**Tamilokus mabinia**, a new, anatomically divergent genus and species of wood-boring bivalve from the Philippines

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Here we describe an anatomically divergent wood-boring bivalve belonging to the family Teredinidae. Specimens were collected off the coast of Mabini, Batangas, Philippines, in February 2018, from sunken driftwood at a depth of less than 2 m. A combination of characteristics differentiates these specimens from members of previously named teredinid genera and species. Most notable among these include: an enlarged cephalic hood which extends across the posterior slope of the shell valves and integrates into the posterior adductor muscle; a unique structure, which we term the ‘cephalic collar’, formed by protruding folds of the mantle immediately ventral to the foot and extending past the posterior margin of the valves; a large globular stomach located entirely posterior to the posterior adductor muscle and extending substantially beyond the posterior gape of the valves; an elongate crystalline style and style sac extending from the base of the foot, past the posterior adductor muscle, to the posteriorly located stomach; calcareous pallets distinct from those of described genera; a prominently flared mantle collar which extends midway along the stalk of the pallets; and, separated siphons that bear a pigmented pinstripe pattern with highly elaborate compound papillae on the incurrent siphon aperture. We used Micro-Computed Tomography (Micro-CT) to build a virtual 3D anatomical model of this organism, confirming the spatial arrangement of the structures described above. Phylogenetic analysis of the small (18S) and large (28S) nuclear rRNA gene sequences, place this bivalve within the Teredindae on a branch well differentiated from previously named genera and species. We propose the new genus and species *Tamilokus mabinia* to accommodate these organisms, raising the total number of genera in this economically and environmentally important family to 16. This study demonstrates the efficacy of Micro-CT for anatomical description of a systematically challenging group of bivalves whose highly derived body plans are differentiated predominantly by soft tissue.
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Title: *Tamilokus mabinia*, a new, anatomically divergent genus and species of wood-boring bivalve from the Philippines.

Short title: *T. mabinia*, a new Philippine shipworm

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Abstract

Here we describe an anatomically divergent wood-boring bivalve belonging to the family Teredinidae. Specimens were collected off the coast of Mabini, Batangas, Philippines, in February 2018, from sunken driftwood at a depth of less than 2 m. A combination of characteristics differentiates these specimens from members of named teredinid genera and species. Most notable among these include: an enlarged cephalic hood which extends across the posterior slope of the shell valves and integrates into the posterior adductor muscle; a unique...
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Introduction

The Teredinidae, commonly referred to as shipworms, are a group of wood-boring bivalves with wide-ranging economic and ecological impacts in coastal marine systems. Their ability to bore
into and digest wood is estimated to cause billions of dollars in damage per year to coastal structures, such as piers, jetties, wharfs, fishing equipment and aquaculture pens (Turner (1966), Distel 2003). Yet this same ability allows them to play a fundamental role in carbon cycling by processing recalcitrant organic carbon trapped in wood and liberating nutrients and energy that would otherwise be less accessible to other organisms (Nishimoto et al. 2015, Charles et al. 2016).

The unique wood-boring life-style of the Teredinidae has led to several anatomical specialisations and as such, this family is believed to exhibit more variation in morphology than any other group in the Mollusca (Turner 1988). In contrast to more typical bivalves, the shell is greatly reduced in size and has prominent pedal and posterior gapes, allowing the protrusion of the foot and elongate body respectively. The location of the shell valves at the extreme anterior end of the body covers little of the visceral mass and offers little or no protection to the animal. Instead, the shell, which features tiny denticulated ridges on its surface, functions as an effective drilling and grinding tool for burrowing into wood (Nair 1966, Turner 1966, Voight 2015). The body is long, vermiform and enclosed by a calcareous tube that is secreted onto the burrow walls, surrounding the animal from the posterior edge of the valves to the burrow opening. The siphons are flanked by a pair of calcareous, paddle or feather-like structures, known as the pallets. These pallets, which are unique to Teredinidae, serve as watertight and predator-resistant plugs to seal the burrow entrance (Nair 1966, Turner 1966, Voight 2015). Currently, there are 16 accepted genera composed of 70 species in the Teredinidae (Bulatov 1933, Turner 1966, Macintosh 2012, BORGES et al. 2018, Velásquez et al. 2018). However, it has been recently suggested that between 80-100 species likely exist (Huber et al. 2015, Shipway et al. 2016).
Using integrative taxonomic methods, including anatomical, morphological and molecular data (Borges et al. 2012, Shipway, O’Connor et al. 2016), we identify and describe a new species and genus of wood-boring teredinid from the Philippines, which we name *Tamilokus mabinia*.

Materials & Methods

Specimen Collection

Nine specimens of *Tamilokus mabinia* were collected in February 2018 off the coast of Balayan Bay, Mabini, Batangas, Philippines (13.758°N, 120.925°E), at a depth of < 2 m, from a log measuring approximately 2m in total length, as part of the Philippine Mollusk Symbiont (PMS) International Collaborative Biodiversity Group (ICBG) expedition. The specimen location map (Fig. 1) was produced and approximate coordinates were determined using the National Oceanographic and Atmospheric Administration Map Viewer (https://www.nauticalcharts.noaa.gov/ENCOnline/enconline.html). Specimens were carefully extracted from their burrows, photographed (Fig. 2), measured, fixed in 4% formaldehyde solution freshly prepared from paraformaldehyde (PFA), washed, and dehydrated through an ethanol series (30%, 50% and 70%, 30 minutes per wash, x2 washes) with final storage in 70% ethanol. Partial specimens, identifiable as *T. mabinia* based on the unique morphology of the pallets, siphons and cephalic collar, were dissected or were fixed in 100% ethanol for DNA preservation. Morphological features (Fig. 3) and calcareous structures (Fig. 4) were imaged using the Keyence VHX-6000 Digital Microscope (Osaka, Japan). Specimens were directly compared with all described teredinid genera from the Harvard Museum of Comparative Zoology. Part of this work was completed under the supervision of the Department of
Agriculture-Bureau of Fisheries and Aquatic Resources, Philippines (DA-BFAR) in compliance with all required legal instruments and regulatory issuances covering the conduct of the research. All Philippine specimens used in this study were obtained using Gratuitous Permit GP-0140-17 issued by DA-BFAR.

**Micro-Computed Tomography**

Specimen PMS-4051L was stained for 20 days in 10% iodine prior to imaging. Imaging was performed using a SkyScan 1173 μCT scanner (Bruker Micro-CT, Kontich, Belgium) equipped with a Hamamatsu 130/300 tungsten X-ray source and a FlatPanel Sensor camera detector with 2240×2240 pixels. Scanning parameters were as follows: source voltage = 60 kV; source current = 100 μA; exposure time = 900 ms; frames averaged = 3; frames acquired over 180° = 960; filter = no; binning = no; flat field correction = activated; scanning time = (x3) 00:28:14; number of connected scans = 3; and, random movement = ON (10). Reconstruction of the raw data was accomplished using the software provided with the scanner (NRecon 1.6.6.0, Bruker Micro-CT, Kontich, Belgium). The following settings were employed to enhance image contrast and to compensate for ring and streak artefacts; smoothing = no, ring artefact correction = 4, and beam hardening correction = activated. Reconstructed scans were then analyzed using CTVox (Bruker Micro-CT, Kontich, Belgium). A Micro-CT 3D rendered model of *Tamilokus mabinia*, including major anatomical structures and transverse cross sections of the model is shown in Figure 5.

**DNA Extraction & Amplification**
DNA was extracted from the siphonal tissue and associated musculature from one specimen of *Tamilokus mabinia* (PMS-3943P), using the DNeasy Blood & Tissue kit (Qiagen, Hilden, Germany) following manufacturer’s protocol. Approximate purity, concentration and yield of DNA were determined by UV spectrophotometry. Genomic DNA was cryo-preserved at -80 °C and archived at the Ocean Genome Legacy Center of New England Biolabs, Northeastern University, Nahant, MA, USA (accession number in Table1). The small (18S) and large (28S) subunit nuclear rRNA genes were amplified from the resultant DNA preparation by polymerase chain reaction (PCR). Amplification reactions were prepared using 12.5 μL of high-fidelity polymerase solution (OneTaq, New England Biolabs, Ipswich, Massachusetts), 0.5 μL of each primer (10 mM), 1–2 μL DNA template (10–20 ng/μL), brought to a total volume of 25 μL with purified water. Fragments of the small (18S) and large (28S) subunit nuclear rRNA genes were amplified using the primer pairs 18S EukF (5’–WAY-CTG-GTT-GAT-CCT-GCC-AGT–3’) and 18S EukR (5’–TGA-TCC-TTC-YGC-AGG-TTC-ACC-TAC–3’) (Medlin et al. 1988); 28S-NLF184-21 (5’–ACC-CGC-TGA-AYT-TAA-GCA-TAT–3’) and 28S-1600R (5’–AGC-GCC-ATC-CAT-TTT-CAG-G–3’) (Distel et al. 2011), resulting in amplicons of approximately 1,686 and 1,416 base pairs, respectively. The PCR amplification proceeded as follows: an initial denaturation step of 94 °C for three minutes, followed by 35 cycles with a denaturation step of 94 °C for 20 s, an annealing step of 64 °C for 40 s for the 18S and 63 °C for 30 s for 28S, an extension step of 68 °C for 60 s and a final extension of 68 °C for five minutes. All reactions were performed on a PTC-200 Thermal Cycler (MJ Research, Quebec, Canada).

For each template, three separate amplicons, all produced under identical conditions, were pooled, cleaned and concentrated using the Zymo Clean & Concentrator Kit (Irvine, CA).
Resulting products were sequenced bidirectionally on a 3730xl DNA Analyzer (Life Technologies, Grand Island, NY) using the Big Dye Terminator 3.1 Cycle Sequencing Kit (Life Technologies, Grand Island, NY) at New England Biolabs (Ipswich, Massachusetts). Primary sequence data from the small (18S) and large (28S) subunit nuclear rRNA genes were submitted to GenBank (NCBI) under accession numbers MH974682 and MH974683, respectively.

**Phylogenetic Analysis**

The 18S and 28S rRNA gene sequences from specimen PMS-3943P were concatenated and aligned as in Distel *et al.* (Distel, Amin *et al.* 2011) with sequences representing 10 of 16 recognized genera (Bulatov 1933, Turner 1966) including: *Bankia* Gray 1842, *Dicyathifer Iredale* 1932, *Kuphus* Guettard 1770, *Lyrodus* Gould 1870, *Nausitora* Wright 1864, *Neoteredo* Bartsch 1920, *Spathoteredo* Moll 1928, *Teredo* Linnaeus 1758, *Teredora* Bartsch 1921, and *Teredothyra* Bartsch 1921. Phylogenetic analysis was performed using MrBayes version 3.2.6 (Ronquist *et al.* 2003) implemented at Geneious version 10.2.2 (https://www.geneious.com) using GTR + I + Γ nucleotide substitution model. *Placopecten magellanicus* was specified as the outgroup, the chain length was set to 5 million, subsampling every 2,000 generations and discarding the first 20% of the results as burn-in.

**Zoobank Registration**

The electronic version of this article in Portable Document Format (PDF) will represent a published work according to the International Commission on Zoological Nomenclature (ICZN), and hence the new names contained in the electronic version are effectively published under that Code from the electronic edition alone. This published work and the nomenclatural acts it
Results

Systematics

Family Teredinidae Rafinesque, 1815

Tamilokus Shipway, Distel & Rosenberg, gen. nov.

urn:lsid:zoobank.org:act:61B916B1-54D1-418D-B1A3-342162C261C2

Type species: Tamilokus mabinia sp. nov.

Type material: Holotype PMS-3916Y; paratypes PMS-3899P, PMS-3915X, PMS-3943P, PMS-3949Y, PMS-4037Y, and PMS-4051L. Smallest to largest specimens measured 6.2 cm – 15.4 cm in total body length. The holotype is currently held at the Academy of Natural Sciences of Philadelphia (ANSP) and will be deposited in the National Museum of the Philippines in Manila pending completion of an ongoing reorganization. Paratypes are deposited at ANSP, at the Marine Science Institute of the University of the Philippines (MSI) and at the Ocean Genome Legacy Center of New England Biolabs, Northeastern University (OGL). For catalog numbers assigned by holding institutions see Table 1.
Type locality: Balayan Bay, off Mabini, Batangas Province, Philippines (coordinates 13.758°N, 120.925°E).

Comparative Material: The following material was examined: *Teredora malleolus* (Turton, 1822) MCZ-350541 (3 specimens); *Teredora princesae* (Sivickis, 1928) MCZ-232090 (5 specimens); *Uperotus clava* (Gmelin, 1791) MCZ-238009 (3 specimens); and, *Uperotus panamensis* (Bartsch, 1922) MCZ-357824 (4 specimens).

Diagnosis: pallets triangular-shaped with an ovate, flattened stalk, prominent cephalic hood and cephalic crest, crystalline style extends beyond posterior adductor muscle, caecum U-shaped doubling back upon itself, siphons with distinct pin-striped pigmentation with incumbent siphon featuring three rows of papillae.

Etymology: *Tamilokus* (masculine), in recognition of the common name for shipworm in the Philippines, ‘tamilok’.

Habitat: Marine, wood-borer.

Description: Pallets very small in relation to body length (Figs. 2A & 3A), formed of a broad, flattened, translucent and ovate stalk, with a solid, white calcareous blade (Fig. 4A); cephalic hood prominent, extending to cover the posterior slope of the shell valves (Figs. 2A, 3A-C & 5A, C); “cephalic collar”, formed from protruding folds of the mantle, extending along the ventral surface from the base of the foot to a point slightly posterior to the posterior margin of the shell valves (Figs. 2A-B, 3A, D-E, 5A-B); large crystalline style and style sac originating within the cephalic collar and extending beyond the posterior adductor muscle into the posteriorly located stomach (Figs. 5A-E); large globular stomach (Figs. 2A-B & 5A, E-F) located posteriorly in relation to posterior adductor muscle and valve pedal gape, composed of three lobes, with
muscular opening into caecum (Figs. 5A, F); large digestive glands (Figs. 2A-C & Figs. 5A, C-E); caecum (Figs. 2A-C & 5A, D-I) large, elongate, U-shaped, doubling back upon itself anteriorly towards the right (Fig. 2B & 5D-I), lacking a typhlosole (Figs. 5D-I); intestine unusually broad and voluminous (Fig. 2B), especially where it passes ventral to the caecum (Figs. 5G-I), containing short ovoid faecal pellets packed in multiple rows across its width (Fig. 2B); intestine extends anteriorly, looping over crystalline style sac (Figs. 5C-E), then doubles back posteriorly, looping under the caecum before extending anteriorly on the dorsal surface of the caecum, finally looping under the posterior adductor muscle before extending a short distance posteriorly and opening into the anal canal; anal canal is open and does not retain faeces; gonad located centrally (Figs. 5A, I-K), beginning posterior to the caecum and ending at the heart; heart located medially (Figs. 5J-K); gill extends from the siphons anteriorly to the posterior tip of the gonads (Figs. 5A, K-L); prominently flared mantle collar around siphons and pallets (Figs. 2B, 3A & 5A).

Remarks: *Tamilokus* gen. nov may be easily differentiated from all other genera within the family Teredinidae based on pallet morphology. The simple triangular cup-shaped pallet and thick ovate stalk of *Tamilokus* may be distinguished from the pallets of: *Bankia* Gray 1842, *Nausitora* Wright 1864, *Nototeredo* Bartsch 1923, and *Spathoteredo* Moll 1928 by the lack of segmentation; from *Lyrodus* Gould 1870, *Teredo* Linnaeus 1758, *Zachia* Bulatoff & Rjabtschikoff 1933, and *Nivanteredo* Velásquez & Shipway 2018 by the absence of a periostracum and presence of a broad ovate stalk; from *Teredothyra* Bartsch 1921 by the presence of a single undivided cup; from *Dicyathifer* Iredale 1932 and *Kuphus* Guettard 1770 by the absence of a medial ridge; from *Bactronophorus* Tapparone Canefri 1877 by the absence of a dagger-like extension; from *Neoteredo* Bartsch 1920, *Psiloteredo* Bartsch 1922, *Teredora*
Bartsch 1921 and *Uperotus* Guettard 1770 by the cup- rather than paddle-shaped blades, and
from the latter three genera by the lack a distinct thumb-nail like depression bearing concentric
or radiating ridges.

Additionally, *Tamilokus* may be easily distinguished from *Bactronophorus* Tapparone Canefri
1877, *Dicyathifer* Iredale 1932, *Neoteredo* Bartsch 1920 and *Teredothyra* Bartsch 1921 by the
absence of a muscular sphincter at the posterior end of the anal canal; from *Lyrodus* Gould 1870,
*Teredo* Linnaeus 1758 and *Zachsia* Bulatoff & Rjabtschikoff 1933 by the absence of brood
pouches on the gill; from *Neoteredo* Bartsch 1920 by the absence of dorsal lappets; and from
*Kuphus* Guettard 1770 by the presence of a caecum and the absence of a strong muscular collar
surrounding the valves.

*Tamilokus* is similar to *Teredora* Bartsch 1921 and *Uperotus* Guettard 1770 in that it possesses a
U-shaped caecum that, after passing posteriorly from the stomach, doubles back upon itself
towards the right and extends anteriorly, terminating near its origin on the right side of the
stomach (Turner, 1966). However, *Tamilokus* can be easily distinguished from these genera
based on pallet morphology (as previously described). Additionally, the gills of *Teredora* and
*Uperotus* extend the entire length of the animal from the base of the siphons to the mouth,
whereas those of *Tamilokus* terminate near the posterior end of the caecum; the labial palps in
both *Teredora* and *Uperotus* are large and free, but are very small and attached in *Tamilokus*; the
crystalline style in both *Teredora* and *Uperotus* is located anteriorly to the posterior adductor
muscle, but extends from the base of the foot well beyond the posterior margin of the adductor
muscle in *Tamilokus*; the stomach of both *Teredora* and *Uperotus* is located anterior to posterior adductor muscle, whereas the stomach of *Tamilokus* is located posterior to the posterior adductor muscle; the heart of both *Teredora* and *Uperotus* is positioned anteriorly, but is medially positioned in *Tamilokus*; the siphons of *Teredora* and *Uperotus* are united along their entire length, but are separate in *Tamilokus*; the incurrent siphons of both *Teredora* and *Uperotus* feature a single primary row of papillae, whereas the incurrent siphon of *Tamilokus* has an additional secondary row that includes compound branched papillae; and, the cephalic hood in both *Teredora* and *Uperotus* is inconspicuous, whereas the cephalic hood in *Tamilokus* is prominent and covers the posterior slope of the shell valves. Finally, *Tamilokus* may be recognized by the presence of the cephalic collar which is unique to this genus. Taxonomic characters differentiating *Tamilokus* from *Teredora* and *Uperotus* are summarized in Table 2.

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*Tamilokus mabinia* Shipway & Distel, *sp. nov.*

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**Etymology:** *Mabinia* (noun in apposition), in honour of Apolinario Mabini, a Philippine national hero, and pertaining to the type location of the specimens from Mabini, Batangas, Philippines.

**Description:** All characteristics of the genus, plus; dorsal anterior slope of shell valves highly elongated (Fig. 4B); incurrent and excurrent siphons ringed with papillae (Fig. 3F); incurrent siphon features outer ring of long papillae, and inner ring of shorter compound or branched papillae (Fig. 3G); siphon tips are ringed with pink to brownish red pigment that extends posteriorly in narrow closely spaced parallel stripes (Figs. 2A-C & 3A).
Definition: Pallets elongate, composed of an unsegmented blade built upon a central stalk (Fig. 4A); blade triangular, single concave U-shaped cup on distal margin with slightly more pronounced curvature on outer face; periostracum absent; mantle collar flared, extending approximately half the length of the pallet; stalk and blade approximately equal in length; stalk translucent, ovate and flattened in both sagittal and transverse section; blade calcarceous and white.

Phylogeny

Bayesian analysis of concatenated small (18S) and large (28S) nuclear rRNA gene sequences (Fig. S1) indicate a basal position for Tamilokus within Teredinidae, sharing a well-supported node with Teredora, but separated from that taxon by a branch length comparable to those separating most accepted teredinid genera. A subtree excerpted from Figure S1 is displayed in Figure 6.

Discussion

Herein, we describe a divergent member of the Teredinidae that, based on anatomical, morphological and molecular characteristics, is distinct from members of described genera. Among the distinguishing features of Tamilokus mabinia, the most striking are the posterior position of the stomach and the elongation of the crystalline style and style sac to accommodate this posterior location. (Fig. 5A). This position of the stomach is found only in Tamilokus but is similar to that found in Kuphus, except that in the latter, the entire visceral mass is located posterior to the posterior adductor. The distinct positions of these two taxa on the phylogenetic tree of Teredinidae suggests that this similarity is due to convergence rather than homology. The
extension of the crystalline style, from a position ventral to and near the anterior end of the foot to a position well posterior to the shell valves, is unique to Tamilokus (Turner 1966). In other bivalves, cilia-mediated rotation of the crystalline style is thought to aid digestion by mixing enzymes and reducing food particle size through mechanical action (Robin et al., Nelson 1918, Edmondson 1920, Yonge 1923, Lavine 1946, Morton 1952, Horiuchi et al. 1982, Alyakrinskaya 2001, Sakamoto et al. 2008, Mackenzie et al. 2014). However, the functional significance of the exceptionally large style of Tamilokus remains to be explained.

Tamilokus also features a characteristic structure absent in other genera, which we term the ‘cephalic collar’. This feature is formed by protruding folds of the mantle and is located directly ventral to the foot and anterior adductor muscle and extends posterior to the shell valves on the ventral mantle surface (Figs. 2A-B, 3A, D-E & 5A). Figures 3D-E show that the cephalic collar can be contracted or expanded, and Micro-CT scans reveal that the style sac is located within the cephalic collar (Figs. 5A-B). These observations suggest that the cephalic collar may serve to accommodate the large size of the crystalline style and may facilitate its movement.

Another unusual feature of Tamilokus is the caecum, which folds sharply to the right at its midpoint, doubling back upon itself to form a compressed U-shape. Among the Teredinidae, only two described genera, Teredora and Uperotus share this feature (Turner 1966). However, Teredora and Uperotus differ from Tamilokus in several important ways. In these genera, the gills extend almost the entire length of the animal from the base of the siphons to the mouth, proportionately the longest gills among the family (Turner 1966). In addition to the extended gills, the labial palps, which help sort and direct filtered particles to the mouth, are prominent
and detached. This observation led to suggestions that these genera are specialized for planktotrophy rather than wood feeding (Turner 1966, Nair et al. 1971). This hypothesis, however, preceded the discovery of cellulolytic symbionts in the gills of shipworms, providing the competing hypothesis that larger gills are adaptive for wood digestion (Distel, Amin et al. 2011).

In contrast to *Teredora* and *Uperotus*, the gills of *Tamilokus* extend only to the posterior end of the caecum (Fig. 5A) and the palps are small, fixed, and appear to be vestigial. Also, in contrast to these genera, the intestine of *Tamilokus* is extremely wide. In all specimens examined, the capacious intestine was tightly packed with light-colored excavated wood fiber (Fig. 2B), matching the color of the wood substrate in which the animals burrowed. No evidence of planktonic organisms could be discerned in caecum contents or faecal material by microscopic examination.

Siphon morphology also distinguishes *Tamilokus* from *Teredora* and *Uperotus*. The siphons of the former are separate while those of the latter two are united along their entire length. The presence of pink pinstripes on both the incurrent and excurrent siphons of *Tamilokus* (Figs. 2 & 3) are also unique among described species. Additionally, the siphons are tipped with two rows of papillae on the incurrent siphon aperture; the primary outer row is simple in structure, while the inner secondary row include shorter compound papillae with multiple branches (Figs. 3F-G). Characteristics of the siphons have been suggested as potentially useful in teredinid identification (Turner 1966, Nair and Saraswathy 1971), but have been described for only a small number of
species (Roch 1940, Turner 1966, Lopes et al. 1998, De Moraes et al. 2003). Although the degree of genetic, ontogenetic and ecophenotypic variation in teredinid siphons has yet to be documented (Turner 1966), characterizing these features could prove valuable for field identification, as specimens could be studied in situ without the necessity of extracting the teredinid from its burrow.

Aside from its unique anatomical and morphological features, the classification of Tamilokus as a new genus is further supported by molecular phylogenetic data. Phylogenetic analysis based on concatenated small (18S) and large (28S) nuclear rRNA gene sequences reveal well supported relationships between Tamilokus and basal taxa within Teredinidae. However, the branch length of Tamilokus is among the longest in the family, indicating divergence comparable to or greater than those separating accepted teredinid genera (Fig. 6). Anatomical and morphological characters are also consistent with the basal position of Tamilokus. As in other basal members of the family, the stomach of T. mabinia is globular (Type 2 of Turner 1966 (Turner 1966)), the intestine loops anteriorly over the crystalline style sac, and brooded larvae are absent in the gills of all specimens examined. This is consistent with the hypothesis that these characteristics are plesiomorphic and that the elongate stomach, absence of the anterior intestinal loop and complex reproductive strategies are apomorphic (Distel, Amin et al. 2011).

In this investigation, Micro-CT has been invaluable for elucidating soft tissue anatomy. This has been especially important in this systematically challenging group, which is characterised by highly derived body parts and few taxonomically informative calcareous features. Previously, Micro-CT has been used to determine quantitative measurements of wood degradation and boring rates of Teredinidae (Amon et al. 2015). Additionally, Micro-CT has been used to reveal
exceptionally well-preserved fossilized wood-boring bivalves in mid-Cretaceous wood, with silicified soft parts and overall body-plan characteristic of the Teredinidae (Robin, Velasquez et al.). Herein, we further demonstrate the power of this technique to reveal the relationships among soft tissue features in three dimensions, and in the evaluation of likely mode of nutrition and life history strategy, while minimizing distortion that can result from manual dissection.

The extent of diversity within the Teredinidae has long been a subject of debate. Historically, as many as 130 species have been recognized (Moll et al. 1931), but many were later synonymized in the first comprehensive work on the taxonomy of this family (Turner 1966). Recently, it has been suggested that teredinid diversity may be greater than previously thought (Huber, Langleit et al. 2015, Shipway, O’Connor et al. 2016). This view is supported by the recent description of one new genus and several new species of teredinid bivalves (Macintosh 2012, BORGES and Merckelbach 2018, Velásquez and Shipway 2018). In addition, further reports suggest several more species (Shipway, O’Connor et al. 2016, Treneman et al. 2018) and at least one more genus (Lozouet et al. 2008) remain to be described.

Conclusions

We describe a new and anatomically divergent genus and species of wood-boring bivalve (Teredinidae) from the Philippines, which we name TAMILOKUS MABINIA. A detailed Micro-CT 3D render of T. mabinia reveals several anatomically divergent characters, including a new structure which we term the ‘cephalic collar’, a prominent crystalline style sac and crystalline style originating within the cephalic collar and extending beyond the posterior adductor muscle into the posteriorly located stomach, and a caecum (wood-storing organ) which doubles back upon
itself. In addition, the unique pallets, the primary taxonomic character of this family, could not be placed in any existing genera. Phylogenetic analysis places *T. mabinia* in a separate taxon among other basal members of the Teredinidae. The addition of *Tamilokus* raises the total number of genera in this economically and environmentally important group of wood-boring bivalves to 17.

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References

Alyakrinskaya, I. (2001). "The dimensions, characteristics and functions of the crystalline style of molluscs." *Biology Bulletin of the Russian Academy of Sciences* 28(5): 523-535.

Amon, D. J., D. Sykes, F. Ahmed, J. T. Copley, K. M. Kemp, P. A. Tyler, C. M. Young and A. G. Glover (2015). "Burrow forms, growth rates and feeding rates of wood-boring Xylophagaidae bivalves revealed by micro-computed tomography." *Frontiers in Marine Science* 2: 10.

Borges, L. M. and L. M. Merckelbach (2018). "Lyrodus mersinensis sp. nov.(Bivalvia: Teredinidae) another cryptic species in the Lyrodus pedicellatus (Quatrefages, 1849) complex." *Zootaxa* 4442(3): 441-457.

Borges, L. M., H. Sivrikaya, A. Le Roux, J. Shipway, S. Cragg and F. Costa (2012). "Investigating the taxonomy and systematics of marine wood borers (Bivalvia: Teredinidae) combining evidence from morphology, DNA barcodes and nuclear locus sequences." *Invertebrate Systematics* 26(6): 572-582.

Bulatov, G. (1933). "Eine neue Gattung aus der Familie der Teredinidae aus dem Japanischen Meer." *Zoologischer Anzeiger* 104: 165-176.

Charles, F., J. Coston-Guarini, J.-M. Guarini and S. Fanfard (2016). "Wood decay at sea." *Journal of Sea Research* 114: 22-25.

De Moraes, D. T. and S. G. Lopes (2003). "The functional morphology of Neoteredo reynei (Bartsch, 1920)(Bivalvia, Teredinidae)." *Journal of molluscan studies* 69(4): 311-318.

Distel, D. L. (2003). "The biology of marine wood boring bivalves and their bacterial endosymbionts." *Wood deterioration and preservation* 845: 253-271.
Distel, D. L., M. Amin, A. Burgoyne, E. Linton, G. Mamangkey, W. Morrill, J. Nove, N. Wood and J. Yang (2011). "Molecular phylogeny of Pholadoidea Lamarck, 1809 supports a single origin for xylotrophy (wood feeding) and xylotrophic bacterial endosymbiosis in Bivalvia." Molecular Phylogenetics and Evolution 61(2): 245-254.

Edmondson, C. H. (1920). "The reformation of the crystalline style in Mya arenaria after extraction." Journal of Experimental Zoology 30(3): 259-291.

Horiuchi, S. and C. E. Lane (1982). "Carbohydrases of the crystalline style and hepatopancreas of Strombus gigas linnê." Comparative Biochemistry and Physiology 74A(3): 673-685.

Huber, M., A. Langleit and K. Kreipl (2015). Compendium of bivalves 2. ConchBooks.

Lavine, T. F. (1946). "A study of the enzymatic and other properties of the crystalline style of clams: evidence for the presence of a cellulase." Journal of cellular and comparative physiology 28(2): 183-195.

Lopes, S. and W. Narchi (1998). "Functional anatomy of Nausitora fusticula (Jeffreys, 1860)(Bivalvia: Teredinidae)." The Veliger 41(3): 274-288.

Lozouet, P. and J. C. Plaziat (2008). "Mangrove environments and molluscs." Abatan River, Bohol and Panglao Islands, Central Philippines. ConchBooks, Hackenheim.

Macintosh, H. (2012). "Lyrodus turnerae, a new teredinid from eastern Australia and the Coral Sea (Bivalvia: Teredinidae)." Molluscan Research 32(1): 36.

Mackenzie, A. L. and C. Marshall (2014). "Proteins in the crystalline styles of the marine mussels Perna canaliculus Gmelin and Mytilus galloprovincialis Lamarck." Journal of Shellfish Research 33(3): 673-685.

Medlin, L., H. J. Elwood, S. Stickel and M. L. Sogin (1988). "The characterization of enzymatically amplified eukaryotic 16S-like rRNA-coding regions." Gene 71(2): 491-499.

Moll, F. and F. Roch (1931). "The Teredinidae of the British Museum, the natural history museums at Glasgow and Manchester, and the Jeffreys collection." Journal of Molluscan Studies 19(4): 201-218.

Morton, J. (1952). "The role of the crystalline style." Journal of Molluscan Studies 29(2-3): 85-92.

Nair, N. B. (1966). "Ecology of marine fouling and wood boring organisms of western Norway." Sarsia 8: 1-88.

Nair, N. B. and M. Saraswathy (1971). The Biology of Wood-Boring Teredinid Molluscs. Advances in Marine Biology. S. R. Frederick and Y. Maurice, Academic Press. Volume 9: 335-509.

Nelson, T. C. (1918). "On the origin, nature, and function of the crystalline style of lamellibranchs." Journal of Morphology 31(1): 53-111.

Nishimoto, A., T. Haga, A. Asakura and Y. Shirayama (2015). "An experimental approach for understanding the process of wood fragmentation by marine wood borers in shallow temperate waters." Marine Ecology Progress Series 538: 53-65.

Robin, N., M. Velasquez, A. Boura, G. Garcia, C. Jauvion, J. M. Boiteau, B. Gomez, V. Daviero-Gomez and X. Valentín "The oldest shipworms (Bivalvia, Pholadoidea, Teredinidae) preserved with soft parts (western France): insights into the fossil record and evolution of Pholadoidea." Palaeontology.

Roch, F. (1940). Die Terediniden des Mittelmeeres, Deutsch-Italienisches Institut für Meeresbiologie zu Rovigno d'Istria.

Ronquist, F. and J. P. Huelsenbeck (2003). "MrBayes 3: Bayesian phylogenetic inference under mixed models." Bioinformatics 19(12): 1572-1574.

Sakamoto, K., S. Uji, T. Kurokawa and H. Toyohara (2008). "Immunohistochemical, in situ hybridization and biochemical studies on endogenous cellulase of Corbicula japonica." Comparative Biochemistry and Physiology Part B: Biochemistry and Molecular Biology 150(2): 216-221.

Shipway, J., R. O’Connor, D. Stein, S. Cragg, T. Korshunova, A. Martynov, T. Haga and D. Distel (2016). "Zachsia zenkewitschi (Teredinidae), a rare and unusual seagrass boring bivalve revisited and redescribed." PloS one 11(5): e0155269.
Treneman, N. C., L. M. Borges, J. R. Shipway, M. J. Raupach, B. Altermark and J. T. Carlton (2018). "A molecular phylogeny of wood-borers (Teredinidae) from Japanese Tsunami Marine Debris." *Aquatic Invasions* **13**(1): 101-112.

Turner, R. D. (1966). "A survey and illustrated catalogue of the Teredinidae (Mollusca: Bivalvia)." *A survey and illustrated catalogue of the Teredinidae (Mollusca: Bivalvia).*

Turner, R. D. (1988). Cellulolytic nitrogen-fixing bacteria in the Teredinidae (Mollusca: Bivalvia). *Biodeterioration* **7**, Springer: 743-748.

Turner, R. T. (1966). "A survey and illustrated catalogue of the Teredinidae." *Spec. publ. Museum of Comparative Zoology.*

Velásquez, M. and J. R. Shipway (2018). "A new genus and species of deep-sea wood-boring shipworm (Bivalvia: Teredinidae) Nivanteredo coronata n. sp. from the Southwest Pacific." *Marine Biology Research: 1*-10.

Voight, J. R. (2015). "Xylotrophic bivalves: aspects of their biology and the impacts of humans." *Journal of Molluscan Studies* **81**(2): 175-186.

Yonke, C. (1923). "Studies on the Comparative Physiology of Digestion: I.--The Mechanism of Feeding, Digestion, and Assimilation in the Lamellibranch Mya." *Journal of Experimental Biology* **1**(1): 15-64.
Figure 1

Specimen collection site:

A, map of collection site and B, overview of geographic location. Scale bar = 0.5 km.

*Note: Auto Gamma Correction was used for the image. This only affects the reviewing manuscript. See original source image if needed for review.
**Figure 2** (on next page)

*Tamilokus mabinia:*

A, Holotype (PMS-3916Y); B, paratype (PMS-3915X); C, paratype (PMS-4051L). Ca, caecum; CC, cephalic collar; CH, cephalic hood; DG, digestive glands; Ft, foot; Gi, gill; In, Intestine; MC, mantle collar; Pa, pallet; Si, siphon; St, Stomach; SV, shell valve. Scale bar = 1.0 cm.
Figure 3 (on next page)

Morphological characters of *Tamilokus mabinia*:

A, *Tamilokus mabinia* major morphological characters (PMS-3949Y). Scale bar = 5.0 mm; B-C, cephalic hood; D-E, cephalic collar; F, siphon features; G, magnified region of the incurrent siphon. Scale bar = 500 μm. CC, cephalic collar; CH, cephalic hood; ES, excurrent siphon; Ft, foot; IP, inner papillae; IS, incurrent siphon; MC, mantle collar; Pa, pallet; OP, outer papillae; Si, siphon.
Figure 4

Calcareous structures of *Tamilokus mabinia*:

A, Inner surface (left) and outer surface (right) of pallet. Specimen PMS-3943P. Scale bar = 1.0 mm; B, shell valve outer surface. Specimen PMS-3915X. Scale bar = 500 μm. The valves are very thin and brittle. Note that the ventral tip of the valve is damaged. Anterior to left, dorsal is toward top.
Figure 5 (on next page)

Microcomputed Tomography 3D render of *Tamilokus mabinia*:

A, major anatomical structures; B – M transverse sections of A from anterior to posterior. AC, anal canal; Ca, caecum; CC, cephalic collar; CH, cephalic hood; CS, crystalline style; CSS, crystalline style sac; DG, digestive gland; EBV, efferent branchial vein; Ft, foot; Gi, gill; Go, gonad; He, heart; In, intestine; MC, mantle collar; MO, mouth; Oe, oesophagus; PAM, posterior adductor muscle; SCA; stomach-caecum aperture; Si, siphon; St, stomach; SV, shell valve. Specimen PMS-4051L. Scale bar = 2.5 mm.
**Figure 6** (on next page)

Phylogenetic position of *Tamilokus mabinia* among the Teredinidae:

A sub tree excerpted from a Bayesian analysis of the concatenated 18S and 28S nuclear rRNA gene sequences obtained from specimen PMS-3943P. The tree was constructed using the taxa presented in Distel *et al.*, (2011). The full tree is presented in Supplemental Figure 1. Numbers at nodes indicate posterior
**Table 1** (on next page)

Specimen details and collection site information.

*The holotype will be transferred to the National Museum of the Philippines pending completion of an ongoing reorganization. †Partial specimen with missing valves.
**Table 1. Specimen details and collection site information.**

| Specimen Number | Holding Institution | Accession Number | Type     | Preparation                                                                 | Figure                  |
|------------------|---------------------|------------------|----------|------------------------------------------------------------------------------|-------------------------|
| PMS-3899P        | MSI                 | PMS-3899P        | Paratype | Preserved in 70% ethanol                                                     | -                       |
| PMS-3915X        | ANSP                | A476699          | Paratype | Preserved in 70% ethanol                                                     | Figs. 2B & 4B           |
| PMS-3916Y*       | ANSP                | A476699          | Holotype | Preserved in 70% ethanol                                                     | Fig. 2A                 |
| PMS-3943P†       | MSI                 | PMS-3943P        | Paratype | Preserved in 70% ethanol                                                     | Figs. 4A & 6            |
| PMS-3943P†       | OGL                 | E28692           | -        | Genomic DNA voucher                                                          | Fig. 6                  |
| PMS-3949Y        | MSI                 | PMS-3949Y        | Paratype | Preserved in 70% ethanol                                                     | Fig. 3                  |
| PMS-4037Y        | ANSP                | A476699          | Paratype | Fixed in 4% PFA, transferred to 70% ethanol                                   | -                       |
| PMS-4051L        | ANSP                | A476699          | Paratype | Fixed in 4% PFA, transferred to 70% ethanol, stained with 10% iodine for Micro-CT. | Figs. 2C & 5            |

*The holotype will be transferred to the National Museum of the Philippines pending completion of an ongoing reorganization.

†Partial specimen with missing valves.
Table 2 (on next page)

Diagnostic characters of all teredinid genera featuring a U-shaped caecum.
Table 2: Diagnostic characters of all teredinid genera featuring a U-shaped caeca.

| Taxonomic character | Tamilokus [Shipway, Rosenberg & Distel, 2018] | Teredora [Bartsch, 1921] | Uperotus [Guettard, 1770] |
|---------------------|-----------------------------------------------|---------------------------|---------------------------|
| Labial palps        | Reduced/absent                                | Free                      | Large and free            |
| Crystalline style   | Extends from base of foot beyond posterior adductor muscle | Located anterior to posterior adductor muscle | Located anterior to posterior adductor muscle |
| Stomach             | Globular, located posterior to posterior adductor muscle | Globular, located anterior to posterior adductor muscle | Globular, located anterior to posterior adductor muscle |
| Caecum              | Doubles back upon itself, to the right         | Doubles back upon itself, to the right | Doubles back upon itself, to the right |
| Caecal typhlosole   | Absent                                        | Rudimentary               | Absent                    |
| Heart               | Median position                               | Located anteriorly        | Located anteriorly        |
| Gills               | Located posteriorly, extending to posterior caecum | Extend from base of siphons to mouth | Extend from base of siphons to mouth |
| Siphons Form        | Separated along entire length, pink pinstriped pigmentation | United to the tip         | United to the tip         |
| Incurrent siphon papillae | Primary row numerous small papillae, secondary row compound papillae | Incurrent siphon numerous large papillae | Incurrent siphon numerous small papillae |
| Excurrent siphon papillae | Numerous small papillae                       | -                         | Excurrent siphon two large papillae on dorsal surface |
| Cephalic hood       | Prominent, covers valve posterior slope        | Inconspicuous, does not cover valve posterior slope | Inconspicuous, does not cover valve posterior slope |
| Cephalic collar     | Present                                       | Absent                    | Absent                    |
| Mantle collar       | Surrounds siphons and pallet stalks            | Absent                    | Absent                    |
| Pallets             | Triangular, cup-shaped, non-segmented, ovate flattened stalk | Paddle shaped, non-segmented, 'thumbnail' depression | Paddle shaped, non-segmented, 'thumbnail' depression, radiating ribs |