Uncovering the shell game with barcodes: diversity of meiofaunal Caecidae snails (Truncatelloidea, Caenogastropoda) from Central America

Christina Egger¹², Timea P. Neusser³, Jon Norenburg⁴, Francesca Leasi⁵, Barbara Buge⁶, Angelo Vannozzi⁷, Regina L. Cunha², Cymon J. Cox², Katharina M. Jörger¹

¹ SNSB-Zoologische Staatssammlung München, Münchhausenstr. 21, 81247 Munich, Germany ² CCMAR, Campus de Gambelas, Universidade do Algarve, 8005-139 Faro, Portugal ³ LMU Munich, Biocenter, Dept. II, Großhaderner Str. 2, 82152 Planegg-Martinsried, Germany ⁴ Department of Invertebrate Zoology, National Museum of Natural History, Smithsonian Institution, Washington, DC 20560, USA ⁵ Department of Biology, Geology and Environmental Science. University of Tennessee at Chattanooga. 615 McCallie Ave. Chattanooga, TN 37403, USA ⁶ Muséum national d’Histoire naturelle, 55 Rue Buffon, 75231 Paris, France ⁷ Independent researcher, Via M.L. Longo 8, Rome, Italy

Corresponding author: Christina Egger (christinaegger@gmx.de)

Abstract

Caecidae is a species-rich family of microsnails with a worldwide distribution. Typical for many groups of gastropods, caecid taxonomy is largely based on overt shell characters. However, identification of species using shell characteristics is problematic due to their rather uniform, tubular shells, the presence of different growth stages, and a high degree of intraspecific variability. In the present study, a first integrative approach to caecid taxonomy is provided using light-microscopic investigation with microsculptural analyses and multi-marker barcoding, in conjunction with molecular species delineation analyses (ABGD, haplotype networks, GMYC, and bPTP). In total 132 specimens of Caecum and Meioceras collected during several sampling trips to Central America were analyzed and delineated into a minimum of 19 species to discuss putative synonyms, and supplement the original descriptions. Molecular phylogenetic analyses suggest Meioceras nitidum and M. cubitatum should be reclassified as Caecum, and the genus Meioceras might present a junior synonym of Caecum. Meiofaunal caecids morphologically resembling C. glabrum from the Northeast Atlantic are a complex of cryptic species with independent evolutionary origins,
likely associated with multiple habitat shifts to the mesopsammic environment. *Caecum invisibile* Egger & Jörger, sp. nov. is formally described based on molecular diagnostic characters. This first integrative approach towards the taxonomy of Caecidae increases the known diversity, reveals the need for a reclassification of the genus *Caecum* and serves as a starting point for a barcoding library of the family, thereby enabling further reliable identifications of these taxonomically challenging microsnails in future studies.

**Keywords**
DNA taxonomy, marine biodiversity, meiofauna, molecular species delineation, Mollusca

**Introduction**

In the past fifteen years molecular barcoding and molecular species delineation have revolutionized the assessment of species diversity and traditional taxonomy, allowing for fast and reproducible species identification and delimitation, and adding to objectivity and reliability in species diagnoses (Leasi et al. 2013; Fontaneto et al. 2015; Scarpa et al. 2016; Martínez-Arce et al. 2020). Molecular data enables testing for morphologically cryptic species as well as phenotypic plasticity, and the evaluation of intra-versus inter-specific variability (Jörger et al. 2012; Leasi et al. 2013, 2016). Given the number of described species and 250 years of taxonomic practice that delimit species based largely on distinct morphologies, it is unsurprising that despite the success of modern molecular approaches, many clades of Metazoa have yet to have their morphological classification tested against molecular markers.

Traditionally, the taxonomy of Gastropoda, one of the most species-rich and better-known clades of invertebrates in the marine environment, is largely based on shell characteristics (Bouchet and Strong 2010). However, this approach is generally problematic as several studies have revealed species exhibiting phenotypic plasticity in shell form due to environmental factors or predation (Trussell 2000; Weigand et al. 2011), and uncovered cryptic species with the aid of molecular data (Haase et al. 2007; Puillandre et al. 2010; Jörger et al. 2012). Consequently, these studies question evolutionary hypotheses based on species delimited by shell characteristics alone and point to the need for an integrative approach using both molecular and morphological data in future research.

Members of the family Caecidae Gray, 1850 can be found in different marine habitats (e.g., among algae or corals) including the marine mesopsammon (i.e., the aqueous interstitial pore spaces of marine sediments). As adults they have uncoiled tubular shells that are likely an adaptation to their infaunal lifestyle (Swedmark 1968). In early descriptions zoologists associated Caecidae snails with tusk-shells (nowadays known as scaphopod molluscs) (see e.g., Montague 1803) or classified them among annelid tube worms (Brown 1827; see Pizzini et al. 2013 for a classificatory history). Even after Caecidae were settled among gastropods (Clark 1849), with current phylogenetic hypotheses placing them among caenogastropod Truncatelloidea (Criscione and Ponder 2013), their unusual tubular shells still posed challenge to taxonomists. Caecid larval shells (protoconch) are usually planispirally coiled with two whorls (Bandel 1996)
and closely resemble related gastropod veliger shells. After settlement of the larvae the adult shell (teleoconch) is formed through differing degrees of uncoiling, with the protoconch either remaining attached (*Panastrophia* de Folin, 1869, *Ctiloceras* R. B. Watson, 1866, *Enigmerces* Iredale & Laseron, 1957, *Jayella* Iredale & Laseron, 1957, *Ponderoceras* Bandel, 1996, *Strebloceras* Carpenter, 1859) or being shed (*Caecum* J. Fleming, 1813, *Meioceras* Carpenter, 1859, *Pizzinia* Vannozzi, 2017, and *Mauroceras* Vannozzi, 2019). In the latter case, the growing teleoconch is closed by a septum (Bandel 1996). The snails continue to shed part of the teleoconch until the fully developed adult shell is formed (Draper 1974). The number of repetitions of shedding likely is variable between species, but unknown for the majority of caecids (Pizzini et al. 1998). This complex shell ontogeny results in highly variable shell morphologies during ontogeny (with a minimum of three different shell morphologies: the larval shell-form, the juvenile shell form(s) and the adult shell form), which hampers species identification and delineation based on single shells if no comparative data is available for the entire morpho-series (i.e., all developmental stages). Moreover, the tubular shells have few taxonomic characters, thus the current taxonomy is largely based on conchological characters such as size, shell shape, ornamentation, construction of the aperture, septum and mucro (i.e., an evagination of the septum, see Fig. 1 for terminology) (Lightfoot 1992a, b, 1993a, b; Pizzini and Raines 2011; Pizzini et al. 2013; Vannozzi et al. 2015; Vannozzi 2017). However, these characters can change for an individual during its lifetime, for instance, young specimens can be entirely smooth and express shell ornamentation only later during maturation, and also shell shape may change as they continue to add shell material at their aperture (i.e., shell opening, see Fig. 1) (Draper 1974; Pizzini 1998b; Lima et al. 2013). Additional difficulties arise in determining whether the septum and mucro are temporary or final (Pizzini et al. 1998).

While the phylogenetic position of the family among truncatelloid gastropods is supported by molecular and morphological data, the taxonomy within the family still is based largely only on shell morphology alone. Indeed, anatomical data is scarce (e.g., Götze 1938; Draper 1974) and thought to offer few diagnostic characters, while molecular barcoding approaches are lacking entirely. Currently, the family Caecidae contains approx. 260 described species in ten genera (MolluscaBase 2019). Most genera (i.e., *Strebloceras*, *Ctiloceras*, *Jayella*, *Enigmerces*, *Ponderoceras*, *Pizzinia*, and *Mauroceras*) are species-poor and limited in distribution to the Indo-Pacific (Iredale and Laseron 1957; Bandel 1996; Pizzini et al. 2013; Vannozzi et al. 2016, 2017). Only *Caecum*, currently with 210 valid species (according to MolluscaBase 2019), shows a circumglobal distribution in temperate and tropical zones. Their abundance is particularly high in tropical waters such as the Indo-Pacific and Central America (Vokes 1983; De Jong and Coomans 1988; Lightfoot 1992a, b, 1993a, b; Díaz Merlano and Puyana Hedegus 1994; Pizzini 1998a; Pizzini and Bonfitto 2008; Discover Life 2020). *Meioceras*, which differs from *Caecum* in the general shape of the shell (i.e., with the widest part towards the middle of the shell), was erected by Carpenter (1858–1859) due to the slightly coiled shape of their juveniles. The genus was recently split into Indo-West Pacific *Mauroceras* and Western Atlantic *Meioceras* (Vannozzi 2019). While recent taxonomic works have de-
scribed the caecid fauna in the Indo-Pacific based on microsculptural investigations of the shell (Pizzini et al. 2013; Vannozzi 2017, 2019), knowledge of caecid diversity in Central American waters is still limited to light-microscopic identification of shells for a large majority of described species.

In this study we present data on caecid diversity based on several recent collecting trips to Central America. We identified the collected Caecidae specimens based on traditional taxonomy and used additional microsculptural observations and molecular barcodes to reliably assign different growth stages to taxa. We applied an integrative experimental approach including multi-marker barcoding and molecular species delineation analyses to test our morphology-based taxonomy, and to identify putative cryptic species.

**Materials and methods**

We collected and microscopically investigated a total of 132 individuals of meiofaunal caecid snails from five different sites in tropical Central America. Of 132 specimens, 67 were selected for further analyses (see Fig. 2 for sampling sites and Tables 1, 2 for details on material and sampling sites). Specimens were extracted from samples of coarse subtidal sands by resting them in buckets for at least 1–2 days to deplete oxygen and accumulate the meiofauna in the surface layer. The surface layer was skimmed off, and the snails extracted by a decantation technique after anesthetization with MgCl₂-seawater solution using a sieve with a mesh size of 100 µm (Jörger et al. 2014). All specimens were documented alive and grouped into preliminary morphotypes based on light microscopic (LM) examination of shell characters in the field and fixed in 75–96% ethanol. Specimens provided by the Muséum national d’Histoire naturelle (MNHN) Paris had previously been removed from their shells in the field by the use of a microwave oven (Galindo et al. 2014), this method is advantageous and recommended over the destructive sampling described below, applied in the beginning of our survey.

**Shell characteristics and microsculptural analyses**

We documented the main taxonomic characters of the tubular shells (Fig. 1), such as the morphology of aperture, septum, and mucro, and measured size and diameter of the shells. Initial species identification in the field was carefully revised in the laboratory, and specimens were assigned to species according to these shell characteristics. The microsculpture of the shell of one representative of each putative morphospecies was investigated via scanning electron microscopy (SEM), whenever a voucher was available (Table 1).

Microscopic debris on the shell was manually removed using an eyelash, and the shell rinsed in 96% ethanol. Specimens were dried by evaporation of the ethanol and transferred onto SEM stubs covered with self-adhesive carbon stickers. We used a sputter coater Polaron SC510 to coat the samples with gold in argon atmosphere. The shells were analyzed with a LEO 1430 VP SEM at a voltage of 15 kV.

All light microscopic images and SEM-micrographs are available through FigShare (https://figshare.com/projects/Central_American_Caecidae/84929).
DNA extraction, amplification, and sequencing

DNA was extracted from 121 of the 132 investigated specimens. The sputter-coated individuals previously investigated by SEM were crushed mechanically using pestles (Bergmeier et al. 2016); specimens investigated only by LM were also crushed if tissue was not already separated. Subsequently, DNA was extracted by the procedure of Knebelsberger and Stöger (2012) combining lysis with 2-mercaptoethanol in CTAB buffer, chloroform-isoamyl precipitation, and recovery using columns with silica-membrane (Nucleo Spin, Macherey-Nagel GmbH & Co. KG, Düren, Germany). The DNA was eluted twice with 25 µl aliquots of pre-heated elution buffer to gain high yield. DNA of specimens deposited to the USMN-Smithsonian Institution were extracted in the Laboratories for Analytical Biology, SI using the standard protocols of the Autogen Prep 956 Extractor (eluting with 100 µl Autogen R9 buffer). Three different markers were partially amplified by PCR: mitochondrial cytochrome oxidase subunit I (COI) and 16S rRNA gene, and nuclear 28S rRNA gene, using the standard PCR primers for gastropods (see Klussmann-Kolb et al. 2008). We either used the Phire polymerase (Thermo Fisher Scientific Inc., Waltham, USA) with the following protocol for PCRs on 16S/COI resp. 28S at the LMU: 98 °C 1 min (98 °C 30 sec; 46–50 °C 20 sec; 72 °C 20 sec) × 36–38 cycles 72 °C 1 min resp. 98 °C 30 sec (98 °C 15 sec; 55–60 °C 5 sec; 72 °C 20 sec) × 35 cycles 72 °C 1 min or the KlenTaq polymerase (AB Peptides, Inc.) with the following program for sequences generated at the SI: 95 °C 3 min, (95 °C 30–45 sec; 48–52 °C 30–45 sec; 72 °C 45–90 sec) × 35–40, 72 °C 7 min. PCR products were either cleaned using a spin column purification kit (Zymo Research, Irvine, California, USA) or were purified with QIAquick (Qiagen Inc.). Samples at the LMU were cycle sequenced on an ABI 3730 48 capillary sequencer (Applied Biosystems, Foster City, USA) using Big Dye 3.1 (Thermo Fisher Scientific Inc.) at the sequencing service of the Ludwig-Maximilians-Universität (LMU) Biocenter, Munich, Germany. At the SI, cycle sequencing was also conducted with BigDye.
Table 1. List of investigated Caecidae specimens, museums numbers (ZSM: SNSB-Bavarian State Collection, MNHN: Muséum National d’Histoire Naturelle, USMN: Smithsonian Institution), and NCBI GenBank accession numbers of sequenced genes and the type of voucher of the material of Caecidae analyzed in the present study. An asterisk (*) marks individuals used for SEM scans.

| Species                  | Field-code | Locality code | Specimen catalog number | Voucher | GenBank number          |
|--------------------------|------------|---------------|--------------------------|---------|-------------------------|
| Caecum imbricatum        | CBC_26     | CBC3          | USNM 1618850             | DNA     | MT727051 MT704281       |
| Caecum imbricatum        | BDT_04     | BRS101        | USNM 1618852             | DNA     | MT704261                |
| Caecum imbricatum        | BDT_07     | BRS103        | USNM 1618854             | DNA     | MT727047                |
| Caecum striatum          | CBC_8      | CBC24         | USNM 1618845             | DNA     | MT727061 MT704275       |
| Caecum invisibile sp. nov. | CBC_1bB   | CBC1b         | ZSM-Mol-20200109         | DNA, paratype | MT727054 MT704267 MT731696 |
| Caecum invisibile sp. nov. | CBC_1bC   | CBC1b         | ZSM-Mol-201000320        | DNA*, holotype | MT727055 MT704268 MT731697 |
| Caecum invisibile sp. nov. | CBC_3a    | CBC1b         | USNM 1618839             | DNA     | MT727056 MT704269 MT731698 |
| Caecum invisibile sp. nov. | CBC_3c    | CBC1b         | USNM 1618840             | DNA     | MT727057 MT704270 MT731699 |
| Caecum invisibile sp. nov. | CBC_3d    | CBC1b         | USNM 1618841             | DNA     | MT727058 MT704271 MT731700 |
| Caecum invisibile sp. nov. | CBC_3e    | CBC1b         | USNM 1618842             | DNA     | MT727059 MT704272 MT731701 |
| Caecum invisibile sp. nov. | CBC_3f    | CBC1b         | USNM 1618843             | DNA     | MT727060 MT704273 MT731702 |
| Caecum invisibile sp. nov. | CBC_13a   | CBC1b         | USNM 1618846             | DNA     | MT727062 MT704276 MT731704 |
| Caecum invisibile sp. nov. | CBC_13b   | CBC1b         | USNM 1618847             | DNA     | MT727063 MT704277 MT731705 |
| Caecum invisibile sp. nov. | CBC_13c   | CBC1b         | USNM 1618848             | DNA     | MT727064 MT704278 MT731706 |
| Caecum invisibile sp. nov. | CBC_13d   | CBC1b         | USNM 1618849             | DNA     | MT727065 MT704279 MT731707 |
| Caecum invisibile sp. nov. | BDT_20    | BRS104        | USNM 1618856             | DNA     | MT727049 MT704264 MT731689 |
| Caecum invisibile sp. nov. | BDT_48    | BRS200        | USNM 1618859             | DNA     | MT727052 MT731694       |
| Caecum regulare           | BDT_22     | BRS108        | USNM 1618883             | DNA     | MT727050 MT731690       |
| Caecum regulare           | CBC_22B    | CBC22         | ZSM-Mol-20100321         | DNA*    | MT704280 MT731708       |
| Caecum donmoorei          | BDT_23     | BRS108        | USNM 1618857             | DNA     | MT704265 MT731691       |
| Caecum donmoorei          | BDT_25     | BRS108        | USNM 1618858             | DNA     | MT704266 MT731692       |
| Caecum donmoorei          | CBC_6      | CBC1b         | USNM 1618844             | DNA     | MT704274 MT731703       |
| MOTU I                    | BDT_17     | ZSM-Mol-20200039 | DNA*                     | MT704263 MT731688 |
| MOTU II                   | BDT_06     | BRS101        | USNM 1618853             | DNA     | MT727046 MT731687       |
| MOTU II                   | BDT_46     | BRS110        | USNM 1618852             | DNA     | MT727051 MT731693       |
| MOTU II                   | BDT_49     | BRS200        | USNM 1618860             | DNA     | MT727053 MT731695       |
| Caecum cf. corrugulatum   | PA_C04     | PA14          | USNM 1618861             | DNA     | MT727069 MT731722       |
| Caecum heptagonum         | PA_28A     | PA23a         | ZSM-Mol-20200030         | DNA*    | MT704283 MT731717       |
| Caecum heptagonum         | PA_G10     | PA23a         | USNM 1618866             | DNA     | MT704291 MT731726       |
| Caecum cf. teres          | PA_E10     | PA23a         | USNM 1618865             | DNA     | MT727070 MT731724       |
| Caecum cf. teres          | PA_30B     | PA23a         | ZSM-Mol-20200033         | DNA*    | MT704284 MT731718       |
| Caecum cf. teres          | PA_30G     | PA23a         | ZSM-Mol-20200037         | DNA*    | MT704286 MT731720       |
| Caecum cf. strangulatum   | PA_G12     | PA23a         | USNM 1618884             | DNA     | MT727072 MT731727       |
Uncovering the shell game with barcodes

chemistry (PerkinElmer) and standard cycles (4 min denaturation at 96 °C, followed by 25 cycles of 10 sec at 96 °C, 5 sec at 50 °C and 4 min at 60 °C), and sequenced on an ABI 3730xl 96-well capillary sequencer. In total, 34%, 43% and 50% of the partial COI, 16S rRNA, and 28S rRNA gene sequences, respectively, were successfully amplified and sequenced. All sequences were edited in Geneious Prime (vers. 11.02.2011, Biomatters, Ltd., Auckland, New Zealand). Primer sequences were removed and base calls checked for misreads against their chromatogram. The sequences were then compared to sequences in the public database NCBI GenBank (http://ncbi.nlm.nih.gov/genbank) by using the BLAST online web service to check for putative contamination. In total 29 COI, 40 16S rRNA and 43 28S rRNA gene sequences were deposited in NCBI GenBank (see Table 1 for accession and voucher numbers).

Phylogenetic analyses

Multiple sequence alignments of the 28S rRNA and COI genes were constructed using Mafft (vers. 7.4.19; Katoh et al. 2002; Nakamura et al. 2018) with default parameter settings. The mitochondrial 16S rRNA sequences were aligned using the program Muscle (vers. 3.8.31; Edgar 2004) with default parameter settings. Alignments were visualized using Seaview (vers. 3.2; Gouy et al. 2009). COI sequences were translated into amino acids. The program Gblocks (vers. 0.91b; Castresana 2000; Talavera and Castresana 2007) was applied to the 16S and 28S rRNA gene alignments to check
Table 2. Details on sampling localities and habitat of the investigated specimens.

| Locality code | Region                   | Station                  | Latitude, Longitude | Depth | Date       | Habitat                                      |
|---------------|--------------------------|--------------------------|---------------------|-------|------------|----------------------------------------------|
| CBC1b         | Carrie Bow Cay, Belize   | House reef               | 16.8015, -88.0790   | 10 m  | 14/01/2010 | open plain                                   |
| CBC3          | Carrie Bow Cay, Belize   | House reef               | 16.8037, -88.0769   | 31 m  | 15/01/2010 | trough inside ridge                          |
| CBC15         | Carrie Bow Cay, Belize   | House reef               | 16.8021, -88.0768   | 31 m  | 22/01/2010 | trough inside ridge                          |
| CBC22         | Carrie Bow Cay, Belize   | Curlew Reef              | 16.7911, -88.0761   | 15 m  | 24/01/2010 | protected sand in patches                    |
| CBC24         | Carrie Bow Cay, Belize   | House reef               | 16.8024, -88.0776   | 19 m  | 25/01/2010 | small sand patches on ridge                  |
| BRS101        | Bocas del Toro, Panama Atlantic | South of Punta Cauro | 9.3609, -82.3467    | 3 m   | 08/06/2010 | small sandy patches, silty, medium coarse sand |
| BRS103        | Bocas del Toro, Panama Atlantic | Solarte Garden | 9.3222, -82.2215    | 4.5 m | 09/06/2010 | exposed, sandy patches, silty, fine          |
| BRS104        | Bocas del Toro, Panama Atlantic | Wild Cane Rock | 9.3503, -82.1723    | 14 m  | 10/06/2010 | deep, sand plain, long ripples, medium coarse sand |
| BRS108        | Bocas del Toro, Panama Atlantic | Near Tiger Rock | 9.2141, -81.9318    | 8.5 m | 10/06/2010 | n/a                                          |
| BRS110        | Bocas del Toro, Panama Atlantic | Wild Cane Reef | 9.3507, -82.1724    | 15 m  | 12/06/2010 | sand plain, medium coarse sand               |
| BRS200        | Bocas del Toro, Panama Atlantic | Wild Cane Reef | 9.3507, -82.1724    | 3 m   | 12/06/2010 | coarse sand 200 µm                           |
| PA4           | Achotines, Panama Pacific | Achotines Bay           | 7.4145, -80.1765    | 2–4 m | 25/02/2016 | sand pits between corals, coarse sand        |
| PA12          | Achotines, Panama Pacific | Back of Achotines Laboratory | 7.4119, -80.1735 | intertidal-subtidal | 28/02/2016 | tide pools, wave action, coarse sand         |
| PA14          | Achotines, Panama Pacific | Isla Iguana south       | 7.6207, -80.0013    | 12 m  | 29/02/2016 | sandy plain around rocks, lots of organic matter, coarse to fine |
| PA15          | Achotines, Panama Pacific | Isla Iguana west        | 7.6301, -80.0022    | 11–16 m | 29/02/2016 | slope with coral rubble, coarse to fine      |
| PA23a         | Achotines, Panama Pacific | Isla Iguana north       | 7.6349, -79.9968    | 10 m  | 06/03/2016 | sand plain, partially with organic matter, gravel and coarse |
| PA23b         | Achotines, Panama Pacific | Isla Iguana north       | 7.6346, -79.9965    | 10 m  | 06/03/2016 | patches next to rocky coral, gravel, course  |
| SL1           | Santa Lucia               | Soufriere Bay           | 13.8494, -61.0675   | 8–9 m | 19/02/2009 |                                             |
| GS32          | Guadeloupe                | west Fajou               | 16.3558, -61.5965   | 2     | 24/05/2012 | lagoon terrace with sandy bottom             |
| GM01          | Guadeloupe                | small marine dead end    | 16.2235, -61.5305   | 1     | 02/05/2012 |                                             |
| AB102         | Martinique               | Anse Noire              | 14.5283, -61.0883   | 6     | 06/09/2016 |                                             |

for unambiguously aligned sites. Proposed exclusion sites were reviewed, adjusted, and subsequently removed (alignments before and after editing are deposited at https://doi.org/10.5281/zenodo.3613958). Sequences available from NCBI GenBank for in-group
taxa (*C. glabrum* (Montagu, 1803) and *C. glabellum* (A. Adams, 1868)), as well as for out-group taxa were added (Table 3). Outgroups were assigned based on the recent molecular phylogenetic analyses by Golding (2014a, b) and Criscione and Ponder (2013).

Two combined data sets were generated: (1) a concatenated alignment of all three marker genes and (2) a concatenated alignment comprising only the mitochondrial 16S rRNA gene, and COI. The data were combined into single matrices using P4 (Foster 2004). The combined data sets were then partitioned by gene and COI codon position.

Maximum likelihood (ML) and Bayesian inference (BI) were used to construct the phylogenetic tree from single genes and from combined and partitioned alignments. For each alignment jModelTest2 (vers. 2.1.10; Darriba et al. 2012) was run and the calculated likelihood scores weighted under the Akaike Information criterion (AICc) (Hurvich and Tsai 1989) which suggested GTR+I+G as the best fitting model. ML was performed using IQ-TREE (multicore vers. 1.6.7.1 for Linux 64-bit; Nguyen et al. 2014) with the GTR+G4+FO model (equivalent to GTR+G in RAxML vers. 8.2; Stamatakis 2014) with 300 bootstrap replicates. Bayesian MCMC analyses were performed using the program MrBayes (vers. v.3.2.6; Huelsenbeck and Ronquist 2001) with the same model. The Bayesian analyses were run in duplicates by default, with each run having four parallel Markov chains (MCMC) to estimate posterior probability support. Each chain was run for 5 million generations, sampling trees every 1000\(^{th}\) generation. Sampled trees were combined into a consensus tree after the first 1000 sampled trees (1000000 generations), considered as ‘burn-in’, were discarded. A general time-reversible model of nucleotide substitutions with a gamma-distribution of among-site rates (GTR+G) was used for the ML analyses. All trees were visualized...
Table 3. List of included Caecidae and outgroup taxa for phylogenetic analyses downloaded from NCBI GenBank (including accession numbers).

| Genus       | Species      | Author        | GenBank number          |
|-------------|--------------|---------------|-------------------------|
| Caecum      | glabrum      | (Montagu, 1803) | FN820514                |
| Caecum      | glabellum    | (A. Adams, 1868) | AB930352, AB930481      |
| Elachorbis  | subtatei     | (Suter, 1907)  | KC110005, KC109953, KC439807 |
| Anemignula  | criscioni    | Golding, 2014  | KC439956, KC439911, KC439788 |
| Pseudomerelina | mahimensis | (Melvill, 1893) | KC439943, KC439894, KC439772 |
| Auricorona  | queenslandica| Golding, 2014  | KC439953, KC439907, KC439786 |
| Nozeha      | topaziaca    | (Hedley, 1908) | KC439952, KC439906, KC439784 |
| Clenchilla  | minutissima  | (Wattebled, 1884) | KC439803, KC109947, KC109999 |
| Calopia     | imitata      | Ponder, 1999  | KC439790, KC439912, KC439957 |
| Calopia     | laseroni     | Ponder, 1999  | KC439792, KC439914, KC439959 |

and annotated using Figtree (vers. 1.4.4; Rambaut 2007). Bootstrap support values (BS) > 85% and posterior probabilities > 0.95 were considered statistically significant.

Species delimitation and characterization based on molecular data

Four different methods of species delineation were used with both the COI and 16S rRNA gene mitochondrial data sets. The Automatic Barcode Gap Discovery (ABGD) webserver was used to partition the data set into putative species based on the calculated gap between intra- and interspecific genetic differences (https://bioinfo.mnhn.fr/abi/public/abgd/abgdweb.html; Puillandre et al. 2012). J ModelTest2 (vers. 2.1.10; Darriba et al. 2012) was applied to the uncorrected COI and 16S rRNA gene alignments and the parameters were weighted under the corrected Akaike Information criterion (AICc) (Hurvich and Tsai 1989). For both alignments the Jukes-Cantor (JC69) as well as Kimura (K80) model showed to be within the 100% confidence interval however, K80 had slightly higher likelihood scores. Both models were applied with the default settings (TS/TV = 2.0, relative gap width = 1.5, Pmin = 0.001, and Pmax = 0.10) on the uncorrected COI and 16S rRNA gene alignments.

To evaluate haplotype connectivity, we generated haplotype networks based on the COI as well as the 16S rRNA gene sequence alignment using the software TCS (vers. 1.21; Clement et al. 2000) using the standard 95% parsimony setting. Ambiguous sites in both alignments were removed to prevent the creation of artificial haplotypes.

The bPTP web server (https://species.h-its.org/) was used to conduct the Bayesian implementation of the PTP model for species delimitation (Zhang et al. 2013) on the optimal ML trees of the individual and the combined COI and 16S rDNA datasets. We applied the default settings with 100000 generations, thinning for each 100th sample with a burn-in of 10% and checked for convergence of the MCMC chains of each run. Posterior probability (PP) support values above 0.95 were considered as strong support.

For the General Mixed Yule-Coalescent model (GMYC) (Pons et al. 2006), ultrametric trees from the COI, 16S rRNA gene, and combined COI and 16S rRNA
gene data were obtained using a time calibrated Bayesian evolutionary analysis in Beast (vers. 1.7.4; Drummond and Rambaut 2007). For the tree prior, we used a Yule process and two fossil records, *Caecum cooperi* and *Caecum imbricatum* [2.58–1.80 myr] (Mansfield 1930; Cooke 1936; Ward and Blackwelder 1987) and the in-group *Caecum* [50–55 myr] (Goedert and Raines 2016) with a lognormal distribution (logL). The analysis was run with the GTR substitution model and under a strict clock assumption. The analysis was started from a random tree and two Markov chains run for 10 000 000 generations with a sampling frequency of 1000. Convergence of the chains was checked in Tracer (vers. 1.7.4.; Rambaut et al. 2018) and effective sampling sizes (ESS) were confirmed as > 200 for all values (Rambaut et al. 2018). The first 10% of sampled trees were removed as burn-in and the trees were combined in TreeAnnotator (vers. 1.7.4.; Drummond et al. 2012) using the maximum clade credibility option and mean node height. The ultrametric trees were uploaded to the web server (https://species.h-its.org/gmyc) for single, as well as, multiple threshold GMYC analyses.

The software QUIDDICH (vers. 1.0.0; Kühn and Haase 2019) was used to identify the diagnostic molecular characters of morphologically cryptic species. We extracted diagnostic characters of type 1 (i.e., characters, which distinguish each individual of the investigated species from other caecids with a fixed character state in the investigated species) and of type 2 (i.e., characters, which distinguish each individual of the investigated species from all other caecids, but vary also within the investigated species) from the COI, 16S rRNA gene and 28S rRNA gene alignments of the same dataset also used for the species delineation and phylogenetic analyses.

**Results**

**Molecular phylogeny and primary species hypothesis**

In our phylogenetic analyses Caecidae form a well-supported clade (1.0 PP, 99% BS; Fig. 3). The two established genera *Caecum* and *Meioceras*, however, are not recovered as reciprocally monophyletic but instead species of *Meioceras* group among *Caecum* species in different parts of the tree (Fig. 3, taxa highlighted in yellow): *M. nitidum* sister to *C. heptagonum* (0.96 PP), and *M. cubitatum* sister to *C. cf. semilaeve* (no statistical support). The phylogeny groups the Caecidae into 21 clades which show moderate to high support values ranging from 0.95 PP/85% BS to full support (Fig. 3). Other clades are only statistically supported by one analysis (*C. regulare*, 92% BS) or do not have statistical support (*C. donmoorei*). The sister group relationships of *C. pulchellum* and *C. regulare* (1.0 PP, 95% BS), and *C. cooperi* and *C. imbricatum* (1.0 PP, 100% BS) are well supported; otherwise, deeper nodes and higher-level relationships among clades are not supported. In agreement with the molecular data, *C. pulchellum* and *C. regulare* as well as *C. cooperi* and *C. imbricatum* show morphological similarities in shell ornamentation and microsculpture. Inconspicuous specimens with smooth shells and few characters that were morphologically ascribable to *C. glabrum* or the American Pa-
cific look-alikes like *C. glabriforme* are polyphyletic and form four lineages separated by branches of comparable length to morphologically distinct species (Fig. 3, highlighted in blue). These lineages are distinct from *C. glabrum* from the North Atlantic (Table 3) included in the analyses, indicating the presence of morphologically cryptic species in this ‘*C. glabrum* species complex’.

**Figure 3.** Optimal ML tree of the concatenated 28S rRNA, 16S rRNA and COI genes partitioned by genes and COI codon positions. Bootstrap values (below nodes) of the ML analysis are shown for values > 80% and posterior probability support (above nodes) of the BI analysis are shown for values > 0.95. Specimens previously classified as *Meioceras* are indicated in yellow color. Smooth, translucent specimens, lacking diagnostic features and summarized in the ‘*Caecum glabrum*-like complex’ are indicated in blue. *C. =* *Caecum, M. =* *Meioceras, MOTU I/ MOTU II =* molecular operational taxonomic unit within the ‘*Caecum glabrum*-like complex’. Figured specimens are all to the same scale. Scale bar: 1 mm.
Molecular species delineation

The methods that were used for species delineation are largely congruent with regard to the assignment of taxa to molecular operational taxonomic units (MOTUs), however individual analyses deviate and evidently differences occur due to incomplete sampling of one of the markers (Fig. 4, Table 1). Both PTP/ bPTP and GMYC (single threshold) delimit 21 MOTUs for the concatenated dataset of COI and 16S rRNA genes (excluding the species whose sequences were retrieved from NCBI GenBank, i.e. North Atlantic *C. glabrum* and Japanese *C. glabellum*). These results are in concordance with the preliminary species hypotheses based on morphological investigation and the molecular phylogenetic tree (Fig. 3) with the exception of additional splits of *C. debile* and *C. regulare* into two distinct MOTUs each. *Caecum cf. teres* resulted in a single species for 16S rDNA alone, and the Bayesian implementation of bPTP split *M. nitidum* into two separate species based on the 16S rRNA genes (however, support value for the split is 0.501%). The multiple threshold analyses in GMYC additionally splits *C. invisibile* sp. nov. of the ‘*C. glabrum* complex’ into two MOTUs, as does TCS but into differing entities. In analyses of individual datasets (numbers not directly comparable due to missing data) ABGD identified 15 MOTUs in our 16S rRNA gene dataset (Fig. 4), while the COI dataset resulted in a hypothesis of 10 MOTUs independent of the application of the JC69 or the K80 model. In comparison to the other methods, TCS appears to oversplit MOTUs (see e.g., TCS analyses of the COI of *C. donmoorei* in Fig. 4). The algorithm of this haplotype-network software splits the 16S rDNA dataset into 19 independent haplotype networks, while it recovered 13 networks for the COI dataset (Fig. 4). Additionally, TCS also splits MOTU II of the ‘*C. glabrum*-like complex’ into two networks and *C. donmoorei* into three independent networks based on 16S rRNA sequence data. Haplotype networks divided *C. cf. teres* and *C. cf. strangulatum* into two unconnected networks. However, the split is not congruent with the two monophyletic sister populations of the species tree (Fig. 4). In summary, we consider only splits relevant, which are supported by at least two different analyses or markers, singular deviating signal might either resemble errors in analyses or might be informative in population analyses (for more details see remarks in Systematics section).

Taxonomy of Central American Caecidae

**Systematics**

**Class Gastropoda** Cuvier, 1797  
**Family Caecidae** Gray, 1850  

**Genus *Meioceras*** Carpenter, 1859  

**Type species.** *Caecum nitidum* Stimpson, 1851 from Florida by subsequent designation, Carpenter (1859): 438.
Based on the molecular phylogeny, specimens identified as *Meioceras nitidum* and *M. cubitatum* both group among *Caecum* species and should therefore be transferred to this genus. However, considering that only one *M. nitidum* is statistically supported, in the interest of taxonomic stability this finding is pending further molecular studies, once additional material is available, preferably including material from the type localities.

**Meioceras nitidum** (Stimpson, 1851)

*Caecum nitidum* Stimpson, 1851 in Stimpson (1851a): 112. Type locality: Florida. *Caecum lermondi* Dall, 1924: 7; *Caecum rotundum* de Folin, 1868: 49, pl. 5, fig. 2; *Meioceras bitumidum* de Folin, 1869: 9, fig. 4; *Meioceras carpenteri* de Folin, 1869: 8, 9, fig. 3; *Meioceras cingulatum* Dall, 1892: 302, pl. 16, figs 6, 7; *Meioceras contractum* de Folin, 1874: 213, t. 2, pl. 4, fig. 7; *Meioceras coxi* de Folin, 1869: 13, fig. 9; *Meioceras crossei* de Folin, 1869: 11, 12, fig. 7; *Meioceras deshayesi* de Folin, 1869: 11, fig. 6; *Meioceras elongatum* de Folin, 1881: 17, pl. 1, fig. 9; *Meioceras fischeri* de Folin, 1870: 188, pl. 26, figsize 3, 4; *Meioceras imiklis* de Folin, 1870: 189, pl. 26, figs 5, 6; *Meioceras leoni* Bérillon, 1874: 251, pl. 5, fig. 7; *Meioceras moreleti* de Folin, 1869: 10, fig. 5; *Meioceras subinflexum* de Folin, 1869: 165, pl. 23, fig. 8; *Meioceras undulosum* de Folin, 1869: 12, fig. 8.

**Material examined.** French Antilles • 1 (Fig. 5A–D); Martinique, Anse Noir; 14.528, -61.088; depth 6 m; 6 Sep 2016; MNHN Madibenthos exped.; Stat. AB102; GenBank: MT704298, MT731714; MNHN-IM-2013-72087a • 1; same collection data as for preceding; GenBank: MT704299, MT731715; MNHN-IM-2013-72087b.

**Shell morphology.** Shell translucent, glossy. Light brown zig-zag pattern covering entire shell in rings with irregular white dorsal patches (Fig. 5A). Bulbous tube, tapering towards aperture and posterior end. Maximum width at about one third of shell length. Slightly more bowed towards aperture. Septum flat, with triangular, pointed mucro (Fig. 5C). No sculpture or microsculpture diagnostic features (Fig. 5D).

**Remarks.** Meioceras and in particular “*M. nitidum*” has a complex taxonomic history involving at present 16 synonyms and several reallocations between *Meioceras* and *Caecum* (MolluscaBase 2019). Vannozzi (2017) highlighted the problems with the ambiguous type specimen of *M. nitidum* (Stimpson 1851a), which encouraged multiple novel descriptions (e.g., Carpenter 1858–1859; De Folin and Périer 1867), nowadays recognized as synonyms. Our two investigated specimens from Martinique are consistent with the meagre original description by Stimpson (1851a) based on specimens from Florida and several redescriptions based on material from the Caribbean and southern America, now all accepted as *M. nitidum* (*M. nitidum* see Bandel 1996: 99, pl. 5, figs 1–7; *M. contractum* see de Folin and Périer 1875: pl. 9, fig. 7). Our specimens differ morphologically from several “*M. nitidum*” specimens from Central American waters all of which were described as a different species in the past but later synonymized with *M. nitidum* acknowledging intraspecific variability (i.e., *M. elongatum*, holotype accessible through the online catalogue of the MNHN (MNHN-
Uncovering the shell game with barcodes

IM-2000-32923) and *M. subinflexum* (see de Folin and Périer 1867: pl. 23, fig. 8). Molecular comparison of specimens spanning the morphological and geographical range is needed to clarify the species status and distribution of the species.

**Meioceras cubitatum** de Folin, 1868

*Meioceras cubitatum* de Folin, 1868 in De Folin and Périer (1867–1871): 50, pl. 5, fig. 4. Type locality: Baie de Bahia [Bahia Bay, Brazil].

Figure 4. Molecular based species delimitation of Central American Caecidae. Guide tree used for PTP and bPTP based on the optimal likelihood tree of the concatenated three-marker dataset. Color codes indicate our preliminary species hypothesis derived from the phylogenetic tree. Color bars reflect the species delimitation suggested by the four consulted species delimitation programs (including ML and Bayesian implementation for PTP and single and multiple threshold for GMYC). Bars are missing where no sequence data obtained.
Figure 5. A–D *Meioceras nitidum*, specimen MNHN-IM-2013-72087  
A light microscopic picture  
B SEM scan  
C SEM close-up of mucro and  
D microsculpture  
E *M. cubitatum*, specimen USNM 1618851  
F, G *C. cf. corrugulatum*, specimen USNM 1618861  
F light microscopic picture  
G light microscopic close-up of microstructure  
H MOTU II, specimen USNM 1618853  
I–L MOTU I, specimen ZSM-Mol-20200039  
I light microscopic picture  
J SEM scan  
K SEM close-up of mucro and  
L microsculpture  
M *C. glabrum*, specimen ZSM-Mol-20200096  
N *C. glabellum* auctt. non Adams, specimen ZSM-Mol-20200074  
O–R *C. invisibile* sp. nov., holotype ZSM-Mol-20100320  
O light microscopic picture  
P SEM scan  
Q SEM close-up of mucro and  
R microsculpture. Scale bars: 10 µm (R); 20 µm (C, K, L, Q); 50 µm (D, E) 100 µm (I, J, O, P); 200 µm (A, B, M, N).
“Caecum cubitatum” (de Folin, 1868): 19; “Meioceras tenerum” de Folin, 1869: 24.

**Material examined.** Belize • 1 (Fig. 5E); Carrie Bow Cay; 16.8021, -88.0767; depth 31 m; 22 Jan 2010; USNM Belize 2010 exped.; Stat. CBC15; DNA voucher; GenBank: MT727067, MT731709; USNM 1618851.

**Shell morphology.** Shell opaque white and solid. Mottled grayish pattern over whole shell, two rows of distinct brown dashes along dorsal side (Fig. 5E). Specimen approx. 2 mm. Tube not evenly curved but appears bulbous and is rounded strongly towards aperture, decreasing towards mucro. Mucro thin and sharp.

**Remarks.** Our molecular phylogenetic results delimited *M. cubitatum* as a separate species, despite similarities to *M. nitidum* in its bulbous shell shape and pattern. Surprisingly, our molecular analyses do not retrieve these morphologically similar *Meioceras* species as a monophyletic entity but suggest independent origin within *Caecum*. Morphological differences towards *M. nitidum* (characterized above) are a more slender shell with more pronounced curvature towards the anterior end and the opaque color of the present individual. We assigned the specimen to *Meioceras cubitatum* sensu de Folin, 1869 from Bahia, Brazil (dos Santos Gomes and Absalão 1996: 523, figs 12–15; De Folin and Périer 1867: pl. 5, fig. 4; Redfern 2001: fig. 178A, B) = *Meioceras cornucopiae* Carpenter, 1859 (from the West Indies, exact type locality unknown) sensu Lima et al. (2015: 3, fig. 7). Nevertheless, this *Meioceras* species likely should also be reallocated to the genus *Caecum* based on the results of our phylogenetic analyses.

**Genus *Caecum* J. Fleming, 1813**

**Type species.** *Dentalium trachea* Montagu, 1803 from England by subsequent designation, Gray 1847: 203.

**Cryptic lineages revealed in molecular analyses**

Twenty-four specimens from Central American waters are smooth and glossy without ornamentation except for occasional growth lines (i.e., possess few shell characteristics), but vary in adult shell length between 0.7 and 2.5 mm (Figs 3, 5F–L, O–R). Morphologically, these specimens all closely resemble *Caecum glabrum* (Montagu, 1803) which is one of the best-known species of caecids, and abundant in the northern Atlantic (Montagu 1803; Wood and Harmer 1848; Götze 1938; Chambers 2009). *Caecum glabrum* was originally described from Biddlesford Bay and Barnstable, Devon, England, the included sequences from GenBank (see Table 3) originates from specimens collected in Norway, but own unpublished data from Roscoff, northern France, supports the wide distribution range of *C. glabrum* along European coastlines based on molecular data. We refer to cryptic species with simple shells lacking characteristic features as ‘*Caecum glabrum*-like’ species complex. In previous works, specimens similar to *C. glabrum* have also been described from the Pacific (*C. glabellum* as *Brochina glabella* A. Adams, 1868 from Akashi, Japan, *C. glabriforme* Carpenter,
1857 and *C. corrugulatum* Carpenter, 1857, both from Mazatlán, Mexico and *C. parvulum* de Folin, 1867 from Panama Bay, Brazil), have been reported and described from Japan, Hawaii, and central America (Carpenter 1855–1857; Adams 1868; Lightfoot 1993b; Pizzini et al. 2007, Takano and Kano 2014). Our study clearly shows an independent evolutionary origin of *C. glabrum* from the northeast Atlantic and *C. glabellum* from Japan, and the cryptic *C. glabrum*-like MOTUs from central America (see Fig. 3, ‘*C. glabrum*-like-complex’ highlighted in blue color). Our molecular species delineation revealed a minimum of four cryptic MOTUs (see above, Fig. 4).

**Caecum cf. corrugulatum** Carpenter, 1857

*Caecum corrugulatum* Carpenter, 1857: 327, pl. 37, figs 375, 1547. Type locality: Mazatlán, 1 sp. off Chama [Mexico].

**Material examined.** Panama • 1 (Fig. 5F, G); Achotines; 7.6207, -80.0013; depth 12 m; 29 Feb 2016; USNM Achotines2016 exped.; Stat. PA14; DNA voucher; GenBank: MT727069, MT731722; USNM 1618861.

**Shell morphology.** Shell color whitish translucent. Tube regularly curved, shape equal in width but bears prominent edge at transition to septum (Fig. 5F). Septum round and blistered lacking a mucro. Aperture equally wide as tube with straight edge. Sculpture appears completely smooth but shows fine concentric ribs at higher magnification (Fig. 5G).

**Remarks.** We assigned the specimen collected in the Pacific coast of Panama to *C. corrugulatum* based on the description of Carpenter (1858–1859) who already highlighted its similarity with another inconspicuous species (*C. glabriforme*). Both species are described from the same geographic area (Mazatlán, Pacific coast of Mexico) and resemble the *C. glabrum*-like type: translucent, blistered septum without mucro, smooth, however slightly bigger than the eponymous *C. glabrum* from European waters. *Caecum corrugulatum*, can be distinguished by microsculptural concentric wrinkles, which could be observed with higher magnification in our specimen. So far, only *C. glabriforme* was recorded in Pacific Panama (Lightfoot 1993b) and recollection at the type locality is needed to 1) confirm the validity of both co-occurring species and reject conspecificity and 2) to confirm their putative distribution range from Mexico to Panama and exclude the possibility of further cryptic species among *C. glabriforme* and *C. corrugulatum* species along the Pacific Coast of Central America (as discovered herein for the Atlantic Coast, see below).

**Caecum invisibile** Egger & Jörger, sp. nov.

http://zoobank.org/4183679F-44F4-4817-A2E1-7325687E5F0A

**Material examined.** Holotype Belize • 1 (Fig. 5O–R); Carrie Bow Cay; 16.8015, -88.0790; depth 10 m; 14 Jan 2010; USNM Belize2010 exped.; Stat. CBC1b; DNA voucher; DNA bank: r462p15f2t91; GenBank: MT727055, MT704268, MT731697; ZSM-Mol-20100320. Paratypes Belize • 1; same data as for holotype; DNA voucher;
Uncovering the shell game with barcodes

DNA bank: r462p14f2t91; GenBank: MT727054, MT704267, MT731696; ZSM-Mol-20200109, Belize • 2; same data as for holotype; ZSM-Mol-20200111, ZSM-Mol-20200112. Other material Belize • 10; same data as for holotype; DNA voucher; GenBank: MT727056–MT727065, MT704269–MT704279, MT731698–MT731707; USNM 1618839, USNM 1618840, USNM 1618841, USNM 1618842, USNM 1618843, USNM 1618846, USNM 1618847, USNM 1618848, USNM 1618849. Panama • 1; Bocas del Toro; 9.2140, -81.9318; depth 8.5 m; 5 Jun 2010; USNM BRS2010 exped.; Stat. BRS104; DNA voucher; GenBank: MT727049, MT704264, MT731689; USNM 1618856. • 1; Bocas del Toro; 9.3507, -82.1724; depth 3 m; 13 Jun 2010; USNM BRS2010 exped.; Stat. BRS200; DNA voucher; GenBank: MT727052, MT731694; USNM 1618859.

Molecular diagnostic characters. see Table 4.

Morphological description. All investigated specimens were very similar in appearance, with little or no variation in shell morphology. Shell completely translucent. Length 0.8 mm long, width 0.2 mm (holotype, Fig. 5O). Tube regularly curved, shape equal in width but bears prominent edge at transition to septum, edge with smaller diameter. Septum round and blistered lacking a distinct mucro. Septum slightly inclining towards the left, dorsal side in holotype with slight variation between the specimens. Aperture equally wide as tube with straight edge. Sculpture appears smooth, only with faint growth lines (Fig. 5R). Whitish translucent body visible through translucent shell. Operculum translucent, slightly tinted yellowish. Radula formula shows taenioglossate pattern 2.1.1.1.2. with very small central rhachidian tooth. Large lateral teeth oriented towards the rhachidian tooth. Marginal teeth finer, outer marginal teeth are scoop-like curved. All the specimens investigated are adults based on the cylindrical shape of the tube and the shape of the aperture showing a reflected lip without cutting edge, which is normally present in immature specimens.

Etymology. The Latin adjective invisibile (invisible, unable to be seen) refers to the minute size of specimens, the translucent color of its shell, its hidden lifestyle between sand grains, and its taxonomic crypsis.

Table 4. Type 1 characters and type 2 characters (Kühn and Haase 2019) for COI, 16S rRNA and 28S rRNA sequence data (no type 2 characters for 16S rRNA and 28S rRNA present) of C. invisibile sp. nov.

| Position | Type 1 | Type 2 | Position | States | Type 1 | Type 2 | Position | States | Type 1 | Type 2 | Position | States |
|----------|--------|--------|----------|--------|--------|--------|----------|--------|--------|--------|----------|--------|
| 15       | A      | 501    | T        | C      | 171    | G      | 450      | G      | 378    | G      | 595      | CA     |
| 171      | G      | 450    | G        | 9      | C      | 32      | T        | 414    | T      | 392    | A       |
| 267      | G      | 93     | T        | 93     | G      | 32      | T        | 426    | C      | 426    | C       |
| 279      | T      | 300    | G        | 97     | T      | 104     | T        | 515    | T      | 515    | T       |
|          |        |        |          | 191    | T      | 612     | G        | 598    | A      | 598    | A       |
|          |        |        |          | 223    | G      | 649     | A        |        |        |        |
|          |        |        |          | 227    | G      | 664     | T        |        |        |        |
|          |        |        |          | 247    | G      |         |          |        |        |        |
Distribution. Type locality: Carrie Bow Cay, Belize. (16.8015°N, -88.0790°W, -10 m). Distributed in Central American Atlantic from Carrie Bow Cay, Belize to Bocas del Toro, Panama. Intertidal in coarse biogenic sediments (calcareous sand and shell hash), shallow subtidal at ten meters’ depth.

Remarks. *Caecum invisibile* sp. nov. is described as a new species based on molecular diagnostic characters, which show it as distinct from the European *C. glabrum* (Fig. 5M), as well as the morphologically similar *C. corrugulatum* (Fig. 5F, G) from the Central American Pacific and *C. glabellum* from Japan (Fig. 5N).

**MOTU I**

**Material examined.** Panamá • 1 (Fig. 5I–L); Bocas del Toro; 2010; USNM BRS2010 exped.; DNA voucher; DNA bank: r462p13f2t91; GenBank: MT704263, MT731688; ZSM-Mol-20200039.

**Morphological characterization.** Shell size 1.3 mm long, 0.3 mm wide. Translucent, with whitish body. Tube regularly curved and equal width. Septum hemispherical (Fig. 5K). Aperture straight, with lip indicating an adult specimen. Operculum brownish. No sculpture or microsculpture diagnostic features (compare Fig. 5L).

**Remarks.** MOTU I is highly similar to the European *C. glabrum* (Fig. 5M) and *Caecum invisibile* sp. nov. (Fig. 5O–R). However, MOTU I shows some small morphological differences such as a bigger shell size and a tiny rim at the aperture (Fig. 5L) which is absent in *C. glabrum*. The septum is further completely round and blistered (Fig. 5K), whereas the one of *Caecum invisibile* sp. nov. slightly inclines (Fig. 5Q). MOTU I is based on a singleton and an incomplete molecular dataset, lacking COI sequence data. Additional material from the same locality is necessary to justify proper species description in future research.

**MOTU II**

**Material examined.** Panamá • 1 (Fig. 5H); Bocas del Toro; 9.4333, -82.347; depth 3 m; 5 Jun 2010; USNM BRS2010 exped.; Stat. BRS101; DNA voucher; GenBank: MT727046, MT704262, MT731687; USNM 1618853. • 1; Bocas del Toro; 9.3507, -82.1724; depth 15 m; 13 Jun 2010; USNM BRS2010 exped.; Stat. BRS110; DNA voucher; GenBank: MT727051, MT731693; USNM 1618852. • 1; Bocas del Toro; 9.3507, -82.1724; depth 3 m; 13 Jun 2010; USNM BRS2010 exped.; Stat. BRS200; DNA voucher; GenBank: MT727053, MT731695; USNM 1618860.

**Morphological characterization.** Shell size unknown. Translucent, with translucent body. Tube regularly curved, slightly increasing in diameter towards aperture. Septum round, slightly flattened (Fig. 5H). Aperture straight. No sculpture visible, microsculptural data missing.

**Remarks.** MOTU II is based on the molecular data of three specimens; however, we unfortunately lack SEM scans and thus microsculptural data of the shell and light
microscopic images are only available for one specimen (Fig. 5H). This specimen is a juvenile, and due to uncertainty with regards to adult ornamentation of the shell, and its possible identity with an already described species, we refrain from providing a formal description based on the available material only.

Adding barcodes to known Central American Caecidae

Caecum heptagonum Carpenter, 1857

Caecum heptagonum Carpenter, 1857: 319, t. 1524. Type locality: Mazatlán [Mexico].

Material examined. Panama • 1 juv. (Fig. 6B–E); Achotines; 7.6349, -79.9968; depth 10 m; 6 Mar 2016; USNM Achotines2016 exped.; Stat. PA23a; DNA voucher; DNA bank: r462p4f2t91, GenBank: MT704283, MT731717; ZSM-Mol-20200030. • 1 juv. (Fig. 6A); same collection data as for preceding; DNA voucher; GenBank: MT704291, MT731726; USNM 1618866. • 1 juv.; same collection data as for preceding; ZSM-Mol-20200116. • 1 juv.; same collection data as for preceding; ZSM-Mol-20200117. • 1 juv.; Achotines; 7.6346, -79.9965; depth 10 m; 6 Mar 2016; USNM Achotines2016 exped.; Stat. PA23b; ZSM-Mol-20200115.

Shell morphology. In juvenile specimens, shell fragile, translucent brownish color. Tube doubles diameter towards aperture, with a moderate curvature in anterior half, increasing distally in curvature. Septum level beneath cutting plane, slightly rising towards mucro (Fig. 6D). Mucro slender finger-like shape (Fig. 6D). Aperture fragile and partly broken. Shell sculptured by seven longitudinal ridges with transverse ribs crossing, knobs at intersections, ridges less prominent towards posterior. Microsculpture of fine rugose longitudinal stripes, noticeably increasing in width on transversal rings in comparison to interspaces (Fig. 6E).

Remarks. Due to the characteristics of the heptagonal tube with the transversal rings, considered unique among caecids (Lightfoot 1993a), the investigated specimen could be unambiguously assigned to C. heptagonum. However, illustrations of C. heptagonum indicate a very thick shell with distinct differentiated aperture with inner bulge forming a round opening instead of the outer polygonal shape (Keen 1974; Pizzini et al. 1998: 142, figs 1–13) including an inner bulge in the aperture, forming a round opening instead of the outer polygonal shape which is absent in the rather thin and fragile investigated specimens. As our samples only comprised juvenile specimen, however, we can attribute this variation to the unfinished shell state.

Caecum imbricatum Carpenter, 1858

Caecum imbricatum Carpenter, 1858: 422, pl. 69, fig. 10. Type locality: “W. Indies [Carribbean].

Caecum coronatum de Folin, 1867: 50–52, pl. 2, fig. 5; Caecum formulosum de Folin, 1869: 24–125, pl. 11, figs 9, 10 (with three varieties paucicostata, simplex and
sulcate); Caecum insigne de Folin, 1867: 52, 53, pl. 2, fig. 4; Caecum sculptum de Folin, 1881: 15, pl. I, figs 1, 2.

**Material examined.** Belize • 1 (Fig. 1, specimen to the left); Carrie Bow Cay; 16.8037, -88.0769; depth 31 m; 15 Jan 2010; USNM Belize2010 exped.; DNA voucher; GenBank: MT727051, MT704281; USNM 1618850. Panama • 1; Bocas del Toro; 9.4333, -82.3467; depth 3 m; 5 Jun 2010; USNM BRS2010 exped.; DNA voucher; GenBank: MT704261; USNM 1618852. • 1; Bocas del Toro; 9.3222, -82.2215; depth 4.5 m; 5 Jun 2010; USNM BRS2010 exped.; DNA voucher; GenBank: MT727047; USNM 1618854. • 1; same collection data as for preceding; GenBank: MT727048; USNM 1618855.

**Shell morphology.** Shell opaque, yellowish. Lighter color in interspaces, darker colored ridges of prominent rhombic pattern (Fig. 1, first specimen from left). Tube evenly narrows towards posterior end, which is approximately half as wide in diameter as aperture. Septum flat, triangular shaped, strongly pointed mucro (Fig. 1, first specimen from left). Rhombic pattern consisting of distinct longitudinal ridges crossed by axial ridges, pattern more distinct towards aperture, last row forms bumps at intersections ((Fig. 1, specimen to the left).

**Remarks.** See remarks on *C. cooperi* “ after paragraph on shell morphology of *C. imbricatum.*

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**Caecum cooperi** S. Smith, 1860

*Caecum cooperi* S. Smith, 1860: 154–155. Type locality: northern part of Gardiner’s Bay, four or five fathoms [United States].

*Caecum costatum* A. E. Verrill, 1872: 283, pl. 6, fig. 6, *Caecum smithi* Cooper, 1872.

**Material examined.** French Antilles • 1 (Fig. 6F–I); Guadeloupe; 16.3558, -61.5965; depth 2 m; 24 May 2012; MNHN KARUBENTHOS exped.; Stat. GS32; GenBank: MT704297, MT731713; MNHN-IM-2019-32.

**Shell morphology.** Shell opaque, with whitish diffuse patterns. Size > 2 mm, tube narrow and elongated, curvature increasing towards posterior end (Fig. 6F, G). Septum flat, triangular shaped, strongly pointed mucro (Fig. 6H) similar to mucro in *C. imbricatum.* Prominent and conspicuous shell ornamentation, minimum of 20 longitudinal strings of beads, more pronounced towards aperture (Fig. 6I).

**Remarks.** Our molecular species delimitation separates *C. cooperi* and *C. imbricatum* into two independent evolving sister species (see Figs 3, 4). This is supported by (minor) morphological differences such as a finer but more pronounced bead-like ornamentation in *C. cooperi* in comparison to flattened squarish and less frequent longitudinal pattern. The included barcodes and distinguishing diagnostic features should help to overcome previous taxonomic uncertainty suggestive of synonymy (see *C. imbricatum* sensu Moore (1972: 888, fig. 6)). The putative synonymy of
Figure 6. A–E *Caecum heptagonum* A specimen USNM 1618866 juvenile specimen with larval shell still attached B–E specimen ZSM-Mol-20200030, juvenile specimen already resembling closely the adult form B light microscopic picture C SEM scan D close-up of mucro and E microsculpture F–I *C. cooperi*, specimen MNHN-IM-2019-32 F light microscopic picture G SEM scan H close-up of mucro and I microsculpture whole specimen and close-up of mucro and microsculpture J–M *C. debile*, specimen MNHN-IM-2019-27 J light microscopic picture K SEM scan L close-up of mucro and M microsculpture N–Q *C. striatum*, specimen ZSM-Mol-20100322 N light microscopic picture O SEM scan P close-up of mucro and Q microsculpture R–U *C. clathratum*, specimen MNHN-IM-2019-17 R light microscopic picture S SEM scan T close-up of mucro and U microsculpture. Scale bars: 10 µm (D, E, I, Q); 20 µm (H, M, P, U); 100 µm (A, B, C, L, N, O, T); 200 µm (F, G, J, K whole specimen); 300 µm (R, S).
**Caecum debile** Verrill & Bush, 1900

*Caecum debile* Verrill & Bush, 1900: 538. Type locality: Bermuda, Ship Channel and Bailey Bay, in 12 to 40 feet.

**Material examined.** French Antilles • 1 (Fig. 6J–M); Guadeloupe; 16.3558, -61.5965; depth 2 m; 24 May 2012; MNHN KARUBENTHOS exped.; Stat. GS32 GenBank: MT704295, MT731711; MNHN-IM-2019-27a • 1; same collection data as for preceding; GenBank: MT704296, MT731712; MNHN-IM-2019-27b.

**Shell morphology.** Color whitish, slightly translucent. Specimens about 2.0 mm long, 0.5 mm wide. Tube of adult specimen 2.1 mm long, 0.5 mm wide. Tube evenly curved and evenly wide over entire length (Fig. 6J, K). Septum separated by sharp rim from tube and hemispherical without protruding mucro (Fig. 6L). Aperture with protruding ring. Shell structured by longitudinal striae clearly observable via LM. Microsculpture of fine wavy striation (Fig. 6M).

**Remarks.** The present specimens were assigned to *C. debile* based on the characteristic microsculpture (see Absalão and Gomes 2001). Morphologically, *C. debile* might present a synonym of *C. infimum*, de Folin, 1867 (de Folin and Périer 1867: 26, pl. 3, fig. 2) but, in awareness of cryptic species we refrain from synonymizing until *C. infimum* from the type locality is available for molecular analyses. Some species delineation analyses, separate *C. debile* into two independent species (Fig. 4), which might indicate a putative speciation, but more data on the genetic variability is needed to exclude the presence of an artefact in analyses.

**Caecum striatum** de Folin, 1868

*Caecum striatum* de Folin, 1868 in De Folin and Périer (1867–1871): 49, pl. 5, fig. 3 (with variety *obsoleta* de Folin, 1874). Type locality: Baie de Bahia [Bahia Bay, Brazil].

**Material examined.** Belize • 1; Carrie Bow Cay; 16.8024, -88.0776; depth 19 m; 25 Jan 2010; USNM Belize2010 exped.; Stat. CBC24; DNA voucher; GenBank: MT727061, MT704275; USNM 1618845. • 1 (Fig. 6N–Q); same collection data as for preceding; ZSM-Mol-20100322.

**Shell morphology.** Shell translucent with mottled ochre and white marbling (Fig. 6N). Size 1.5 mm in length and 0.3 mm in width with thick shell (= 10 µm at aperture). Tube curved regularly with equal width at posterior and anterior end. Blistered, dome-shaped septum with prominent ring separating tube from septum
(Fig. 6P). Mucro central, flat and rounded, pointing slightly dorsal, hardly separated from septum. Aperture simple, straight. Sculpture not visible under the light microscope, i.e., shell appears rather smooth despite two faint transversal rings slightly noticeable close to aperture. SEM examination reveals longitudinal and slightly wavy structure (Fig. 6Q).

Remarks. *Caecum striatum* was identified based on a comparison with the material collected in the sampling region and dedicated as lectotypes by Absalão and Gomes (2001). Their microsculptural description of the type material does correspond to the longitudinal striaion of our investigated specimen. Furthermore, shape, mucro, and the noticeably sharp aperture are identical to our specimen. A comparison with a specimen of *C. striatum* pictured by Pastorino and Chiesa (2014: figs 10–16), also shows the same fine-lined microsculpture as our specimen. Type specimens of three highly similar species, namely *C. johnsoni* Winkley, 1809, *C. antillarum* Carpenter, 1858 and *C. strigosum* de Folin, 1867, are described from Central American waters. Differences can be compared in the reinvestigation of Absalão and Gomes (2001: 20, figs 39–41, 12, figs 11, 12 and figs 7, 8 respectively).

**Caecum cf. clathratum** Carpenter, 1857

*Caecum clathratum* Carpenter, 1857 in Carpenter (1855–1857): 322, pl. 34, figs 269, 1528. Type locality: Mazatlán [Mexico].

**Material examined.** French Antilles • 1 (Fig. 6R–U); Guadeloupe; 16.2235, -61.5305; depth 1 m; 2 May 2012; MNHN KARUBENTHOS exped.; Stat. GM01; GenBank: MT704294, MT731710; MNHN-IM-2019-17.

**Shell morphology.** Large, thick shell (3.0 mm length and 0.8 mm width) with and even curvature (Fig. 6R). Color opaque yellow brownish, entire specimen covered in dense dark periostracum. Septum triangular merged with pointed mucro (Fig. 6T). Aperture oblique and constricted. Shell with 21 strong and protruding sharp ribs and deep interspaces narrowing at aperture (Fig. 6R, S). Ribs and interspaces smooth without microsculptural diagnostic features (Fig. 6U).

**Remarks.** The specimen corresponds to *C. clathratum*, which differs from other ribbed *Caecum* species by its exceptional size, golden color and lack of microsculpture (compare with Lightfoot 1993a: 15, fig. 1 and a syntype collected by Carpenter available through the online catalogue of the Natural History Museum London catalogue number 1857.6.4.1528). However, the specimen is described and known only from the Eastern Pacific. Our herein investigated specimen from the Atlantic might thus present a (morphologically cryptic) sister species new to science, which potentially originated when populations were separated via the formation of the Isthmus of Panama. But molecular data of specimens collected from the Eastern Pacific is required to confirm the molecular identity or justify the description of a new species.
**Caecum pulchellum** Stimpson, 1851

*Caecum pulchellum* Stimpson, 1851 in Stimpson (1851b): 36, pl. 2, fig. 3. Type locality: New England, Buzzard’s Bay [New Bedford Harbor, Massachusetts].

*Caecum capitatum* de Folin, 1874: 227, 228, pl. 9, fig. 8; *Caecum conjunctum* de Folin, 1867: 46, pl. 4, figs 5, 6; *Caecum curtatum* de Folin, 1867: 20, pl. 2, figs 4, 5

**Material examined.** Saint Lucia • 1 (Fig. 7A, C, E, F); Soufriere Bay; 13.8494, -61.0675; depth 8–9 m; 19 Feb 2009; ZSM stuff leg.; DNA voucher; DNA bank: r462p-19f2t91; GenBank: MT727074, MT704300, MT731729; ZSM-Mol-20090485. • 1, juv. (Fig. 7B, D); same collection data as for preceding; DNA voucher; DNA bank: r462p20f2t91; ZSM-Mol-20200118.

**Shell description.** Color opaque whitish, slightly translucent, shell thick (Fig. 7A). Adult specimen 2.1 mm long, 0.5 mm wide. Tube slightly tapering, constricted at aperture, with thickened lip (Figs 7C). Septum slightly lower than posterior end of tube, rising, small and peaked mucro (Fig. 7E). Shell bears 27 squares transverse ribs of even width and equal interspaces, except two or three ribs which meld close to the aperture. Topmost ring sloped towards septum and smaller than others. Fine inconspicuous longitudinal microstriae cover ribs (Fig. 7F). Interspaces covered with organic material, therefore no microsculptural pattern visible.

**Remarks.** Our investigated specimens agree well with recent descriptions and geographical records of *C. pulchellum* (e.g., Bandel (1996): 112, 113, pl. 6, figs 1–4, 6, pl. 7, figs 1, 2 from Columbia and Curacao; Lightfoot (1992a): 143, fig. 2). The type specimen, however, is originally described from Buzzard’s Bay in Massachusetts, USA. A molecular comparison of specimen from the Caribbean and northeast America is needed to confirm the distribution range based on morphology.

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**Caecum regulare** Carpenter, 1858

*Caecum regulare* Carpenter, 1858 in Carpenter (1858–1859): 428–429, pl. 69. Type locality: W. Indies (Woodward) [Caribbean].

**Material examined.** Panama • 1, juv.; Bocas del Toro; 9.2141, -81.9318; depth 8.5 m; 5 Jun 2010; USNM BRS2010 exped.; Stat. BRS108; DNA voucher; GenBank: MT727050, MT731690; USNM 161888. Belize • 1, juv. (Fig. 7H–K); Carrie Bow Cay; 16.7911, -88.0761; depth 15 m; 24 Jan 2010; USNM Belize2010 exped.; Stat. CBC22; DNA voucher; DNA bank: r462p17f2t91; GenBank: MT704280, MT731708; ZSM-Mol-20100321. • 1, juv.; same collection data as for preceding; ZSM-Mol-20200114.

**Shell morphology.** Shell translucent, color white to yellowish (Fig. 7H). Size 1.0 mm long, 0.4 mm wide. Tube curved regularly in moderate angle, tapering towards posterior, widening at aperture, thick lip (10 µm) (Fig. 7H, I). Septum slightly domed. Mucro slender and sharply pointed (Fig. 7J, but tip broken in specimen ZSM-
Mol-20100321). Sculpture consists of 32 marked and rounded transverse ribs, narrowing towards septum, with narrow deep interspaces (Fig. 7I). Microsculpture shows longitudinal fusiform lobes covering the ribs (Fig. 7K).

**Remarks.** We identified the specimens as *C. regulare* by referring to the drawings of Carpenter’s original description (Carpenter 1857; 1858–1859), the syntype material from the Natural History Museum, London, UK accessed through the online catalogue (catalogue numbers 1858.12.9.19, 1858.12.9.20, 1858.12.9.21) and the re-description of Moore (1972): 888, fig. 7. The conspicuous widening of the shell very close to the aperture in our specimen investigated can be interpreted as a character of a sub-adult growth stage (Bandel 1996), thus not contradicting the assignment to *C. regulare*. A literature survey suggests at least one putative synonym of *C. regulare*, resp. *Caecum planum*, de Folin, 1874 (de Folin and Périer 1875: 277, t. 2, pl.10, figs 8, 9).

However, the geographical distribution differs (*C. regulare* was originally described from the Caribbean and *C. planum* from Brazil), and there are no evident diagnostic differences to *C. regulare*. Hence, molecular data is needed for clarification. Further data also is needed for the complementary gene sequences (16S rRNA and COI) for the two investigated specimens. We consequently attribute the split of our two investigated specimens into distinct molecular species to missing data in our analyses.

**Caecum donmoorei** Mitchell-Tapping, 1979

*Caecum donmoorei* Mitchell-Tapping, 1979: 104, 105, figs 21, 22, 31, 32 Type locality: In 5 m of water in Sprat Baz, Water Island, USVI.

**Material examined.** Panama • 1, juv.; Bocas del Toro; 9.2141, -81.9318; depth 8.5 m; 5 Jun 2010; USNM BRS 2010 exp.; Stat. BRS108; DNA voucher; GenBank: MT704265, MT731691; USNM 1618857. • 1; same collection data as for preceding; DNA voucher; GenBank: MT704266, MT731692; USNM 1618858. Belize • 1 (Fig. 7G); Carrie Bow Cay; 16.8015, -88.0790; depth 10 m; 14 Jan 2010; USNM Belize2010 exped.; Stat. CB-C1b; DNA voucher; GenBank: MT704274, MT731703; USNM 1618844.

**Shell morphology.** Shell opaque white and solid. Size large > 2.0 mm. Tube moderately curved and curvature stronger towards aperture (Fig. 7G). Septum blistered with pointed, sharp mucro (Fig. 7G). Aperture surrounded by three narrow thick rings. Sculpture consists of 25 distinct squarish ribs with wide, deep interspaces. No microsculpture available.

**Remarks.** The collected specimen USNM 1618844 closely resembles the description of *C. donmoorei* from the Virgin Islands (Mitchell-Tapping 1979), however microsculptural data for comparison is missing. Our molecular data suggests the species identity among specimens with less separated, strongly flattened rings and almost vanishing interspaces (e.g., in juvenile USNM 1618857). The same has been observed for the confusingly similar species *C. quadratum*, Carpenter (1855–1857: 322, pl. 34, fig. 370). It can also exhibit a considerable variety of shell morphologies (Lightfoot
1993a), however it origins from the Eastern Pacific. And a second highly similar species, *C. regulare* (see above) is clearly distinguished herein based on molecular data.

**Caecum cf. strangulatum de Folin, 1867**

*Caecum strangulatum* de Folin, 1867: 82 (with variety *acuta* de Folin, 1867). Type locality: Iles aux Perles, dans la baie de Panama [Pearl Islands, Panama].

**Material examined.** Panama ♦ 1 juv.; Achotines; 7.6349, -79.9968; depth 10 m; 6 Mar 2016; USNM Achotines2016 exped.; Stat. PA23a; DNA voucher; GenBank: MT727072, MT731727; USNM 1618884. ♦ 1 juv. (Fig. 7O, P, R, T); same collection data as for preceding; ZSM-Mol-20200038. ♦ 1 juv.; Achotines; 7.6207, -80.0013; depth 12 m; 29 Feb 2016; USNM Achotines2016 exped.; Stat. PA14; DNA voucher; GenBank: MT727068, MT704287, MT731721; USNM 1618864.

**Shell morphology.** Shell fragile, color frosted translucent (Fig. 7O). Shape and size the same as for *C. cf. teres* juvenile (Fig. 7M, N). Sculpture however appears rough and annular using light microscopy, growth lines more distinct and wavy (Fig. 7T). Septum flat, mucro narrow, finger-like (Fig. 7R). Aperture fringed as typical for a juvenile, growing specimen (Fig. 7P). Striped microsculpture consists of narrow longitudinal and interrupted emarginations, shifted against each other. Almost identical to *C. cf. teres* (Fig. 7S).

**Remarks.** We assign the examined specimen to *Caecum strangulatum* (in the juvenile form), which was described from Pacific Panama (de Folin 1867) (holotype MNHN-IM-2000-4586, accessed through the online catalogue of the MNHN), due to the narrow mucro and the annular sculpture, which probably can be interpreted as the transition to a ribbed ornamentation in later life stage. The separate species status of *C. strangulatum* and *C. teres* (see below) is not supported via our molecular species delineation (see Fig. 4). Both are sister clades in phylogenetic analyses (see Fig. 3), but the monophyly of both lineages is not reflected on mitochondrial 16S rRNA (Fig. 4). Both species can be distinguished by morphological features (i.e., a noticeably more pronounced shell structure and a slimmer mucro), but share a unique microsculptural pattern (Fig. 7D). We currently lack comparative data on COI to evaluate whether the results in species delineation on 16S rRNA present a case of incomplete lineage sorting in this recent split between the two sister species or both belong to one genetically and morphologically diverse species. Due to this lack of data, we currently refrain from synonymizing the two yet existing species until more data is available to text species boundaries and the degree of intraspecific variability in shell morphology.

**Caecum cf. teres** Carpenter, 1857

*Caecum teres* Carpenter, 1857: 329, pl. 37, figs 378, 1550. Type locality: Mazatlán [Mexico].
**Material examined.** Panama • 1 juv.; Achotines; 7.6349, -79.9968; depth 10 m; 6 Mar 2016; USNM Achotines 2016 exped.; Stat. PA23a; DNA voucher; GenBank: MT727070, MT704289, MT731724; USNM 1618865. • 1 juv. (Fig. 7L); same collection data as for preceding; DNA voucher; DNA bank: r462p7f2t91; GenBank: MT704284, MT731718; ZSM-Mol-20200033. • 1 juv.; same collection data as for preceding; DNA voucher; DNA bank: r462p11f2t91; GenBank: MT704286, MT731720; ZSM-Mol-20200037. • 1 juv. (Fig. 7M, N, Q, S); same collection data as for preceding; ZSM-Mol-20200032. • 1 juv.; Achotines; 7.6207, -80.0013; depth 12 m; 29 Feb 2016; USNM Achotines 2016 exped.; Stat. PA14; ZSM-Mol-20200027.

**Shell morphology.** Thin, fragile shell. Color whitish translucent. Length varies from 1.2 to 1.5 mm. Tube elongated, uniformly cylindrical, narrowing towards posterior (Fig. 7M, N). Septum clearly set off from edge of tube with large, triangular mucro with rounded tip (Fig. 7Q). Aperture round with sharp and thin rim. Sculpture appears smooth using light microscopy except for numerous fine horizontal growth lines. Microsculpture composed of numerous narrow longitudinal stripes consisting of serial fine indentations (Fig. 7S). Stripes of indentations are slightly shifted, when intersecting a growth line.

**Remarks.** Our material from Pacific Panama closely resembles *Caecum teres* (lectotype, NHMUK catalog number 1857.6.4.1550). However, all investigated specimens are juveniles in different growth-stages and identification remains therefore to be confirmed when adult specimens are available for molecular analyses.

**Caecum cf. semilaeve** Carpenter, 1857

*Caeccum semilaeve* Carpenter, 1857: 319, pl. 33, figs 1526. Type locality: Mazatlán [Mexico].

**Material examined.** Panama • 1 juv.; Achotines; 7.6207, -80.0013; depth 12 m; 29 Feb 2016; USNM Achotines 2016 exped.; Stat. PA14; DNA voucher; DNA bank: r462p2f2t91; GenBank: MT704282, MT731716; ZSM-Mol-20200028. • 1 juv. (Fig. 7U–X); Achotines; 7.6349, -79.9968; depth 10 m; 6 Mar 2016; USNM Achotines 2016 exped.; Stat. PA23a; DNA voucher; DNA bank: r462p8f2t91; GenBank: MT704285, MT731719; ZSM-Mol-20200034.

**Shell morphology.** Shell very thin, delicate and highly translucent, glossy (Fig. 7U). Size 1.0 mm long, 0.5 mm wide. Tube gradually narrowing towards posterior end and evenly curved. Septum blistered, entirely below posterior tube end (Fig. 7W). Posterior end fringed and in specimen ZSM-Mol-20200034 still connected partly with mucro indicating a recent shedding of transitional septum. Mucro with elongated, rounded tip, only slightly extending from tube (Fig. 7W). Aperture bordered by very tiny sharp lip, otherwise fragile (Fig. 7X). Shell surface smooth, no sculpture visible apart from regular growth lines. Shell covered by organic layer (periostracum).

**Remarks.** The examined shells all belong to juveniles due to their fragile character and the unfinished aperture. Therefore, it will be critical to reassess these observations
Figure 7. A–F *Caecum pulchellum*  A, C, E, F specimen ZSM-Mol-20090485 B, D juvenile specimen ZSM-Mol-20200118 A, B light microscopic pictures C, D SEM scans E close-up of mucro and F microsculpture  H–K *C. donmoorei*, specimen USNM 1618844 H–K *C. regulare*, specimen ZSM-Mol-20100321, juvenile H light microscopic picture I SEM scans J close-up of mucro and K microsculpture  L–T *C. cf. teres* and *C. cf. strangulatum* L specimen ZSM-Mol-20200033, juvenile specimen with larval shell still attached M, N, Q, S specimen ZSM-Mol-20200032, juvenile M light microscopic picture N SEM scan Q close-up of mucro and S microsculpture O, P, R, T *C. cf. strangulatum*, specimen ZSM-Mol-20200038, juvenile O light microscopic picture P SEM scan R close-up of mucro and T microsculpture  U–X *C. cf. semilaeve*, specimen ZSM-Mol-20200034 U light microscopic picture V SEM scan W close-up of mucro and X microsculpture. Scale bars: 10 µm (F, S, T, W, X); 20 µm (J, K, Q, R); 50 µm (E); 100 µm (H, I, L–P, U, V); 200 µm (A–D).
based on mature shell structures, as sculpturing is known to be variable during development (see e.g., *C. metamorphosicum* S. Lima, Santos & Absalão, 2013 in Lima et al. 2013). The specimens investigated here build and shed transitional septa as described by Pizzini et al. (1998). Specimens that show a similar micro are *Caecum lineicinctum* de Folin, 1880 (compare Absalão and Gomes 2001: 10, figs 1, 2), *C. liratocinctum* Carpenter, 1857; however, both occur in the Western Atlantic (de Folin 1880; Moore 1972; Lightfoot 1992a; Absalão and Gomes 2001). *Caecum semilaeve* is a species described as similar to *C. liratocinctum* (Carpenter 1855–1857) and its type locality is Mazatlán, Mexico, Eastern Pacific, thus with geographic proximity to the localities of our investigated specimens (Achotines, Panama, eastern Pacific). We therefore assign our material to *C. semilaeve*. However, identification remains uncertain without having observed the manifestations of the shell sculpture as described for *C. semilaeve* in later developmental stages (compare syntypes *C. elongatum* var. *semilaeve* NHMUK 1857.6.4.1526).

**Discussion**

**Taxonomic consequences for Caecidae and the fate of Meioceras**

The Caecidae are currently classified in ten genera (MolluscaBase 2019) of which two, *Caecum* and *Meioceras*, can be found in the Central American region. We investigated two species classified as *Meioceras* and 15 *Caecum* species, including one species new to science (*C. invisibile* sp. nov.), and two candidate species (MOTU I and II). Three individuals that were originally assigned to *Meioceras* are resolved among species of the genus *Caecum* in our molecular phylogenetic analyses and, moreover, have independent evolutionary origins within *Caecum* (Fig. 3). Our data confirms the existence of at least two valid *Meioceras* species (i.e., *M. nitidum* and *M. cubitatum*), which however, based on our data should be reassigned genus *Caecum*. For taxonomic stability, we refrain to reallocate these species at present, until the molecular sampling can be expanded and further data is available supporting our initial results. Unfortunately, we lack material of *M. cornucopiae* (the type species by subsequent designation) and of a putative fourth species, *M. tumidissimum*, both described from Brazil (de Folin and Périer 1869). These taxa are needed to settle the debate on the number of species and to clarify the validity of the genus *Meioceras*. Our findings indicate, however, that the more bulbous shell in *Meioceras* when compared to a more tube-like shell in *Caecum* might not justify generic subdivision. Moreover, the diagnostic spiral growth pattern in the larval and juvenile shell of *Meioceras* (Bandel 1996) might not be used for unambiguous discrimination as our phylogeny indicates that such patterns have evolved independently at least two times within the genus *Caecum*. The bulbous adult shell with an oblique constricted aperture was thought to develop from the preceding ontogenesis of a helicoidal shell section, observable at the beginning of the second growth stages (De Folin 1880; Absalão and Pizzini 2002). This diagnosis has been problematic because in the past other species (Absalão and Pizzini 2002) that also express a similar shell-shape had been classified as *Caecum* due to a lack of observations of the juvenile stadia or the lack of the afore-
mentioned growth pattern (e.g., *Caecum ryssotitum*, Bandel 1996). By contrast, species with typical tube-shaped *Caecum*-shells are known to also have curved growth axes (e.g., *C. antillarum* and *C. japonicum* referred to as *C. glabellum* in Bandel 1996: 65, figs 8, 9). Differences between these growth patterns and those of *Meioceras* seem to be negligible. Thus, it remains to be tested whether the remaining two *Meioceras* species form a monophylum separate form *Caecum* and whether alternative diagnostic morphological or molecular characters can be found to justify the generic subdivision of Western Atlantic Caecidae. Alternatively, such studies may confirm that *Meioceras* is a junior synonym of *Caecum*. The present study might also have consequences for the recently established genus *Mauroceras*, which unites Indo-Pacific Caecidae formerly classified as *Meioceras* (Vannozzi 2019). But, in contrast to *Caecum* and Western Atlantic *Meioceras*, which cannot be clearly separated based on variable growth patterns, *Mauroceras* is diagnosed by a planorbid protoconch with a clear sinusigera, which at present justifies its generic status.

**Phylogenetic interpretation of shell morphologies and general insights for shell-based taxonomy**

The taxonomy of Central American Caecidae has been based on macroscopic shell characters and, consequently, type-species are often poorly defined, and has made the established taxonomy prone to multiple descriptions of synonyms and the establishment of ambiguous species-complexes that are typical for many clades of micromolluscs (Golding 2014b). Modern microsculptural analyses have greatly increased the reliability of shell-based taxonomy and the availability of diagnostic characters in the otherwise largely featureless caecid shells (Pizzini et al. 2013; Vannozzi 2017). However, distinct shells based on coarser diagnostic features can have a similar microsculpture (Vannozzi 2017), suggesting that shell microsculpture should be co-evaluated with traditional diagnostic features and, indeed, that it might be especially valuable to discriminate closely related species. In Central American Caecidae the presence of a series of morphologically highly-similar ribbed taxa (i.e., *C. compactum* Carpenter, 1857, *C. quadratum*, *C. clathratum*, *C. gurgulio* Carpenter, 1858, *C. pulchellum*, *C. regulare*, *Caecum planum* de Folin, 1874) with controversial species status and inconsistent synonymization (Moore 1972) are especially problematic. Here we report SEM-based shell microsculpture that can distinguish taxa and justify the independent species status of *C. regulare* and *C. pulchellum* (compare fusiform lobes (Fig. 7C) with fine longitudinal lamellae (Fig. 7A). However, *C. clathratum* does not possess a unique microsculpture and we are lacking SEM data for our investigated specimen of *C. donmoorei*. Nevertheless, molecular species delineation analyses confirm the existence of four genetically distinct species among those Central American ribbed caecids (i.e., *C. pulchellum*, *C. regulare*, *C. donmoorei* and *C. cf. clathratum*), highlighting the value of complementary molecular analyses to detect possible synonyms or confirm the validity of existing species in taxonomically problematic species complexes.

The different growth stages of caecid development present an additional problem for taxonomic circumscription, which cannot be overcome easily by microsculptural
analyses because, the shape and some patterns of ornamentation appear late in development. This often results in the incorrect assignment of different growth stages even at the generic level (Absalão and Pizzini 2002). In consequence, it requires time consuming comparisons of hundreds of shells for reliable species description and identification (Lightfoot 1992a, 1993a), unfeasible in modern times of taxonomic impediment. The molecular analyses presented in this study show that barcoding markers (i.e., partial mitochondrial COI and 16S rRNA genes) are a valuable tool to address the challenges of caecid taxonomy and that molecular species delineation analyses can reliably identify groups of closely related specimens, therewith providing objective data on intraspecific variability of shell characters. Above all, they enable an unambiguous assignment of juvenile forms in different growth stages to their fully developed adult morphologies (see e.g., C. heptagonum in Fig. 5G, C. pulchellum and C. cf. teres in Fig. 7A, D). Based on a purely morphological approach, these juveniles would have remained unidentified and unaccounted for in biodiversity data, and their contribution to caecid diversity would have been lost. However, in some cases, juveniles could not be matched to their adult counterparts using molecular data since we had no adult animals in our sample. These taxa identified by the molecular data could not be named (e.g., Caecum sp. MOTU II). These examples highlight, how the successful identification of juveniles lacking morphological diagnostic features by means of their genetic fingerprints requires an extensive barcode library of Central American Caecidae as a taxonomic reference. The barcodes of the morphospecies investigated here are the first contribution to such a reference library that can help to provide a baseline and enhance future identification. In general, the poor taxonomic coverage of gastropods and marine invertebrates in public molecular databases such as NCBI GenBank has been identified as a major obstacle to making effective use of molecular barcoding approaches (e.g., to assign spawn to adult specimens; Puillandre et al. 2009). Thus, it is hoped that in the future the scientific community will be able to invest more of its financial and personnel capacities in integrative faunistic approaches that strengthen fundamental biodiversity research.

In biodiversity assessment and conservation biology, molecular species delineation has also demonstrated its potential for identifying cryptic species (Bickford et al. 2007; Jörger et al. 2012; Lemer et al. 2014; Leasi et al. 2016). In revealing cryptic taxa, our study indicates that the species diversity of caecids may have been underestimated until now. Unsurprisingly, the cryptic species, which we identified, are those of particularly small, feature-poor, caecids with few diagnostic characters (see Fig. 5C–F). Indeed, our analyses suggest that meiofaunal character-poor caecids (assigned to the ‘Caecum glabrum-like’ species complex) have evolved several times independently from the larger ornamented caecids in the Central American region. The same may have happened in the northern Atlantic C. glabrum and Northwest Pacific C. glabellum Adams, 1868 from Japan. The evolution of a tubular shell marks the origin of Caecidae and likely correlates with a transition to an infaunal lifestyle (e.g., among corals and coral rubble or algae; Bandel 1996). However, interstitial habitats are very variable, differing with regards to the available space between the sand grains which influences the mobility, light intensity and therefore visibility and protection from predators.
In the ‘Caecum glabrum-like’ microsnails, the morphological similarity among taxa (i.e., minute, slim shell, lack of ornamentation and coloration) likely correlates with a habitat shift into the mesopsammon and the consequent habitat restrictions of this special interstitial environment. ‘Regressive evolution’ leading to simplified and highly adapted body plans are typical for the mesopsammon (Swedmark 1968) and consequently the associated meiofauna is prone to cryptic speciation (Jörger et al. 2012; Meyer-Wachsmuth et al. 2014; Leasi et al. 2016).

Conclusions

Our study of Central American Caecidae shows that traditional taxonomic shell characters cannot sufficiently describe the diversity of these microsnails. Microsculptural investigations add valuable additional information for correct taxonomic assignment, species delineation, and the evaluation of gross shell morphological variation within and among species. However, its effectiveness in allocating juvenile growth stages or morphologically rather cryptic species with few diagnostic shell characters into the classificatory system remains limited. This limitation in morphology-based approaches was overcome by integrating genetic barcoding data and molecular species delineation which revealed a complex of cryptic lineages that were potentially associated with a habitat shift from an epibenthic to (temporary) mesopsammic lifestyle among the interstices of sand grains and shell hash. Integrative biodiversity assessments help contribute to a barcoding library of genetic fingerprints of the targeted fauna which enable rapid identification of new samples and is linked to the existing taxonomic history by morphological identification of the voucher specimens. Thus, beyond documenting the shell in microstructural detail, whenever possible a shell voucher should remain intact available for future investigation when novel methods approach. Nevertheless, the vast accumulation of potential synonyms and old names in gastropod taxonomy is problematic, and species need to be taxonomically revised prior to establishing names for newly discovered species. Re-collecting at type localities might not always be feasible for each species, especially when revising large groups with many described species. Additionally, it bears the risk of false identification when cryptic species co-occur at small geographical ranges. However, genetic barcodes have been generated successfully from old mollusk samples in natural history collections – wet material (Jaksch et al. 2016) and dried shells (Der Sarkissian et al. 2017) alike – and hopefully advances in genetic methodology will soon provide cost-efficient and reliable workflows to also adapt them to microsnails as a complement towards ongoing biodiversity studies.

Acknowledgements

CE received funding by the Malacological Society (Early Career Research Grant) and by the Linnaean Society of London and the Systematics Association (Systematics
Uncovering the shell game with barcodes

This study received Portuguese national funds from FCT – Foundation for Science and Technology through project UIDB/04326/2020, and from the operational programmes CRESC Algarve 2020 and COMPETE 2020 through projects EMBRC.PT ALG-01-0145-FEDER-022121 and BIODATA.PT ALG-01-0145-FEDER-022231.

The workgroup of Prof. G. Haszprunar (LMU Munich) is acknowledged for support and critical discussion of the project, Roland Melzer, Enrico Schwabe, Dirk Neu- mann and Bastian Brenzinger (all ZSM) for their help with scanning electron microscopy and collection management. Many thanks to João Brazão and Gianluca De Moro at the Centro de Ciências do Mar (CCMAR) Portugal for their support in molecular phylogenetic analyses.

The SNSB-Zoological State Collection (ZSM), Munich, Germany, the Smithsonian Natural History Museum, Washington D.C., USA and the Muséum national d’Histoire naturelle Paris, France generously contributed specimens, DNA extracts and sequences. Sampling in Belize and Panama was supported by facilities of the Smithsonian Institution and funded by a Smithsonian Institution Marine Science Network award to JLN (Belize 2010, Panama 2010, 2011) and by Global Genome Initiative (GGI) Award (Grant No. GGI-Rolling-2015-020) to FL. For the material contributed by the Muséum national d’Histoire naturelle Paris we thank the crew of the Madibenthos Survey for collecting the material and preserving the specimens. The Madibenthos Survey was spearheaded by the French Marine Protected Areas Agency (now part of the French Agency for Biodiversity), the Regional Directorate for the Environment (DEAL), and the Martinique Water Bureau (ODE), with support from the Directorate of the Sea (DM) and the Martinique Natural Regional Park (PNRM). It was implemented by the Muséum national d’Histoire naturelle (MNHN, Principal Investigator Philippe Bouchet), with funding from the European Regional Development Fund (ERDF), the Territorial Collectivity of Martinique (CTM). Furthermore, we thank the KARUBENTHOS expedition, which was a joint project of Muséum national d’Histoire naturelle (MNHN; Principal Investigator: Philippe Bouchet), the National Park of Guadeloupe, Université des Antilles et de la Guyane (UAG), and Université Pierre et Marie Curie (UPMC), with funding from Fonds Européen de Développement Régional (FEDER) and Port Autonome de la Guadeloupe. For lists of stations, resulting publications, and other documents on the expedition, see http://expeditions.mnhn.fr/campaign/karubenthos2012. We wish to thank two anonymous reviewers for their detailed comments which considerably helped to improve the manuscript.

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39

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