinae (solid lines) in part of species.
synapomorphies of a single clade or if they evolved independently in separate clades.

Among 21 currently accepted species of the Ephyrina-Meningodora-Notostomus group (WoRMS, 2020), none has been included in previous morphological phylogenetic analyses and only a few were included in molecular analyses: a single one (Bracken et al., 2009), two (Chan et al., 2010) or seven species (Wong et al., 2015). The previous molecular analyses targeted higher level relationships within Caridea and/or Oplophoroidea, and did not cover the proper diversity of the Ephyrina-MeningodoraNotostomus group. Here we present the first comprehensive phylogenetic analysis based on the simultaneous use of morphological characters and molecular markers. We used 95 morphological characters to encode all valid species of the target genera, and six gene markers for 15 species ( $71 \%$ of currently accepted species). We also included in the analyses representatives of all three genera of Oplophoridae (outgroups). Acanthephyra A. MilneEdwards, 1881, which is morphologically variable and probably polyphyletic (Chace, 1986), was represented in our analysis by 12 species from morphologically different groups.

## MATERIAL AND METHODS

## MORPHOLOGICAL ANALYSIS

Oplophoridae encompasses three genera (Janicella Chace, 1986, Oplophorus Milne-Edwards, 1837 and Systellaspis Spence Bate, 1888) and is considered as a sister-clade to Acanthephyridae (Wong et al., 2015), which includes, in addition to the three analysed genera, Acanthephyra, Heterogenys, Hymenodora and Kemphyra. We chose as the outgroups representatives of the three genera of Oplophoridae: Janicella spinicauda (A. Milne-Edwards, 1883) (Analysis 1), Oplophorus gracilirostris A. Milne-Edwards, 1881 (Analysis 2) and Systellaspis pellucida (Filhol, 1884) (Analysis 3).

We included as the ingroups all valid species of Ephyrina (six species), Meningodora (six) and Notostomus (nine), and representatives of all other genera of Acanthephyridae; the highly diverse and probably polyphyletic (Chace, 1986) genus Acanthephyra was represented by 12 species from potentially different clades (Table 1).

For each included taxon we identified and encoded 95 morphological characters (not weighted, Supporting Information, Appendix S1), which were combined into four morphological groups (Fig. 2): carapace (characters $0-32$ in Supporting Information, Appendix S1), pleon + telson (33-55), mouthparts (58-74) and pereopods (75-92).

The dataset (Supporting Information, Appendix S2) was handled and analysed using a combination of programs using maximum parsimony settings: WINCLADA/NONA and TNT (Nixon, 1999; Goloboff et al., 2000). Trees were generated in TNT with 30000 trees in memory, under the 'traditional search' (branch-and-bound) algorithms. Relative stability of clades was assessed by standard bootstrapping (sample with replacement) with 10000 pseudoreplicates and by Bremer support (algorithm TBR, saving up to 10000 trees up to 12 steps longer). In all analyses, clades were considered robust if they had synchronously Bremer support $\geq 3$ and bootstrap support $\geq 70$.

## MOLECULAR ANALYSIS

We used both original data ( 15 species across six genera) and sequences from GenBank ( 22 species across six genera) (Table 2). All seven genera of the family are thus represented in the molecular dataset. Outgroups and ingroups were the same as in the morphological analysis (Table 2).

We selected six molecular markers: a mitochondrial ribosomal gene (16S), a mitochondrial protein-coding gene (cytochrome $c$ oxidase subunit I, COI), a nuclear ribosomal gene (18S) and three nuclear protein-coding genes: histone H 3 , sodium-potassium ATPase alphasubunit ( $\mathrm{NaK}, \sim 565 \mathrm{bps}$ ) and phosphoenolpyruvate carboxykinase (PEPCK). These markers have been widely applied in decapod phylogenetic analyses and proven to be informative at fine and coarse evolutionary scales (Bracken et al., 2009; Felder \& Robles, 2009; Robles et al., 2009; Toon et al., 2009; Bracken-Grissom et al., 2014; Ditter et al., 2020).

Total genomic DNA was extracted from the pleopods or abdomen using the Qiagen DNeasy Blood and Tissue Kit in accordance with the manufacturer's protocol. Polymerase chain reaction (PCR) amplification of the COI gene was performed with the primers: COL6/ COH6 (~ 650 bps; Schubart \& Huber, 2006; Schubart, 2009) or LCOI 1490/HCOI 2198 (~ 650 bps, Folmer et al., 1994). The mitochondrial large subunit 16 S rRNA was amplified by 16L2/16H3 primers ( $\sim 550$ bps;Schubart et al., 2002; Reuschel \& Schubart, 2006), and the nuclear small subunit 18 S rRNA was amplified by A/L, C/Y, O/B primers ( $\sim 1800 \mathrm{bps}$; Apakupakul et al., 1999). Nuclear H 3 gene fragment was amplified by H3A/H3B primers ( $\sim 330$ bps; Colgan et al., 1998), NaK with primers for-b/rev2 ( $\sim 660 \mathrm{bps}$; Tsang et al., 2008) and PEPCK ( $\sim 510 \mathrm{bps}$ ) with the 5' primer PEPCK for (Tsang et al., 2008) and newly designed for this study PEPCK acant-rev2 ( ${ }^{\prime}$ '-RCCR AAGTTGTARCCAAAGAAGGG-3') as the 3 ' primer. Polymerase chain reaction amplification reactions were performed in $25 \mu \mathrm{~L}$ containing $1 \times$ PCR buffer, $1 \mu \mathrm{~L}$ of $10 \mu \mathrm{~mol} / \mathrm{L}$ of primer pair mix, $1 \mu \mathrm{~L}$ of DNA
Table 1. Individuals used in morphological analyses. MNHN, National Museum of Natural History (Paris, France); NMNH, National Museum of Natural History, Washington, D.C., United States; ZMUK, National History Museum, Copenhagen, Denmark; IO RAN, Institute of Oceanology, Russian Academy of Scienses

| Species | Coordinates | Other information | Museum, number |
| :---: | :---: | :---: | :---: |
| Acanthephyra acutifrons | $14^{\circ} 43^{\prime} \mathrm{N} ; 45^{\circ} 02^{\prime} \mathrm{W}$ | 'Professor Logatchev' 39 cruise St 215 RT, RTAK | IO RAN 39L 215 RT №1 |
| Acanthephyra acutifrons | $8^{\circ} 53 ' \mathrm{~S}, 159^{\circ} 23^{\prime} \mathrm{E}$ | Océanie, Salomon, New Georgia sound, SALOMONBOA 3, N.O. ‘Alis', CP2783, prof. 1501-1545 m. 13.09.2007 | MNHN-IU-2016-9247 |
| Acanthephyra armata | $06^{\circ} 56^{\prime} \mathrm{N}, 52^{\circ} 35^{\prime} \mathrm{W}$ | N.O. 'Hermano Gines' GUYANE 2014 Stn CP4405 555-597 m, MNHN-convention APA-973-1, 09.08.2014 | MNHN-IU-2013-2686 |
| Acanthephyra armata | $06^{\circ} 36^{\prime} \mathrm{N}, 52^{\circ} 35^{\prime} \mathrm{W}$ | N.O. 'Hermano Gines' GUYANE 2014 Stn CP4405 555-597 m, MNHN-convention APA-973-1, 09.08.2014 | MNHN-IU-2016-9261 |
| Acanthephyra armata | $06^{\circ} 36^{\prime} \mathrm{N}, 52^{\circ} 35^{\prime} \mathrm{W}$ | N.O. 'Hermano Gines' GUYANE 2014 Stn CP4405 555-597 m, MNHN-convention APA-973-1, 09.08.2014 | MNHN-IU-2016-9261 |
| Acanthephyra carinata | $0^{\circ} 27^{\prime} \mathrm{S}, 145^{\circ} 56^{\prime} \mathrm{E}$ | Papouasie Nouvelle-Guinée: Astrolabe Bay, N.O. 'Alis', BIOPAPUA. Stn CP3717, 850-945 m. 06.10.2010 | MNHN-IU-2016-9274 |
| Acanthephyra cucculata | $16^{\circ} 04 \mathrm{~N}^{\prime} ; 46^{\circ} 41^{\prime} \mathrm{W}$ | 39 cruise RV 'Logatchev', 14-15.03.2018, 1500-0 m | IO RAN 39L233RT №65 |
| Acanthephyra curtirostris | $35^{\circ} 37^{\prime} 96{ }^{\prime} \mathrm{E}, 21^{\circ} 36{ }^{\prime} 54^{\prime} \mathrm{S}$ | Afrique, Mozambique, Canal du Mozambique, Indien, Mainbasa, 'Vizconde de Eza', CP3147. Chalut a perche, prof. 990-996 m, 12.04.2009 | MNHN-IU-2016-9280 (MNHN-Na-17146) |
| Acanthephyra curtirostris | $14^{\circ} 43^{\prime} \mathrm{N} ; 45^{\circ} 02^{\prime} \mathrm{W}$ | 39 cruise RV 'Logatchev' | IO RAN39L215RT, №1 |
| Acanthephyra fimbriata | $12^{\circ} 09^{\prime} \mathrm{N}, 122^{\circ} 14^{\prime} \mathrm{E}$ | MUSORSTOM 3, Philippines. St. CP 136, 1404 m | MNHN-IU-2018-1565 |
| Acanthephyra indica | $8^{\circ} 11^{\prime} \mathrm{N}, 79^{\circ} 03^{\prime} \mathrm{E}$ | N.O. 'Marion Dufense' SAFARI II, St. 04 CP06, 1035 m | MNHN-IU-2018-1566 |
| Acanthephyra indica | $12^{\circ} 57{ }^{\prime} \mathrm{S}, 48^{\circ} 03^{\prime} \mathrm{E}$ | Campagne MIRIKY Madagascar, 'Miriky', entre Nosy-be et Banc du Leven, Stn CP3219, 01.07.09, 906-918 m. | MNHN-IU-2009-1905 |
| Acanthephyra indica | $21^{\circ} 36{ }^{\prime} 54^{\prime} \mathrm{S}, 35^{\circ} 57^{\prime} 96{ }^{\prime} \mathrm{E}$ | Afrique, Mozambique, Canal du Mozambique, Indien. Mainbaza, N.O. 'Vizconde de Eza’, Campagne Mainbaza. Stn. CP3147, 990-996 m. 12.04.2009. | MNHN-IU-2008-10188 |
| Acanthephyra media | $13^{\circ} 05^{\prime} \mathrm{N}, 122^{\circ} 25^{\prime} \mathrm{E}$ | MUSORSTOM 2, Philippines. St. CP 42, 1580-1610 m | MNHN-IU-2018-1567 |
| Acanthephyra pelagica | $36^{\circ} 45^{\prime} \mathrm{N}, 0^{\circ} 16^{\prime} \mathrm{E}$ | DANA 1920-1922. St. 1128(1). S 200. 01.10.1921, 21.50. | ZMUK |
| Acanthephyra quadrispinosa | $29^{\circ} 39^{\prime} \mathrm{S}, 44^{\circ} 16^{\prime} \mathrm{E}$ | Expedition ATIMO VATAE. SUD MADAGASCAR, Sud Pointe Barrow. Chaultier 'Nosy Be 11', Stn. CP 3596, 986-911 m. 12.05.2010. | MNHN-IU-2010-4285 |
| Acanthephyra quadrispinosa | $29^{\circ} 39^{\prime} \mathrm{S}, 44^{\circ} 16^{\prime} \mathrm{E}$ | Expedition ATIMO VATAE. SUD MADAGASCAR, Sud Pointe Barrow. Chaultier 'Nosy Be 11', Stn. CP 3596, 986-911 m. 12.05.2010. | MNHN-IU-2010-4285 |
| Ephyrina benedicti | $2^{\circ} 39^{\prime} 5 \mathrm{~S}, 5^{\circ} 43^{\prime} 2 \mathrm{E}$ | Campagne WALDA Prélèvement 142Engin CY20, Chalut Blake Prof. 4088m. 27.07.1971. | MNHN-IU-2018-1582 |
| Ephyrina bifida | $28^{\circ} 41^{\prime} \mathrm{N}, 60^{\circ} 57^{\prime} \mathrm{W}$ | Anton Dohrn 92, St. 5780, 2000 m | ZMUK |
| Ephyrina bifida | $39^{\circ} 39,1^{\prime} \mathrm{N}, 15^{\circ} 00,2^{\prime} \mathrm{W}$ | ABYPLANE St CP15, chalutage $5320 \mathrm{~m}, 9.06 .1981$, 8h07-9H30 | MNHN-IU-2018-1580 |
| Ephyrina figueirai | $05^{\circ} 27{ }^{\prime} \mathrm{S}, 146^{\circ} 09^{\prime} \mathrm{E}$ | Bismarck Sea: Basamuk Bay, N.O.'Alis', Expedition PAPUA NIUGINI, Stn CP4082, 800-1065 m, 26/12/2012 | MNHN-IU-2013-8725 |
| Ephyrina ombango | $10^{\circ} 23,17^{\prime} \mathrm{N}, 46^{\circ} 45,34^{\prime} \mathrm{W}$ | DEMERABY, CP07, chalutage 4850 m .20 .09 .80 | MNHN-IU-2018-1579 |
| Ephyrina ombango | $9^{\circ} 18{ }^{\prime} \mathrm{S}, 11^{\circ} 10^{\prime} \mathrm{E}$ | 'Ombango', C14, St. 325 ,midwater traul, 0-725 m, 02.03.1961, 23h00-23h15 | MNHN-IU-2014-11098 |
| Hymenodora glacialis | $02^{\circ} 03{ }^{\prime} \mathrm{S}, 118^{\circ} 45^{\prime} \mathrm{E}$ | Indonésie, CORINDON -Makassar. St CH286, 1710-1730 m | Na 10655 |

Table 1. Continued

| Species | Coordinates | Other information | Museum, number |
| :---: | :---: | :---: | :---: |
| Hymenodora glacialis | $73^{\circ} 28^{\prime} \mathrm{N}, 10^{\circ} 07^{\prime} \mathrm{W}$ | Mer de Norvège, Campagne NORBI, N.O. ‘Jean Charcot’, Stn CP16, 2937 m, 07.08.1975 | MNHN-IU-2008-16833 |
| Hymenodora gracilis | $37^{\circ} 39^{\prime} \mathrm{S}, 77^{\circ} 26^{\prime} \mathrm{E}$ | Ile Amsterdam, Campagne Jasus (MD 50), N.O. ‘Marion Dufresne’, Stn CP193, 2800-3075 m. 27.06.1986 | MNHN-IU-2008-16839 |
| Janicella spinicauda | $1^{\circ} 28^{\prime} \mathrm{S}, 48^{\circ} 06^{\prime} \mathrm{E}$ | ROV 'Vityaz', 17th cruise, St. 2604,13.11.88, 670-690 m | ZMUK |
| Janicella spinicauda | $8^{\circ} 44{ }^{\prime} \mathrm{S}, 43^{\circ} 54^{\prime} \mathrm{E}$ | Dana Expedition, St. 3939-1, 23.12.1929, 500 meter wire | ZMUK |
| Kemphyra corallina | $37^{\circ} 54{ }^{\prime} \mathrm{S}, 77^{\circ} 22^{\prime} \mathrm{E}$ | Iles St Paul et Amsterdam, 'Marion Dufresne’ Cne MD Jasus Stn CP 56. $2280-2310 \mathrm{~m} .14 .07 .1986$. 20h02-22H31 | MNHN-IU-2018-1581 |
| Kemphyra corallina | $33^{\circ} 59{ }^{\prime} \mathrm{S}, 43^{\circ} 55^{\prime} \mathrm{E}$ | Indian Ocean: Walters shoal, Plaine Sud. N.O. 'Marion Dufresne', Campagne MD208(Walters Shoal). Stn CP4915б, 1865-2058 m, 12.05.2017 | MNHN-IU-2016-9402 |
| Meningodora compsa | $16^{\circ} 16^{\prime} \mathrm{N}, 22^{\circ} 16^{\prime} \mathrm{W}$ | Service de l'élevage du Sénégal. M.W.T. 0-1000 m. 16.01.1959 | MNHN-IU-2018-1577 |
| Meningodora longiscula | $9^{\circ} 55^{\prime} \mathrm{N}, 142^{\circ} 00^{\prime} \mathrm{E}$ | Nouvelle-Calédonie, Campagne Caride V. Stn 15, 1000 m., 12.09.1969 | MNHN-IU-2011-5635 |
| Meningodora mollis | $34^{\circ} 06^{\prime} \mathrm{N}, 17^{\circ} 06^{\prime} \mathrm{W}$ | North Atlantic, Campagne Abyplane, N.O. 'Cryos',Stn. CP11, 4270 m, 30.05.1981 | MNHN-IU-2011-5640 |
| Meningodora vesca | $39^{\circ} 59^{\prime} \mathrm{N}, 15^{\circ} 00^{\prime} \mathrm{W}$ | North Atlantic, Campagne ABYPLANE, N.O. 'Cryos', STN CP15, 5320 m | MNHN-IU-2011-5634 |
| Notostomus auriculatus | $33^{\circ} 59{ }^{\prime} \mathrm{S}, 43^{\circ} 55^{\prime} \mathrm{E}$ | Indian Ocean: Walters shoal, Plaine Sud. N.O. 'Marion Dufresne', Campagne MD208(Walters Shoal). Stn CP4915, 1865-2058 m, 12.05.2017 | MNHN-IU-2016-9404 |
| Notostomus elegans |  | 37 cruise RV Logatchev, St 156 TS | IO RAN |
| Notostomus gibbosus | $21^{\circ} 46{ }^{\prime} \mathrm{S}, 36^{\circ} 35^{\prime} \mathrm{E}$ | Afrique, Mozambique, Canal du Mozambique, Indien, Campagne Mainbasa, N.O. 'Vizconde de Eza', Stn CC3156, 1810-1820, 14.04.2009 | MNHN-IU-2008-10189 |
| Notostomus japonicus | $54^{\circ} 11^{\prime} 19^{\prime} \mathrm{N}, 166^{\circ} 45^{\prime} 37{ }^{\prime} \mathrm{W}$ | North Pacific Ocean, Bering Sea, Alaska. 19.05.2001. NOAA Expedition 2001, R/V Vesteraalen. Depth 628 m. | USNM 1164652 |
| Notostomus murrayi | $29^{\circ} 50,9^{\prime} \mathrm{S}, 48^{\circ} 35,5^{\prime} \mathrm{E}$ | SUD/SUD-EST MADAGASCAR, Campagne Safari I(MD20), N.O. 'Marion Dufresne, St.18, CP10, 04.09.1979, 7:36-8:20, 3668-3800 m | MNHN-IU-2011-5686 |
| Notostomus murrayi | $31^{\circ} 12.0 \mathrm{~S}, 39^{\circ} 18.0 \mathrm{~W}$ | ASV46, St 2719, 800-200 m, 25.10.2018 | IO RAN 9-D2 |
| Notostomus robustus | $37^{\circ} 08^{\prime} 24^{\prime} \mathrm{N}, 074^{\circ} 17^{\prime} 42^{\prime} \mathrm{W}$ | North Atlantic ocean, United States, North American slope, Off Virginia. Trawl. Depth 2933 ms. Gilliss R/V GI-75-08-35. 14.09.1973 | USNM 222196 |
| Oplophorus gracilirostris | $25^{\circ} 11^{\prime} \mathrm{N}, 122^{\circ} 35^{\prime} \mathrm{E}$ | Dana Expedition, St. 3722-3, 300 meter wire | ZMUK |
| Oplophorus gracilirostris | $20^{\circ} 08^{\prime} \mathrm{N}, 82^{\circ} 59^{\prime} \mathrm{W}$ | Dana Expedition, St. 1218, 800 meter wire | ZMUK |
| Oplophorus gracilirostris | $12^{\circ} 30^{\prime} \mathrm{S}, 48^{\circ} 16^{\prime} \mathrm{E}$ | ROV 'Vityaz', 17th cruise, St. 2597, 12.11.88, 360-555 meter wire. | ZMUK |
| Oplophorus gracilirostris | $22^{\circ} 06^{\prime} \mathrm{N}, 84^{\circ} 58^{\prime} \mathrm{W}$ | Dana Expedition, St. 1223, 500 metre wire | ZMUK |
| Pasiphaea sivado | $35^{\circ} 47^{\prime} \mathrm{N}, 05^{\circ} 17^{\prime} \mathrm{W}$ | Detroit de Gibraltar, N.O. ‘Cryos', BALGIM, St. CP150, 280-300 m, 18.06.1984 | MNHN-IU-2018-1611 |
| Penaeus monodon | No data | Andrea 1877, Singapore | ZMUK |
| Solenocera membranaceum | $7^{\circ} 55^{\prime} \mathrm{S}, 12^{\circ} 38^{\prime} \mathrm{E}$ | Atlantide Exp. West Africa 1945-1946. St. 135, depth 460-235 m, Gear: ET, 17.03.1946, 13:40-15:40. | ZMUK |
| $\underline{\text { Systellaspis debilis }}$ | $2^{\circ} 15^{\prime} \mathrm{S}, 98^{\circ} 55,5^{\prime} \mathrm{E}$ | Dana Expedition, St. 3817-3, 11.09.1929, 300 metre wire | ZMUK |

Table 1. Continued

| Species | Coordinates | Other information | Museum, number |
| :--- | :--- | :--- | :--- |
| Systellaspis debilis | $17^{\circ} 10^{\prime} \mathrm{N}, 46^{\circ} 25 \mathrm{~W}$ | ROV ‘Professor Logachev', 37 cruise, St. 99 IKMT | IO RAN 37L99 IKMT |
| Systellaspis debilis | $21^{\circ} 57^{\prime} \mathrm{N} ; 22^{\circ} 58^{\prime} \mathrm{W}$ | Dana Expedition, St. 1157, E 300, 27.10.1921 | ZMUK |
| Systellaspis debilis | $31^{\circ} 12.0 \mathrm{~S}, 39^{\circ} 18.0 \mathrm{~W}$ | R.V. Akademik Sergey Vavilov, 46 cruise, St. 2719, 25.10.2018, 200-800 m | IO RAN 13-D1 |

template, $0.2 \mathrm{mmol} / \mathrm{L}$ of each dNTP and 0.5 units of Taq polymerase. The thermal profile used an initial denaturation for 3 min at $95{ }^{\circ} \mathrm{C}$ followed by 35-40 cycles of 20 s at $94{ }^{\circ} \mathrm{C}, 30 \mathrm{~s}$ at $45-57{ }^{\circ} \mathrm{C}$ depending on primer pair, 1 min at $72^{\circ} \mathrm{C}$ and a final extension of 7 min at $72^{\circ} \mathrm{C}$. Polymerase chain reaction products were purified using the PCR Purification Kit protocol (Promega) and sequenced in both directions using BigDye Terminator v.3.1 (Applied Biosystems). Each sequencing reaction mixture, including $0.5 \mu \mathrm{~L}$ of BigDye Terminator v.3.1, $0.8 \mu \mathrm{~L}$ of $1 \mu \mathrm{~mol} / \mathrm{L}$ primer and $1-2 \mu \mathrm{~L}$ of purified PCR template, was run for 30 cycles of $96^{\circ} \mathrm{C}(10 \mathrm{~s}), 50^{\circ} \mathrm{C}(5 \mathrm{~s})$ and $60^{\circ} \mathrm{C}$ ( 4 min ). Sequences were purified by ethanol precipitation to remove unincorporated primers and dyes. Products were re-suspended in $14 \mu \mathrm{~L}$ formamide and electrophoresed in ABI Prism-3500 sequencer (Applied Biosystems). The nucleotide sequences were cleaned and assembled using CodonCode Aligner v.7.1.1. Protein-coding sequences ( $C O I, H 3, \mathrm{NaK}$ and PEPCK) were checked for indels and stop codons to prevent the inclusion of pseudogenes. All sequences were then compared to genes reported in GenBank using BLAST (National Center for Biotechnology Information, NCBI) to check for potential contamination.

For each gene-fragment, the sequences were aligned using MUSCLE (Edgar, 2004) implemented in MEGA v.X (Kumar et al., 2018), and the alignment accuracy was adjusted by eye. Missing data were designated with a '?' for any incomplete sequences. All obtained sequences were submitted to the NCBI GenBank database (Table 2).

To assess phylogenetic relationships between species, Bayesian inference (BI) and maximum likelihood (ML) analyses were run. The BI analysis was conducted in MrBayes v.3.2.6 (Ronquist et al., 2012) for the concatenated dataset of all genes. The combined dataset was partitioned and analysed using models selected by PartitionFinder2 (Lanfear et al., 2016). Akaike information criterion (AICc modification for small sample size) metric implemented in PartitionFinder2 was used to obtain the optimal partitioning scheme. Two independent runs, each consisting of four chains, were executed for this analysis. A total of 10000000 generations were performed for the combined dataset, with sampling every 1000 generations, and the first $25 \%$ trees (i.e. 2500 trees for combined dataset) were discarded as 'burn-in'. A $1 \%$ average standard deviation of split frequencies was reached after about 1.1 million generations.

The maximum likelihood (ML) analysis was run in RAxML GUI v.2.0 (Stamatakis, 2014; Edler et al., 2020), and the GTR+G model was used. Bootstrap resampling with 1000 replicates was run using the thorough bootstrap procedure to assign support to


Figure 2. Grouping of morphological characters (schematic white lines) in Ephyrina, Meningodora and Notostomus exemplified by Notostomus elegans.
branches in the ML tree. Final ML tree was generated using the partitioned dataset of all concatenated genes.

We considered the clades statistically supported if they had a synchronous support of posterior probabilities $\geq 0.9$ in the BI analysis and bootstrap value $\geq 70 \%$ in the ML analysis.

## RESULTS

## MORPHOLOGICAL ANALYSES

Analysis 1 with Janicella spinicauda as outgroup retrieved a single most-parsimonious (MP) tree (Fig. 3A; Supporting Information, Appendix S3) with a score of $109(\mathrm{Ci}=88, \mathrm{Ri}=96)$. The tree shows that Hymenodora is a sister-clade to the rest of the genera, the latter clade includes two sister-clades: Ephyrina and Heterogenys + Kemphyra + Acanthephyra + Notostomus + Meningodora. There is also a wellsupported clade Meningodora + Notostomus within which both genera are robust sister-clades. Acanthephyra shows polytomy.
Analysis 2 with Oplophorus gracilirostris as outgroup retrieved a single MP tree (Fig. 3B; Supporting Information, Appendix S3) with a score of $116(\mathrm{Ci}=82$, Ri = 95). Analysis 3 with Systellaspis pellucida as outgroup also retrieved a single MP tree (Fig. 3C; Supporting Information, Appendix S3) with a score of $104(\mathrm{Ci}=92, \mathrm{Ri}=98)$. Both trees are similar in topology to each other and to the tree retrieved in Analysis 1, with the same set of statistically supported clades.

## MOLECULAR ANALYSES

We successfully obtained 84 sequences across six gene fragments for 15 out of 21 species from the genera Ephyrina, Meningodora and Notostomus. In order to retrieve phylogenetic reconstructions, we also included all species of Acanthephyridae from GenBank with at least two selected gene markers. Prior to analyses, all sequences from GenBank were checked for contamination or possible misidentification using BLAST search and preliminary phylogenetic reconstruction with each gene separately. A total of 35 species from seven genera of Acanthephyridae and three genera of Oplophoridae were thus put in the data matrix. The concatenated six-marker dataset comprised 4525 bp . Results from PartitionFinder2 recommended a 12 -partition scheme by gene and codon (H3, COI, NaK, PEPCK), which was used in the final analyses. Substitution models for each partition are listed in Table 3.

Molecular analysis retrieved Bayesian and ML trees, which are similar to each other in topology but significantly differ in support of two major clades (Fig. 4).

On the BI tree, Hymenodora is a sister-clade to the rest of the genera, the latter clade includes two sister-clades: Ephyrina and 'Heterogenys + Kemphyra + Acanthephyra + Notostomus + Meningodora'. There is also a well-supported clade Meningodora + Notostomus, within which Notostomus is monophyletic and Meningodora shows polytomy. Although the BI tree shows polytomy of Acanthephyra, some clades
Table 2. Individuals used in phylogenetic reconstruction with localities, voucher numbers, and GenBank accession numbers. New sequences obtained are in bold,
' N ' - sequences not acquired ' N ' - sequences not acquired

| Species | Voucher No | Locality/year | GenBank accession numbers |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | COI | 16S | 18S | H3 | NaK | PEPCK | References |
| Ephyrina |  |  |  |  |  |  |  |  |  |
| Ephyrina benedicti | ACP1 | Central Atlantic / 2018 | MW043002 | MW043446 | MW043461 | N | MW052302 | MW052315 | This study |
| Ephyrina benedicti | HBG6799 | Gulf of Mexico / | MH572612 | MH542965 | N | N | N | N | Published in GenBank, Wilkins \& BrackenGrissom, 2020 |
| Ephyrina bifida | HBG160 | North Atlantic | KP076186 | N | KP075779 | N | KP076039 | N | Wong et al., 2015 |
| Ephyrina figueirai | MNHN-IU-2013-8725 <br> / ACP4 | Papua New Guinea / 2012 | MW043003 | MW043447 | MW043462 | MW052288 | MW052303 | MW052316 | This study |
| E. figueirai spinicauda | HBG933 | Taiwan | KP076189 | KP075911 | KP075800 | KP076105 | KP076038 | N | Wong et al., 2015 |
| Ephyrina ombango | ACP6 | Central Atlantic / 2018 | MW043004 | MW043448 | MW043463 | MW052289 | MW052304 | MW052317 | This study |
| Ephyrina ombango | HBG1230 | Gulf of Mexico | KP076188 | KP075914 | KP075802 | KP076107 | N | N | Wong et al., 2015 |
| Heterogenys |  |  |  |  |  |  |  |  |  |
| Heterogenys microphthalma | HBG937 | Taiwan | KP076183 | KP075898 | KP075787 | KP076124 | KP076035 | N | Wong et al., 2015 |
| Hymenodora |  |  |  |  |  |  |  |  |  |
| Hymenodora glacialis |  | Arctic Ocean / 1975 |  | GQ131896 | GQ131915 | N | N | N | Chan et al., 2010 |
| Hymenodora gracilis | ACP12 | Central Atlantic / 2018 | MW043005 | MW043449 | MW043464 | MW052290 | MW052305 | MW052318 | This study |
| Kemphyra |  |  |  |  |  |  |  |  |  |
| Kemphyra corallina | MNHN-IU- <br> 2016-9402/ACP46 | South-West Indian Ocean / 2017 | MW043006 | MW043450 | MW043465 | MW052291 | N | N | This study |
| Meningodora |  |  |  |  |  |  |  |  |  |
| Meningodora compsa | $\begin{gathered} \text { HBG1241 and } \\ \text { HBG7260 } \end{gathered}$ | Gulf of Mexico | Unpublushed, H. BrackenGrissom | KP075907 | KP075791 | KP076114 | N | N | Wong et al., 2015 |
| Meningodora longisulca | ACP19 | Central Atlantic / 2016 | MW043007 | MW043451 | MW043466 | MW052292 | MW052306 | MW052319 | This study |
| Meningodora miccyla | ACP17 | $\begin{aligned} & \text { Central Atlantic / } \\ & 2018 \end{aligned}$ | MW043008 | MW043452 | MW043467 | MW052293 | MW052307 | MW052320 | This study |
| Meningodora mollis | HBG901 | Gulf of Mexico | KP076192 | KP075910 | KP075783 | KP076115 | KP076033 | N | Wong et al., 2015 |
| Meningodora mollis | HBG1170 | Spain | KP076193 | N | KP075813 | KP076116 | KP076034 | N | Wong et al., 2015 |
| Meningodora vesca | ACP32 | $\begin{aligned} & \text { Central Atlantic / } \\ & 2018 \\ & \hline \end{aligned}$ | MW043009 | MW043453 | MW043468 | MW052294 | MW052308 | MW052321 | This study |

Table 2. Continued

| Species | Voucher No | Locality/year | GenBank accession numbers |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | COI | 16S | 18 S | H3 | NaK | PEPCK | References |
| Meningodora vesca | HBG4178 | Gulf of Mexico | MF197254 | MF197197 | N | N | N | N | Published in GenBank, Wilkins \& Bracken-Grissom, 2017 |
| Notostomus |  |  |  |  |  |  |  |  |  |
| Notostomus auriculatus | MNHN-IU- <br> 2016-9404/ACP20 | South-West Indian Ocean / 2017 | MW043010 | MW043454 | MW043469 | MW052295 | MW052309 | MW052322 | This study |
| Notostomus elegans | ACP23 | $\begin{aligned} & \text { Central Atlantic / } \\ & 2016 \end{aligned}$ | MW043011 | MW043455 | MW043470 | MW052296 | MW052310 | MW052323 | This study |
| Notostomus elegans | HBG1232 and HBG1169 | Gulf of Mexico | KP076194 | KP075900 | KP075803 | KP076119 | KP076034 | N | Wong et al., 2015 |
| Notostomus gibbosus | ACP35 | Central Atlantic / 2015 | MW043012 | MW043456 | MW043471 | MW052297 | MW052311 | MW052324 | This study |
| Notostomus gibbosus | $\begin{gathered} \text { HBG903A and } \\ \text { HBG4220 } \end{gathered}$ | Gulf of Mexico | MH572685 | KP075905 | KP075795 | KP076120 | N | N | Wong et al., 2015; Published in GenBank, Wilkins \& BrackenGrissom, 2020 |
| Notostomus japonicus | USNM 1164652/ ACP25 | Bering Sea / 2001 | DQ882094 | MW043457 | MW043472 | MW052298 | N | N | This study; Published in GenBank, Costa et al., 2018 |
| Notostomus murrayi | ACP30 | South Atlantic / $2018$ | MW043013 | MW043458 | MW043473 | MW052299 | MW052312 | MW052325 | This study |
| Notostomus robustus | ACP24 | Central Atlantic / 2016 | MW043014 | MW043459 | MW043474 | MW052300 | MW052313 | MW052326 | This study |
| Acanthephyra |  |  |  |  |  |  |  |  |  |
| Acanthephyra acutifrons | HBG1254 | Gulf of Mexico | KP076167 | KP075874 | KP075817 | KP076084 | KP076037 | N | Wong et al., 2015 |
| Acanthephyra armata | MNHN-IU-2011-3081 | Papua New Guine <br> / 2010 | KP759353 | KP725471 | KP725668 | KP726043 | N | N | Aznar-Cormano et al., 2015 |
| Acanthephyra carinata | BSM110 | Philippines | KP076184 | KP075896 | KP075798 | KP076093 | N | N | Wong et al., 2015 |
| Acanthephyra cucullata | HBG925 | Taiwan | KP076160 | KP075893 | KP075809 | KP076110 | N | N | Wong et al., 2015 |
| Acanthephyra curtirostris | HBG1407 | Gulf of Mexico | KP076161 | KP075889 | KP075807 | KP076088 | KP076029 | N | Wong et al., 2015 |
| Acanthephyra eximia | MNHN-IU-2008-16779/ <br> NTOU M00709 | Pacific Ocean: Southern Archipelago / 2002 | KP759360 | KP725477 | KP725675 | KP726049 | EU427181 | EU427250 | Aznar-Cormano et al., 2015; Tsang et al., 2008 |
| Acanthephyra fimbriata | HBG927 | Philippines | KP076185 | KP075895 | KP075788 | KP076092 | N | N | Wong et al., 2015 |
| Acanthephyra indica | MNHN-IU-2008-10188/ ACP16 | Mozambique Channel / 2009 | MW043001 | MW043445 | MW043460 | MW052287 | MW052301 | MW052314 | This study |
| Acanthephyra media | HBG930 | Philippines | KP076166 | KP075892 | KP075805 | KP076086 | N | N | Wong et al., 2015 |

Table 2. Continued

| Species | Voucher No | Locality/year | GenBank accession numbers |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | COI | 16 S | 18S | H3 | NaK | PEPCK | References |
| Acanthephyra pelagica | HBG153 | North Atlantic | KP076182 | KP075880 | KP075789 | KP076100 | KP076027 | N | Wong et al., 2015 |
| Acanthephyra purpurea | HBG899A | Gulf of Mexico | KP076170 | KP075882 | KP075782 | KP076095 | KP076023 | N | Wong et al., 2015 |
| Acanthephyra quadrispinosa | HBG931 | Taiwan | KP076178 | KP075886 | KP075821 | KP076099 | KP076025 | N | Wong et al., 2015 |
| Outgroups |  |  |  |  |  |  |  |  |  |
| Janicella spinicauda | HBG7002, HBG905 and MNHN-IU-2014-18783/ Op11 | Gulf of Mexico, Papua New Guinea | MH572546 | KP075932 | MH100869 | MH107256 | N | N | Wong et al., 2015; Lunina et al., 2019b; Published in GenBank, Wilkins \& BrackenGrissom, 2020 |
| Oplophorus gracilirostris | HBG909A and HBG904A | Gulf of Mexico | KP076150 | KP075920 | KP075847 | KP076072 | KP076045 | N | Wong et al., 2015 |
| Systellaspis pellucida | $\begin{aligned} & \text { HBG944 and } \\ & \text { NTOU:M01001 } \end{aligned}$ | Taiwan | KP076147 | KP075924 | JF346250 | JF346319 | JF346355 | JF346391 | Li et al., 2011; <br> Wong et al., 2015 |

[^1]within the genus are robust: 'Acanthephyra armata', 'Acanthephyra media' and 'Acanthephyra purpurea' species groups (Fig. 4). The ML tree shows lesser support (slightly below accepted 70) of the two deepest nodes indicated by arrows in Fig. 4. In other respects, ML and BI trees are similar.

## MORPHOLOGICAL SYNAPOMORPHIES

MP trees are similar in Analyses 1-3 and we, therefore, mapped morphological synapomorphies in a single picture Fig. 5 for all analyses. In addition to robust clades shown in Fig. 3, all morphological analyses retrieved three minor clades within Acanthephyra, which do not receive statistical support but are identical to species groups retrieved in molecular analyses: 'Acanthephyra armat', 'Acanthephyra media' and 'Acanthephyra purpurea' (Fig. 5). Unlike its position on the molecular trees, the first species group was combined with Kemphyra.

The clade 'Acanthephyridae without Hymenodora' is supported by the presence of the postorbital dorsal teeth on the rostrum (character 5, see Supporting Information, Appendix S1), a submarginal papilla and a lamina on the second maxilla (63), three-segmented endopod on the first maxilliped $(65,66)$ and a reduced dactyl of the fifth pereopods attached transversely to the propodus (91, 92). Ephyrina is supported by a rostrum shaped as an unarmed crest $(0,4)$, a postorbital ridge from the orbit to the posterior margin of the carapace and a blunt ridge ventral to the postorbital ridge ( 20,23 ), a mandible unarmed along the distal margin (61), greatly compressed and expanded meri and ischia on all pereopods ( $75,76,81-86,88,89$ ). The clade Meningodora + Notostomus is supported by a net of sharp lateral carinae along the whole carapace length ( 7,8 ), including a sharp postorbital carina from the orbit to the posterior margin of the carapace (18) and a sharp oblique transverse carina ventral of the postorbital carina (24). In addition, Meningodora + Notostomus have a mandible unarmed along the distal margin (61), similar to that in Ephyrina. Meningodora is supported by a reduction of the dorsal carina on the second pleonic segment (35) and a blunt, indistinct carina on the third pleonic segment (38, 39). Notostomus is supported by a long branchiostegal carina, which is $0.7-1.0$ of the carapace length (13), a supraorbital carina extending from the rostrum to the postorbital region (15), an additional lateral carina on posterior part of the carapace parallel to the postorbital carina (21) and a strong mesial teeth on the posterior margin of the third and fourth pleonic segments $(44,47)$. All these synapomorphies are stable within clades, except the presence of the postorbital dorsal teeth on the rostrum in the clade 'Acanthephyridae without Hymenodora': the teeth posteriorly disappear


Figure 3. Morphological MP trees with Janicella spinicauda (A), Oplophorus gracilirostris (B) and Systellaspis pellucida (C) as the outgroups. Different colours indicate different genera. Only clades supported by both Bremer values (in bold, below branches) and bootstrap values (blue, above branches) are shown.

Table 3. Partitioning scheme and best models selected by PartitionFinder2

| Partition | Best Model |
| :--- | :--- |
| 16 S | GTR+I+G |
| $18 \mathrm{~S}, 1^{\text {st }}$ codon of $H 3$ | TRN+G |
| $2^{\text {nd }}$ codon of H 3 | K80 +I |
| $3^{\text {rd }}$ codon of H 3 | GTR+G |
| $1^{\text {st }}$ codon of $C O I$ | SYM $+\mathrm{I}+\mathrm{G}$ |
| $2^{\text {nd }}$ codon of $C O I$ | TVM+G |
| $3^{\text {rd }}$ codon of $C O I$ | GTR+G |
| $1^{\text {st }}$ codon of $\mathrm{NaK}, 1^{\text {st }}$ codon of | TVM $+\mathrm{I}+\mathrm{G}$ |
| $\quad$ PEPCK |  |
| $2^{\text {nd }}$ codon of NaK | GTR+I |
| $3^{\text {rd }}$ codon of NaK | HKY+G |
| $2^{\text {nd }}$ codon of PEPCK | TVMEF+I |
| $3^{\text {rd }}$ codon of PEPCK | TVM+G |

in Ephyrina. The mandible, unarmed along the distal margin, is a homoplasy found in the Ephyrina and Meningodora + Notostomus clades.

We grouped morphological synapomorphies into four types (Fig. 2) and calculated the contribution of each type in the support of major clades (Table 4 based on Fig. 5 and Supporting Information, Appendix 4). Average contribution of each type of synapomorphies ranged between $14 \%$ and 37\% (last line in Table 4), but supporting synapomorphies were unevenly distributed in the analysed clades. The support of the Ephyrina clade was mainly provided by
synapomorphies linked to the pereopods (oar-like meri and ischia): their contribution was exceptionally high ( $67 \%$ vs. $14 \%$ on average). Meanwhile, the clades Meningodora + Notostomus, Meningodora and Notostomus were mainly supported by synapomorphies linked to strengthening of the carapace, pleon and telson (carinae and teeth), their combined contribution was greater than on average ( $80-100 \%$ vs. 32-37\%).

## DISCUSSION

## Ephyrina, Meningodora and Notostomus on

 PHYLOGENETIC TREES AND THEIR STATUSThe most comprehensive analysis of Acanthephyridae hitherto done (Wong et al., 2015) encompassed seven species of the target genera and 14 other species of the family: Hymenodora (two species), Ephyrina (three), Meningodora (two), Notostomus (two), Heterogenys (one) and Acanthephyra (11); the analysis was based on seven gene markers and did not include morphological evidence. Here we use six gene markers and significantly extended the number of analysed species of the target group (to 15) and the rest of Acanthephyridae (to 17, including a representative of the genus Kemphyra not sequenced before this study). In order to improve the power of the analyses, we also included morphological evidence.

Overall, our study makes the phylogenetic results shown in Fig. 2 by Wong et al. (2015) statistically significant. First, the major clades 'Acanthephyridae


Figure 4. Molecular BI tree with supported clades, the horizontal scale bars mark the number of expected substitutions per site. Statistical support indicated as Bayesian posterior probabilities (black, above branches) and ML bootstrap analysis (blue, below branches). Different colours indicate different genera. Arrows indicate deep nodes perfectly resolved on the BI tree and insufficiently resolved on the ML tree.
without Hymenodora' and 'Heterogenys + Kemphyra + Acanthephyra + Notostomus + Meningodora' are not robust on the BI tree in Wong et al. (2015) but gain support here. Both clades have great support on the BI molecular tree ( 0.98 and 1, Fig. 4) and high Bremer and bootstrap support on all morphological trees in Analyses 1-3 (Fig. 3). Having this in mind, we consider both deepest nodes on the tree (arrows in Fig. 4) resolved, although bootstrap values on the ML molecular tree are below the generally accepted 70 ( 68 and 63). Our results thus confirm that Hymenodora is a sister-clade to the rest of Acanthephyridae and that Ephyrina is a sister-clade to 'Heterogenys + Kemphyra + Acanthephyra + Notostomus + Meningodora'.

As in Fig. 2 by Wong et al. (2015), the clade Notostomus + Meningodora is robust; this clade gains greater support on both of our molecular trees and is robust on all morphological trees. As in the previous studies, Notostomus is monophyletic in all trees and Meningodora is not resolved on both molecular trees. However, Meningodora is monophyletic and gains bootstrap and Bremer support on our morphological trees, which shows the resolving power of morphological methods in this particular case. The
current phylogenetic status of Meningodora match the status of Systellaspis (Lunina et al., 2019b) from the sister-clade Oplophoridae: both genera are robust on the morphological trees and do not receive support on the molecular trees. As in Lunina et al. (2019b), we maintained a conservative approach and did not change the taxonomic status of Meningodora. We hope to solve the problem of a possible polyphyly of Meningodora and Systellaspis after completing a revision of the whole superfamily Oplophoroidea. Ephyrina and Notostomus are monophyletic genera on all trees.

Molecular methods, in turn, show the resolving power in a retrieving of statistical support for three species group clades within Acanthephyra (Fig. 4), which do occur (Fig. 5) but are not robust on the morphological trees (Fig. 3). Future use of the combination of morphological and molecular methods based on richer datasets and focused on Acanthephyra is needed to justify the taxonomic status of these species groups.

We conclude that the target group Ephyrina-Notostomus-Meningodora is not monophyletic on all phylogenetic trees and the unusual mandible and strengthened carapace observed in these genera thus


Figure 5. Synapomorphies on identical morphological MP trees with Janicella spinicauda, Oplophorus gracilirostris and Systellaspis pellucida as the outgroup. Different colours indicate different genera. Synapomorphies retrieved in analyses 1-3 are similar and mapped in Supporting Information, Appendix S4. Character coding see in Supporting Information, Appendix S1.

Table 4. Contribution (\%) of different groups of synapomorphies supporting major clades of Acanthephyridae, results of morphological analyses $1-3$ combined

| Clades | Synapomorphies and their numerical order in parenthesis (see Appendix S1) |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
|  | Carapace (0-32) | Pleon + telson (33-55) | Mouthparts (58-74) | Pereopods (75-92) |
| Hymenodora | 83 | 0 | 17 | 0 |
| Acanthephyridae without | 17 | 0 | 50 | 33 |
| $\quad$ Hymenodora |  |  |  |  |
| Ephyrina | 0 | 7 | 67 |  |
| Acanthephyridae without | 0 | 71 | 29 | 0 |
| $\quad$ Hymenodora and Ephyrina |  | 0 | 20 | 0 |
| Meningodora + Notostomus | 80 | 100 | 0 | 0 |
| Meningodora | 0 | 50 | $\mathbf{0}$ | $\mathbf{1 4}$ |
| Notostomus | $\mathbf{3 2}$ | $\mathbf{1 7}$ |  |  |
| Average for clades | $\mathbf{3 7}$ |  |  |  |

exemplify parallel evolution. We reject the hypothesis of the monophyly of the target group.

## MORPHOLOGICAL TRAITS IN EPHYRINA, MENINGODORA AND NOTOSTOMUS

Evolutionary traits in the clades Ephyrina and Meningodora + Notostomus are different. Ephyrina is mostly supported by synapomorphies linked to the pereopods: contribution of these characters is
nearly five times higher than on average in the major clades of Oplophoroidea (Table 4). All meri and ischia in Ephyrina are greatly compressed, expanded and resemble oars (Fig. 1E) adapted to locomotory function. Unlike the usual spear-like and serrate shrimp rostra (Fig. 1C, 1D), the rostrum in Ephyrina is a smooth, wide lamina (Fig. 1E), possibly adapted to stabilize forward motion, as does a stempost of a cruiser. The forward motion requires strengthening of the carapace but not in the form of sharp carinae, which may cause
turbulent flow along carapace and pleon. Ephyrina has smooth ridges on the carapace extending from the orbit to the posterior margin, a blunt ridge ventral to the postorbital ridge (Fig. 1E) and no more ridges, carinae or teeth on the pleon. Ephyrina benedicti Smith, 1885 and E. bifida Stephensen, 1923 have 'dorsomedial teeth' on the third pleonic segment but these teeth are soft protuberances flattened dorsoventrally, adjacent to the pleon and do not prevent an active forward motion.

Our data suggest that the set of characters above evolved as a single morphological unit and contributed to the evolutionary success of the genus. Once evolved, this set of adaptations remained conservative and the six known species of Ephyrina are similar externally. Morphological traits within the genus mainly encompass development and the shape of dorsal protuberances on the third abdominal segment in two species (entire in $E$. benedicti and bifid in $E$. bifida) and spination of the telson: position of spines (dorsolateral in E. bifida, E. childressi Chace, 1986 and E. hoskynii Wood-Mason, 1891 or marginal in E. benedicti, E. figueirai Crosnier \& Forest, 1973 and E. ombango Crosnier \& Forest, 1973), number of spines and additional rows of spines (E. figueirai).

The Meningodora + Notostomus clade is mostly supported by synapomorphies linked to the strengthening of the carapace; contribution of these characters is two to three times higher than on average in the major clades of Oplophoroidea (Table 4). The strengthening is provided by means of sharp lateral carinae along the whole carapace length (a postorbital carina from the orbit to the posterior margin, an oblique transverse carina ventral of the postorbital carina; Fig. 1C, 1D), which are coupled with a thin, half-membranous integument. The strengthening has probably evolved to keep the body firm and rigid. Strong carinae on the carapace are likely analogous to stiffening members in ships and serve to reduce vibrations and flexing during fast movement (especially escape flips). Sharp carinae, which are absent in shrimps with a firm carapace, such as Oplophoridae and most Acanthephyridae, become indispensable for Meningodora and Notostomus having a thin and half-membranous integument.

The main evolutionary traits within the clade are linked to a further strengthening of the carapace and the pleon. Notostomus is supported by an impressive set of such synapomorphies (Fig. 1C) (the last two may also be defensive):

- A sharp branchiostegal carina along 0.7-1.0 of carapace length.
- An additional lateral carina on the posterior part, parallel to the postorbital carina.
- A supraorbital carina, extending from the rostrum to the postorbital region.
- A denticulate dorsal carina on the carapace.
- Strong and firm posteromesial teeth on the third to fifth abdominal somites.

The basic morphological trait within Notostomus is also linked to further strengthening of the carapace: N. auriculatus Barnard, 1950, N. crosnieri Macpherson, 1984, N. elegans A. Milne-Edwards, 1881, $N$. japonicus Spence Bate, 1888 and N. murrayi Spence Bate, 1888 have an additional lateral carina along the entire carapace length between the branchiostegal carina and the ventral margin of the carapace, the three latter species also have a transverse oblique carina extending dorsally from the postorbital carina (Fig. 1C). In other respects, all species of Notostomus are similar on the exterior and variations encompass proportions of carinae, denticulation of the carapace and the first abdominal somite.
Meningodora is supported by synapomorphies linked to a reduction of the dorsal pleonic carinae (absent on the second segment, blunt and indistinct on the third segment). The genus encompasses species smaller than Notostomus, which may partly explain the absence of the further strengthening observed in Notostomus. However, some strengthening is still observed in larger species: a long, sharp branchiostegal carina on the carapace ( $\sim$ half of the carapace length) in M. mollis Smith, 1882 and M. compsa (Chace, 1940) (Fig. 1D), and an armament of the third pleonic somite [posterodorsal tooth in M. marptocheles (Chace, 1940) and M. miccyla (Chace, 1940)]. There is an interesting trend linked to M. mollis, which occurs deeper than other Meningodora (Crosnier \& Forest, 1973): this shrimp has a soft body and reduced cornea owing to the deep-living mode of this species. Another trait in Meningodora concerns relative length of the sixth and the fifth pleonic somites: the ratio is $1-1.5$ in M. compsa, 1.5-2 in M. longisulca Kikuchi, 1985 and $>2$ in the rest of the genus. We suggest that this row mirrors increasing movability of the species.

Overall, in the revised group, we observe different evolutionary traits. The first one is linked to an armament of the carapace and pleon with strong and numerous spines and ridges. This trait, likely associated with a defensive function and recorded here in Notostomus, was previously found in other pelagic crustaceans, such as Euphausiacea (Vereshchaka et al., 2019a) and Oplophoridae (Lunina et al., 2019b). The second trait is morphologically opposite to the first one and is linked to a 'smoothening' of the body (reduction of the spines and carinae). This trait, found here in Ephyrina, was previously recorded in the pelagic branch of Benthesicymidae
(Lunina et al., 2019a; Vereshchaka et al., 2020). Analyses of the 'Notostomus + Meningodora' clade retrieved a novel evolutionary trait associated with keeping the body firm and rigid. The shrimps of this clade occur in the deep-sea and have a halfmembranous carapace and pleon, which provide nearly zero buoyancy and, consequently, a reduction of energy loss. Strong carinae on their carapace and pleon may serve as a compensatory structure to provide a necessary supporting structure for locomotion.

A key to species of Ephyrina, Meningodora and Notostomus may be found in Chace (1986); the only species described since then is M. longisulca, which differs from all other Meningodora in the absence of the branchiostegal carina and in the unique (for Meningodora) ratio between the sixth and the fifth pleonic somites (1.7).

## The Unusual mandible: AN EXAMPLE OF A PARALLEL EVOLUTION

In addition to the synapomorphies discussed above, the clades Ephyrina and Meningodora + Notostomus are supported by such a character as the unusual mandible (Fig. 1B), which is easily distinguishable from the mandibles of other decapods (Fig. 1A). We suggest that the characteristic mandibles have evolved in the Ephyrina and Meningodora + Notostomus clades independently as adaptations to feeding on an unusual prey. Indeed, most caridean pelagic shrimps (families Pasiphaeidae, Oplophoridae and Acanthephyridae) are voracious predators, living on small fish, decapods and euphausiids (e.g. review in: Burukovsky, 2009). In particular, Burukovsky (2009) studied in detail the gut content of such representatives of Oplophoroidea as Systellaspis [S. debilis (A. Milne-Edwards, 1881), S. pellucida], Acanthephyra [A. acanthitelsonis Spence Bate, 1888, A. eximia Smith, 1884, A. fimbriata Alcock \& Anderson, 1894, A. kingsleyi Spence Bate, 1888, A. pelagica (Risso, 1816) and A. purpurea A. MilneEdwards, 1881] and Oplophorus [O. gracilirostris, O. novaezealandiae (de Man, 1931), O. spinosus (Brullé, 1839) and O. typus H. Milne-Edwards, 1837], and found that the most common and voluminous dietary items of all these decapods are fish and crustaceans. Shrimps have typical oplophoroid mandibles (reduced molar process and subtriangular incisor process armed with teeth along the entire inner margin; Fig. 1A), which can crush crustacean carapaces and fish bones and further cut tissues.
The prey of Ephyrina, Meningodora and Notostomus is different. Although the feeding of these genera is underexplored, scattered information confirms our suggestion about their different trophic specializations. Examination of the gut content of Notostomus (N. crosnieri and N. elegans), Ephyrina figueirai and

Meningodora vesca shows that the most common and voluminous dietary items of these species differ from those found in other pelagic decapods and are represented by pelagic cnidarians (Burukovsky, 2009). Other studies also indicate that cnidarian tissue is the most common dietary item of Notostomus japonicus (Nishida et al., 1988). Notostomus robustus Smith, 1884 has even been observed from a submersible feeding on the medusa Atolla wyvillei Haeckel, 1880 (Moore et al., 1993). Our study is first to emphasize a link between this type of mandible (Fig. 1B) and feeding on gelatinous organisms but no direct observations on feeding procedure of deep-sea Notostomus, Meningodora or Ephyrina have been made (if possible at all). We may only hypothesize that a sharp, smooth blade is more efficient for the destruction of voluminous soft tissues (feeding objects are large enough) than a thickened serrate margin (teeth are buttressed by a relief). A sharp, smooth blade likely chops tissues (as we use a smooth acute knife to cut butter), whereas a serrated blade saws and crushes tissues (like when we slice bread).

Overall, during colonization of the pelagic realm, the main trophic trend in the evolution of pelagic decapods, including Oplophoroidea, was linked to feeding on crustaceans and fish that was mirrored in the mandible with subtriangular and an entirely serrated incisor process. Two clades of Oplophoroidea, Ephyrina and Meningodora + Notostomus, followed another trophic pathway. They feed, presumably, on gelatinous animals, mainly cnidarians, thus filling a separate ecological niche. Gelatinous animals, which significantly contribute to the pelagic biomass in all depth zones (Vereshchaka et al., 2016), are consumed by a limited number of predators and thus represent a potentially strong food source. Noteworthy, Notostomus is dominant in the meso- and upper bathypelagic of the Subequatorial and Equatorial Atlantic in terms of biomass (Vereshchaka et al., 2019b). The unusual mandibles that evolved in the clades Ephyrina and Meningodora + Notostomus, therefore, represent a remarkable example of parallel evolution.

## CONCLUSIONS

Here we present the first comprehensive phylogenetic revision of the genera Ephyrina, Meningodora and Notostomus based on the synchronous use of 95 morphological characters (all valid species included) and six gene markers ( $71 \%$ of valid species belonging to the target genera included). These three genera have an unusual carapace strengthened with a set of ridges and carinae, and a one-sided serrated mandible, which suggest possible monophyly of this group; a hypothesis we test here.

It is noteworthy that both morphological and molecular analyses retrieve trees with similar topology and a set of statistically supported clades. We show that Ephyrina and Meningodora + Notostomus are separate clades and thus reject the hypothesis of group monophyly. The genera Ephyrina and Notostomus are monophyletic, both on the morphological and on molecular trees; Meningodora gains support only on the morphological trees.
Basic evolutionary traits in the Ephyrina and Meningodora + Notostomus clades are different. In Ephyrina, they are mostly linked to the pereopods (oarlike) and shape of the rostrum (smooth lamina possibly acting as a stempost) favouring active forward motion. The Meningodora + Notostomus clade is predominantly supported by synapomorphies coupled with the carapace and pleon strengthened with ridges and carinae, which is indispensable for Meningodora and Notostomus with their half-membranous integument. Carapace strengthening further evolved into an even more elaborate net of sharp carinae in large Notostomus as a possible response to increasing carapace loads.
Our results suggest that unusual mandibles evolved in the clades Ephyrina and Meningodora + Notostomus independently and represent convergent trophic adaptations. Unlike most pelagic decapods feeding on other crustaceans and fish, both clades follow an alternative pathway and are adapted to feeding on gelatinous organisms, mostly cnidarians. Living on this prey appears to be ecologically advantageous, as species of Notostomus dominate in the low-latitude Atlantic communities in terms of biomass.

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## DATA ACCESSIBILITY

Our data will be deposited in the Dryad Digital Repository.

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## SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article at the publisher's web-site.
Appendix S1. Character scoring.
Appendix S2. Data matrix.
Appendix S3. Retrieved trees.
Appendix S4. Synapomorphies.


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[^1]:    Sequences obtained in this work are indicated in bold. An ' $N$ ' designates gene sequences we were unable to acquire

