Hyper-Cryptic Marine Meiofauna: Species Complexes in Nemertodermatida

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Abstract

Nemertodermatida are microscopically small, benthic marine worms. Specimens of two nominal species, Sterreria psammicola and Nemertinoides elongatus from 33 locations worldwide were sequenced for three molecular markers. Species delimitation and validation was done using gene trees, haplotype networks and multilocus Bayesian analysis. We found 20 supported species of which nine: Nemertinoides glandulosum n.sp., N. wolfgangi n.sp., Sterreria boucheii n.sp., S. lundini n.sp., S. martindalei n.sp., S. monolithes n.sp., S. papuensis n.sp., S. variabilis n.sp. and S. ylva n.sp., are described including nucleotide-based diagnoses. The distribution patterns indicate transoceanic dispersal in some of the species. Sympatric species were found in many cases. The high level of cryptic diversity in this meiofauna group implies that marine diversity may be higher than previously estimated.

Introduction

More than 70% of the earth’s surface is covered by oceans, and sediment covers most of the ocean floor. Marine infauna thus inhabits one of the earth’s largest ecosystems. Sediment meiofauna is a diverse assemblage with representatives from many animal phyla. Despite the vast size of the marine benthic ecosystem, the marine meiofauna is poorly known, and even in well-studied areas numerous undescribed species exist [1–4]. Nominal meiofauna species are often reported to have cosmopolitan distributions in concordance with the “Everything is Everywhere (EiE)” hypothesis stating that animals below 1 mm body size are easily dispersed. EiE was originally applied to microorganisms [5] and later extended to organisms up to 1 mm size [6,7]. However, species identification of meiofauna requires time-consuming microscope studies, which is often only possible when a specialist brings equipment to the field to examine live specimens. Such detailed taxonomic studies have shown a high level of endemcity for some groups, e.g. Platyhelminthes and Acoela [3], thus contradicting the EiE hypothesis; whereas other groups such as gastrotrichs of the genus Turbellanella seem to conform to a pattern of large distributions [8].

The diversity of the marine worms of the taxon Nemertodermatida that are part of the meiofauna in clean sandy sediments was reviewed by Sterrer [9] who recognized eight broadly circumscribed species with a potential for further subdivision as some of them were known only from few specimens, many of which were incomplete. Morphological identification of nemertodermatid species is complicated by the fact that a large number of specimens are juveniles where the diagnostic reproductive organs cannot be studied. In total Sterrer [9] reported that 229 specimens of nemertodermatids were studied by him since 1964. The nominal species with the largest distribution range was Sterreria psammicola Sterrer, 1970. Sterrer studied 43 specimens of S. psammicola from the North Sea area, the Mediterranean, Caribbean, Australia and Papua New Guinea and considered it “remarkably homogeneous throughout its global distribution range” and regarded Nemertodermatida rubra (Faubel 1976) [10] as its junior synonym. There is, however, some morphological variation in this cosmopolitan species, most apparent in the pigmentation, which can range from non-existent, with the worms appearing glossy silvery, over a narrow, often only anterior, reddish or brownish “spinal stripe” to a more or less uniform bright red colour (Fig. 1a, h). Pigmented and unpigmented specimens have been recorded from the same site, e.g. around the island of Helgoland, North Sea. The nominal species Nemertinoides elongatus Rher, 1907 [11], which is known only from relatively few specimens, is similar in shape to S. psammicola and juvenile specimens of the two species cannot be distinguished although adults differ in reproductive anatomy and the morphology and distribution of epidermal gland cells.
While current taxonomy suggests that *Sterreria psammicola* has a cosmopolitan distribution, there are biological factors that indicate limited dispersal ability and consequently a high degree of endemism in these small interstitial marine worms: none of the known nemertodermatid species have dormant eggs or a planktonic stage; small and fragile juveniles hatch from thin-shelled eggs shortly after they have been deposited [12].

A recent estimate of marine eukaryote biodiversity based mainly on expert opinion concluded that there may be 0.7–1.0 million marine species including the 226 000 currently known nominal species. The proportion of cryptic species remaining to be identified was approximated to range between 11% and 43% of the currently known number [13]. Here we follow Bickford et al. [14] in regarding as cryptic those species that are or have been classified as the same nominal species due to morphological similarity. Some groups, including Nemertodermatida, were considered too poorly known to allow an estimate of the incidence of cryptic species by Appeltans et al. [13]. Costello et al. [15] estimated the number of marine species based on the rate of descriptions of new species using data from the World Register of
Marine Species (WoRMS) and concluded that there are 300 000 marine species. The latter study did not discuss the effect that undetected cryptic species would have on the estimate. Neither of the two studies defined their concept of species, instead a “legacy species concept” based on the numbers entered into WoRMS was used.

Currently, developments in DNA-sequencing technology and bioinformatics are unleashing the potential for broader and deeper sampling of marine biodiversity. Poorly known meiofauna taxa that may exhibit low morphological complexity or be fragile may thus become better known through metagenetic studies such as those by Fonseca et al. [16]. Next-generation sequencing is also likely to reveal additional diversity in the form of cryptic species; see Emerson et al. [17] for an example from soil fauna. Metagenetic studies of diversity will be immensely more valuable when a populated database of sequences from known species with revised taxonomy is available.

Here we aim to test the hypothesis that the nominal species Sterreria psammicola and Nemertinoides elongatus are complexes of cryptic species. Our data is based on 172 specimens that were collected during seven years from 33 different locations (Tab. 1). We sequenced complete or near-complete ribosomal large and small subunit (LSU and SSU) genes and a fragment of the protein coding Histone 3 (H3) gene, and computed separate gene trees under Maximum Likelihood and Bayesian approaches as well as parsimony networks and pairwise distances to generate primary species hypotheses. Clades identified as putative species were tested for genetic isolation using a multilocus Bayesian approach with the software BP&P [18] to generate secondary species hypotheses. Clades with at least three specimens that were supported in at least two of the three gene trees, present as separate haplotype networks under statistical parsimony, had an averaged interspecific pairwise distance at least twice the averaged intraspecific distance, and that were validated by multilocus Bayesian analysis, are formally described and named in this paper. We operate with a species concept in accordance with the “unified species concept” of de Queiroz [19] emphasizing that species are independently evolving lineages that can be diagnosed in a multitude of ways.

Materials and Methods

Permits

Taxa used in this study are interstitial invertebrates, which do not need special sampling permits, as they are not subject to regulations of species protection and are collected within small amounts of sediment.

For sampling around Helgoland, Germany, at Waimanolo, Hawaii, in Norway, Sweden and most of the Mediterranean, no specific or additional sampling permits for the collection of small amounts of marine sediments were required. Geographic coordinates for each site are given in table 1 of the manuscript. A sampling permit for Bermuda was granted by the Department of Conservation Services, Bermuda; the permit for New Caledonia by the Direction de l’environnement, Nouvelle Caledonie. The permit for sampling in the Parco Nazionale dell’Arcipelago di La Maddalena, Sardinia, was granted by the National Park authority. Sampling in Papua New Guinea took place under a permit delivered by the Papua New Guinea Department of Environment and Conservation.

Specimens

Specimens were extracted from sediments using isotonic magnesium chloride solution [20] and identified under a dissecting microscope sometimes in combination with a compound microscope. Specimens were photographed using a compound microscope, if possible equipped with differential interference contrast optics, before fixing in ethanol or RNAlater. Their microscopic size necessitates use of whole specimens for DNA extraction. To ensure a direct link between morphology and gene sequences all type specimens were photographed prior to preservation for DNA extraction and images are deposited as illustrations of the type material, see table 2 for museum and genbank accession numbers.

For the description of the position of morphological characters, a relative scale (U) is used with the anterior tip of the animal corresponding to 0 U and the posterior tip to 100 U [21]. Measurements, however, are difficult to take as animals seldom lie straight and relaxed for a sufficiently long time and in many cases specimens are incomplete, as the worms are fragile.

DNA extraction, amplification and sequencing

DNA was extracted using the Qiagen Micro Tissue Kit. The microscopic size and corresponding low yield of extracted DNA from the specimens as well as the unavailability of prior sequence data severely limited the choice of nucleotide markers. We were able to consistently amplify and sequence rRNA genes as well as the nuclear protein coding Histone 3 gene. The large ribosomal subunit gene was obtained from 168 specimens with an alignment length of 3583 bp, the small ribosomal subunit gene from 166 specimens (1792 bp) and H3 from 166 specimens (328 bp). All markers were amplified and sequenced using several different primer combinations (Tab. 3), and, in the case of SSU, a nested PCR approach.

Sequence editing, alignment (MAFFT [22]), translation into amino acids and checks for open reading frames were performed using the Geneious Pro 7.0.4. software package created by Biomatters available from http://www.geneious.com. The alignments were tested for random similarity with the program Aliscor [23,24] using the default settings. jModelTest v. 2.1.1. [25] analyses were performed for each dataset in order to test the datasets for the use of the proportion of invariable sites (I, propinvar) and the rate variation across sites (G) and to obtain values to set useful priors. Evolutionary neutrality of the coding gene H3 was tested using Tajima’s D calculated with the software MEGA 5 [26]. Saturation of the H3 gene was detected through plotting the uncorrected p-distances versus the phylogenetic distance using an R-script [27].

We chose two other nemertodermatid species, Nemertoderma westbladi, Steinbock 1930 and Meara stichopi, Westblad 1949 as outgroup taxa.

Phylogenetic “species discovery”

The Geneious package (v. 7.0.4.) was also used to calculate pairwise distances between sequences within and between putative species. For this the LSU and SSU alignments were trimmed by eye to 2009 bp and 1502 bp respectively in order to have sequences of similar lengths but keep most of the information. Those specimens represented by less than half of the alignment length were excluded (s. Supplementary table ST1 for details).

Parsimony haplotype networks were computed using the software TCS 1.21 [28] with the reduced and trimmed datasets for LSU (further reduced to 154 specimens and 2009 bp) and SSU. Gaps were considered a fifth state. For relatively fast evolving mitochondrial genes, a 95% threshold has been shown to recover known species reliably [29]. To account for a slower evolutionary rate the connection limit was set to 98% for the LSU and SSU genes; an additional analysis with the connection limit of 90% was performed for the higher resolving Histone 3 dataset.
**Table 1.** List of localities with geographic coordinates, the number of specimens per gene included in this study.

| Country       | Locality                  | Abbr. | Latitude     | Longitude    | H3 | 18S | 28S | present in clades                |
|---------------|---------------------------|-------|--------------|--------------|----|-----|-----|----------------------------------|
| **Ingroup**   |                           |       |              |              |    |     |     |                                  |
| Bermuda       | John Smith’s Bay          | Ber   | 32°19’8.54”N| 64°42’39.51”W| 4  | 4   | 4   | S. variabilis n.sp.              |
| Croatia       | Cherso                    | CC    | 45°5’43.26”N| 14°17’58.14”E| 3  | 8   | 8   | N. elongatus, N. wolfgangi n.sp., S. rubra, S. variabilis n.sp. |
|               | Umag                      | CU    | 45°25’37”N   | 13°31’18”E   | 0  | 1   | 1   | S. psammicola                   |
| France        | Banyuls-sur-Mer           | FB    | 42°28’53.98”N| 3°7’56.55”E  | 7  | 8   | 8   | N4, N. elongatus, N. wolfgangi n.sp., N. elongatus n.sp., S. lundini n.sp. |
| Germany       | Helgoland; Nordostmauer  | GHn   | 54°11’19.32”N| 7°53’6.83”E  | 3  | 6   | 6   | S. rubra                        |
|               | Helgoland; Tonne 2        | GHT   | 54°10’48.63”N| 7°55’55.49”E| 3  | 4   | 4   | N. elongatus, N. glandulosum n.sp., S. lundini n.sp. |
| Italy         | Acireale                  | IAc   | 37°36’13.47”N| 15°10’40.56”E| 0  | 1   | 1   | S3                              |
|               | Agnone                    | IAg   | 37°18’37.09”N| 15°6’21.03”E | 2  | 2   | 3   | N. wolfgangi n.sp., S. lundini n.sp. |
|               | Budelli Island            | IB    | 41°17’36.21”N| 9°21’39.65”E | 0  | 2   | 3   | S. rubra, S. lundini n.sp.      |
|               | Castello                  | ICa   | 42°45’5”N    | 10°52’2”    | 3  | 3   | 3   | N. elongatus, N. glandulosum n.sp., S. lundini n.sp. |
|               | Formica                   | IF    | 42°34’18.48”N| 10°53’4.92”E| 3  | 4   | 5   | S. rubra                        |
|               | Ischia                    | II    | 40°43’52.06”N| 13°57’46.72”E| 1  | 1   | 1   | S. rubra                        |
|               | La Maddalena              | IS    | 41°16’50.88”N| 9°19’14.52”E| 3  | 11  | 11  | S. rubra, S. lundini n.sp., S. variabilis n.sp. |
|               | La Maddalena cave         | ISc   | 41°13’30.76”N| 9°22’35.36”E| 1  | 2   | 2   | S. rubra                        |
|               | Marchiato                 | IM    | 42°48’0.4”N  | 10°44’6.8”E  | 1  | 4   | 4   | S. rubra, S. lundini n.sp., S. psammicola |
|               | Mimare                    | AM    | 47°42’37.00”N| 13°42’48.48”E| 0  | 2   | 1   | S. rubra, S. psammicola         |
|               | Torre Civette             | IC    | 42°51’17.71”N| 10°46’23.90”E| 5  | 16  | 16  | N2, N. elongatus, N. glandulosum n.sp., S. rubra, S. lundini n.sp., S. psammicola |
|               | Castiglione della Pescaia| IR    | 42°45’58.98”N| 10°51’16.99”E| 4  | 7   | 7   | N1, N3, N. glandulosum n.sp., S. rubra, S. lundini n.sp. |
|               | Punta Ala                 | IW    | 42°48’24.87”N| 10°44’34.37”E| 10 | 15  | 15  | N. elongatus, N. wolfgangi n.sp., S. rubra, S. lundini n.sp. |
| New Caledonia | Amédee                    | NCA   | 22°28’39.58”S| 166°28’21.54”E| 1  | 1   | 1   | S. variabilis n.sp.              |
|               | Poe Beach                 | NCP   | 21°30’50.72”S| 165°23’46.82”E| 3  | 3   | 2   | S. variabilis n.sp.              |
| Papua New Guinea | Sir Island             | PNGS  | 05°11’11.94”S| 145°48’15.12”E| 1  | 1   | 1   | S. papuensis n.sp.               |
|               | Tab Island                | PNGT  | 05°10’16.84”S| 145°50’18.29”E| 1  | 3   | 3   | S. papuensis n.sp.               |
|               | Panab Island              | PNGP  | 05°10’18”S  | 145°48’29”E | 6  | 7   | 7   | S. papuensis n.sp., S. monolithes n.sp., S. boucheii n.sp., P3 |
|               | Wanad Island              | PNGW  | 05°08’07”S  | 145°49’16”E | 11 | 13  | 13  | S. papuensis n.sp., S. monolithes n.sp., S. boucheii n.sp., S. variabilis n.sp. |
| Portugal      | Faro                      | PF    | 36°57’32.1”N | 3°7’53.78”E  | 9  | 11  | 12  | N. elongatus, N. glandulosum n.sp., S. rubra, S2, S. psammicola, S. variabilis n.sp. |
|               | Ilha da Culatra           | PC    | 36°58’55.2”N | 7°52’1.2”E   | 2  | 3   | 3   | S. rubra                        |
| Sweden        | Grisbôdarna              | SG    | 58°55’22.15”N| 10°49’48.79”E| 2  | 3   | 3   | N. elongatus, S. rubra           |
|               | Kallgrund                 | SK    | 58°55’22.94”N| 11°2’42.86”E | 4  | 5   | 5   | N. elongatus, S2                 |
| USA, Hawaii   | Waimanalo                 | H     | 21°19’35.68”N| 157°40’57.93”W| 8  | 8   | 8   | S. martindei n.sp., S. ylvae n.sp., S. variabilis n.sp. |

**Note:** The table provides a comprehensive list of localities with geographic coordinates, the number of specimens per gene included in this study, along with the presence of specific nemertodermatid species.
Maximum Likelihood (ML) and bootstrap support calculations were performed by raxmlGUI [30] using the GTR+G+I evolutionary model and the rapid bootstrap algorithm with 1000 bootstrap reiterations.

Bayesian analyses were performed using the program MrBayes 3.2.1. [31]. No evolutionary model was set and the program was allowed to sample the entire model space of the GTR model by defining nst = mixed. The proportion of invariable sites and G were applied with the prior set to shapepr = Uniform(0.05,1.00) for SSU and LSU and shapepr = Uniform(0.05,2.00) for H3; the pinvarpr was left at the default. Analyses were stopped when the standard deviation of split frequencies was between 0.01 and 0.05, indicating sufficient convergence and a relative burn-in of 25% was used. No concatenated analyses for all three genes combined were conducted. This would conceal incongruences between the gene trees and therefore possibly lead to subsequent errors in the validation of species using BP&P [32].

Trees were visualized using FigTree v1.3.1. [33]. Alignments and tree-files are deposited with Treebase (http://purl.org/phylo/treebase/phylows/study/TB2:S15809).

“Species” validation

Those clades that consisted of at least three specimens, showed an averaged interspecific pairwise distance at least two times higher than the intraspecific averaged pairwise distance (relative threshold distance [34]), formed separate parsimony networks and were present in at least two of the three gene trees, were tested using a multilocus Bayesian approach with the program BP&P to generate secondary species hypotheses [18,35], species validation sensu [36]. The program relies on a user-defined tree and only tests for the presence of nodes in the input-tree; the input of an incorrect guide tree will corrupt the results [32]. In order to create unambiguous input trees the dataset was divided into three subsets and the putative species Sterreria martindalei n.sp. and Sterreria papuensis n.sp. were excluded (different colours in Fig. 2, 3) because the gene trees could not resolve all deeper nodes with high support. Both excluded species, however, are highly supported in all species discovery methods, thus we think that further validation in these cases was not necessary. The subgroups within Sterreria variabilis n.sp. were not validated because of the unresolved topology (polytomies) of the group. Two analyses with the gamma priors set to G(1, 100) and G(1, 1000) for the population size $h$ and G(1, 1000) for the root age $t$ were conducted while the other divergence time parameters are assigned the Dirichlet prior ([18]: equation 2). An additional analysis with an older root age with the G of $h$ (1, 100) and the G of $t$ (1, 10000) was also conducted.

The species we describe are diagnosed based on unique differences in the nucleotide sequences following Jo¨rger and Schrödl [4] in addition to morphological diagnostic characters, which are provided where available.

Nomenclatural acts

The electronic edition of this article conforms to the requirements of the amended International Code of Zoological Nomenclature, and hence the new names contained herein are available under that Code from the electronic edition of this article. This published work and the nomenclatural acts it contains have been registered in ZooBank, the online registration system for the ICZN. The ZooBank LSIDs (Life Science Identifiers) can be resolved and the associated information viewed through any standard web browser by appending the LSID to the prefix “http://zoobank.org/”. The LSID for this publication is: urn:lsid:zoobank.org:pub:A306F670-B4B4-4376-A859-48A9735E1593. LSIDs for new species are given...
**Table 2.** List of all individuals used in this study sorted by clade, with Zoobank Life Science Identifiers (LSID) where applicable, connecting collection code (used in the scratchpads database for Acoela and Nemertodermatida at http://acoela.myspecies.info/), genbank accession numbers per gene and the museum collection numbers for type material.

| species/clade | ZooBank LSID | collection code | SMNH type number | Genbank accession number |
|---------------|--------------|----------------|-----------------|-------------------------|
| N. elongatus   |              |               |                 |                         |
|               | 07–010       | KM062712      |                 | KM062546               |
|               | 07–011       | KM062713      | KM062547        | KM194610               |
|               | 07–013       | KM062714      | KM062548        | KM194612               |
|               | 07–030       | KM062716      | KM062550        | KM194614               |
|               | 07–040       | KM062719      | KM062553        | KM194616               |
|               | 07–051       | KM062720      | KM062554        | KM194617               |
|               | 07–074       | KM062722      | KM062556        | KM194618               |
|               | 07–076       | KM062723      | KM062557        | KM194619               |
|               | 07–078       | KM062724      | KM062558        | KM194620               |
|               | 08–090       | KM062728      | KM062563        |                         |
|               | 08–110       | KM062740      | KM062575        |                         |
|               | 08–120       | KM062745      | KM062580        |                         |
|               | 09–001       | KM062749      | KM062584        | KM194635               |
|               | 11–143       | KM062799      | KM062633        | KM194665               |
|               | 13–170       | KM062814      | KM062648        | KM194676               |
|               | 13–176       | KM062815      | KM062652        |                         |
|               | 13–180       | KM062816      | KM062649        | KM194677               |
|               | 13–441       | KM062824      | KM062657        |                         |
|               | 13–442       | KM062825      | KM062658        | KM194683               |
|               | 13–446       | KM062826      | KM062659        |                         |
|               | MCG04        | KM062834      | KM062661        | KM194684               |
| N. glandulosum | urn:lsid:zoobank.org:act:DFBD9E91-83E2-4567-91ED-BF279F16CB24 | 07–001 | KM062705 | KM062539 | KM194607 |
|               |              | KM062706      | KM062540        | KM194608               |
|               |              | KM062707      | KM062541        |                         |
|               |              | KM062709      | KM062543        | KM194609               |
|               |              | KM062741      | KM062576        | KM194632               |
|               |              | KM062747      | KM062582        | KM194634               |
|               |              | KM062792      | KM062626        | KM194660               |
|               |              | KM062793      | KM062627        | KM194661               |
|               |              | KM062817      | KM062650        | KM194678               |
|               |              | KM062819      | KM062652        | KM194679               |
|               |              | MCG05         | KM062668        | KM194685               |
|               |              | MCG07         | KM062670        | KM194687               |
|               |              | MCG08         | KM062671        | KM194688               |
| N. wolfgangi  | urn:lsid:zoobank.org:act:CC4AC7FC-SCAD-4DD0-9C0E-039390D11356 | 09–041 | KM062757 | KM194640 |
|               |              | KM062732      | KM062567        | KM194627               |
|               |              | KM062733      | KM062568        | KM194628               |
|               |              | KM062739      | KM062574        | KM194631               |
|               |              | KM062763      | KM062597        | KM194642               |
|               |              | 13–453        | 8632            | KM062828               |
|               |              | MCG10         | KM062673        | KM194690               |
|               |              | MCG13         | KM062843        | KM194691               |
|               |              | MCG15         | KM062845        | KM194692               |
|               | MCG08        | KM062838      | KM062671        | KM194688               |
|               | N1           | 08–102        | KM062736        | KM062571               |
Table 2. Cont.

| species/clade | ZooBank LSID | collection code | SMNH type number | Genbank accession number |
|---------------|-------------|----------------|------------------|-------------------------|
|               |             |                |                  | LSU         | SSU         | H3           |
| N2            |             | 08–121         | KM062746         | KM062581    |
|               |             | 08–123         | KM062748         | KM062583    |
| N3            |             | 08–098         | KM062734         | KM062569    | KM194629    |
|               |             | 08–100         | KM062735         | KM062570    | KM194630    |
| N4            | MCG06       | KM062836       | KM062669         | KM194686    |
|               | MCG09       | KM062839       | KM062672         | KM194689    |
| P3            | PNG60       | KM062858       | KM062690         | KM194699    |
|               | PNG61       |                 | KM194700         |             |
| S. boucheti n.sp. | urn:lsid:zoobank.org:act:65760DAD-F39F-4B29-9539-F091D45774FA | PNG70 | KM062863 | KM062695 | KM194705 |
|               |             | PNG54          | KM062854         | KM062686    | KM194697    |
|               |             | PNG68          | KM062862         | KM062694    | KM194704    |
|               |             | PNG72          | KM062864         | KM062696    |             |
|               |             | PNG75          | 8633             | KM062866    | KM194707    |
|               |             | PNG83          | KM062869         | KM062701    | KM194709    |
|               |             | PNG87          | KM062872         | KM062704    | KM194712    |
| S. lundini n.sp. | urn:lsid:zoobank.org:act:F05F5C93-D3C5-4AEA-969D-F1AD2ADE8C20 | 08–093 | KM062730 | KM062565 | KM194626 |
|               |             | 08–094         | KM062731         | KM062566    |             |
|               |             | 08–117         | 8634             | KM062743    | KM062578    | KM194633    |
|               |             | 09–013         | KM062753         | KM062588    |             |
|               |             | 09–035         | KM062756         | KM062591    | KM194639    |
|               |             | 09–053         | KM062761         | KM062595    | KM194641    |
|               |             | 10–076         | KM062779         | KM062613    |             |
|               |             | 10–110         | KM062784         | KM062618    |             |
|               |             | 11–073         | KM062794         | KM062628    |             |
|               |             | MCG01          | KM062831         | KM062665    |             |
|               |             | MCG03          | KM062833         | KM062667    |             |
|               |             | MCG11          | KM062841         | KM062674    |             |
|               |             | MCG14          | KM062844         | KM062677    |             |
| S. martindalei n.sp. | urn:lsid:zoobank.org:act:AD07EBF4-F151-4139-A3FC-88B548E488D6 | 10–055 | KM062771 | KM062605 | KM194648 |
|               |             | 10–056         | 8635             | KM062772    | KM062606    | KM194649    |
|               |             | 10–060         | KM062774         | KM062608    | KM194651    | |
| S. monolithes n.sp. | urn:lsid:zoobank.org:act:638DA2C2-4120-4270-8442-C8D857ED78F6 | PNG57 | KM062856 | KM062688 | KM194698 |
|               |             | PNG84          | 8636             | KM062870    | KM062702    | KM194710    |
|               |             | PNG85          | KM062871         | KM062703    | KM194711    | |
| S. papuensis n.sp. | urn:lsid:zoobank.org:act:B5470A6B-3F8F-432A-84A0-5B980E9469A | PNG48 | KM062849 | KM062681 | KM194694 |
|               |             | PNG49          | KM062850         | KM062682    |             |
|               |             | PNG50          | KM062851         | KM062683    | KM194695    |
|               |             | PNG51          | KM062852         | KM062684    |             |
|               |             | PNG52          | KM062853         | KM062685    | KM194696    |
|               |             | PNG56          | KM062855         | KM062687    |             |
|               |             | PNG58          | KM062857         | KM062689    |             |
|               |             | PNG62          | KM062859         | KM062691    | KM194701    |
|               |             | PNG66          | KM062860         | KM062692    | KM194702    |
|               |             | PNG77          | 8637             | KM062868    | KM062700    |
| species/clade | ZooBank LSID | collection code | SMNH type number | Genbank accession number |
|--------------|-------------|-----------------|------------------|-------------------------|
| S. psammicola|             | 07-006          |                  | KM062708 KM062542       |
|              |             | 09-012          |                  | KM062752 KM062587       |
|              |             | 13-155          |                  | KM062810 KM062644 KM194673 |
|              |             | 13-186          |                  | KM062820 KM062653 KM194680 |
|              |             | 13-483          |                  | KM062829 KM062662       |
|              |             | 13-508          | 8640             | KM062830 KM062663       |
|              | S. rubra    | 07-008          |                  | KM062710 KM062544       |
|              |             | 07-009          |                  | KM062711 KM062545       |
|              |             | 07-031          |                  | KM062717 KM062551       |
|              |             | 08-092          |                  | KM062729 KM062564 KM194625 |
|              |             | 08-103          |                  | KM062737 KM062572       |
|              |             | 08-116          |                  | KM062742 KM062577       |
|              |             | 08-118          |                  | KM062744 KM062579       |
|              |             | 09-002          |                  | KM062750 KM062585 KM194636 |
|              |             | 09-005          |                  | KM062751 KM062586 KM194637 |
|              |             | 09-028          |                  | KM062754 KM062589 KM194638 |
|              |             | 09-029          |                  | KM062755 KM062590       |
|              |             | 09-049          |                  | KM062758 KM062592       |
|              |             | 09-051          |                  | KM062759 KM062593       |
|              |             | 09-052          |                  | KM062760 KM062594       |
|              |             | 09-054          |                  | KM062762 KM062596       |
|              |             | 09-059          |                  | KM062764 KM062598       |
|              |             | 09-060          |                  | KM062765 KM062599 KM194643 |
|              |             | 09-061          |                  | KM062766 KM062600       |
|              |             | 10-073          |                  | KM062776 KM062610       |
|              |             | 10-074          |                  | KM062777 KM062611       |
|              |             | 10-075          |                  | KM062778 KM062612 KM194653 |
|              |             | 10-090          |                  | KM062780 KM062614       |
|              |             | 10-092          |                  | KM062781 KM062615 KM194654 |
|              |             | 10-093          |                  | KM062782 KM062616       |
|              |             | 10-098          |                  | KM062783 KM062617       |
|              |             | 10-117          |                  | KM062785 KM062619       |
|              |             | 10-184          |                  | KM062787 KM062621 KM194656 |
|              |             | 10-188          |                  | KM062788 KM062622       |
|              |             | 10-247          |                  | KM062789 KM062623 KM194657 |
|              |             | 11-139          |                  | KM062795 KM062629 KM194662 |
|              |             | 11-140          |                  | KM062796 KM062630 KM194663 |
|              |             | 11-141          |                  | KM062797 KM062631 KM194664 |
|              |             | 11-142          |                  | KM062798 KM062632       |
|              |             | 11-144          |                  | KM062800 KM062634       |
|              |             | 11-184          |                  | KM062801 KM062635       |
|              |             | 13-094          |                  | KM062806 KM062640 KM194670 |
|              |             | 13-096          |                  | KM062807 KM062641 KM194671 |
|              |             | 13-097          |                  | KM062808 KM062642       |
|              |             | 13-148          |                  | KM062809 KM062643 KM194672 |
|              |             | 13-158          |                  | KM062813 KM062647       |
|              |             | 13-182          |                  | KM062818 KM062651       |
in table 2. The electronic edition of this work was published in a journal with an ISSN, and has been archived and is available from the following digital repositories: PubMed Central, LOCKSS and DiVA [http://www.diva-portal.org/smash/search.jsf].

### Results

When testing for random similarity between sequences, Aliscore highlighted 419 of 3583 aligned sites in the LSU dataset and 110 of 1792 sites in the SSU dataset. No random similarities were indicated in the H3 dataset. Consensus and best trees resulting from analyses of the original and Aliscore-filtered alignments had
identical topologies with the exception of four specimens of Sterreria rubra that grouped with the specimen S7 in the Aliscore pruned LSU analysis. There were small differences in branch support when comparing original and filtered alignments (greatest difference 6% BS in the LSU dataset and 55% in the SSU dataset, but generally no or less than 10% BS difference). The gene trees shown as supplementary data are based on the original alignments (Figures S2–7).

A summary of the results in terms of putative species is given in figure 2, which shows a 75% majority rule consensus tree (MF75) of the LSU ML analysis performed with RAxML. Support (bootstrap for ML analyses, posterior probabilities for Bayesian analyses) for the 20 putative species (excluding outgroups) is shown to the right of the node (or in case of a few long branches over these) in the order LSU/SSU/H3 (for all gene trees see Figures S2–7). The colours refer to the subsets used for species validation (green: Nemertinoides-group, red: European Sterreria-group, blue: extra-European Sterreria-group, orange: untested Sterreria species).

In the uncorrected pairwise distances matrix several groups with at least twice the intraspecific distance to their sister group could be identified (Tables S2–4). The LSU and SSU gene datasets each had 19 distinct putative species groups (excluding outgroups), and in the Histone 3 gene dataset we found 28 such groups. The respective distances never exceeded 0.8% and 0.5% respectively in the LSU and SSU data partitions, with the exception of N4 with pairwise distance 0.2%. In the H3 partition intraspecific distances reached 9.9% in Sterreria variabilis n.sp. Of the 27 groups in the H3 data, 15 correspond to the same putative species as seen in the LSU and SSU gene datasets (Table 4). Eight groups in the H3 dataset did not correspond to putative species supported by the other two genes. This may be a saturation artefact (Figure S1).

The TCS software defined different numbers of parsimony haplotype networks for each of the three loci. Analyses of the LSU, SSU and H3 gene datasets with a connection limit of 90% found 25, 20 and 42 networks respectively (excluding outgroups, Fig. 3a, b). When the Histone 3 gene analysis was relaxed with a connection limit of 90% only 30 networks were found (Fig. 3c). In the LSU dataset N. glandulosum n.sp., S. papuensis n.sp., S. psammicola and S. variabilis were recovered as two and three separate networks respectively. Putative species S7 was excluded from the dataset due to its short sequence. In the SSU analysis, S. boucheti n.sp. and S. ylvae n.sp. were recovered as one network with two steps between the two species. One specimen of the diverse S. variabilis n.sp. formed a separate network not connected to the other specimens of the species. The H3 gene analyses split S. lundini n.sp. and S. papuensis n.sp. into two networks each and N. elongatus into three different networks. S. rubra was recovered in seven networks most of them consisting of only one or two specimens, corresponding with the observed pairwise distances. S. variabilis n.sp. formed five networks and one network connecting with S. boucheti n.sp. S. ylvae n.sp. and S. monolithes n.sp. formed one network connected by ten steps. In summary the network assemblages discovered with TCS are highly congruent with the groups identified in the pairwise distance matrix between genes, especially in the LSU and SSU genes. Tajima’s D for the H3 dataset is D = 1.931983, which

### Table 3. Primers used in this study for sequencing of SSU, LSU and H3.

| Gene | Name     | sequence                     | direction |
|------|----------|------------------------------|-----------|
| SSU  | TimA[65] | AMCTGTTGTGATCCCTGCGAG     | forward   |
|      | TimB[65] | TGATGCTACTCGAGGGTCACCT     | reverse   |
|      | S30[66]  | GCTTGTCCTCAAGATAGCGCG     | forward   |
|      | S5K[65]  | TTCTGGCAAAATGCCTGGCG      | reverse   |
|      | 4FB[65]  | CCACGACCGCGGGAGATCCCG     | forward   |
|      | 11806R[66]| CCTTGTGATACCTCTCTCT       | reverse   |
| LSU  | U178[67] | GCACCCCACTGAATYTAAG       | forward   |
|      | L1642[67] | CACGCGCCATCCATTTTCA    | reverse   |
|      | 1200F[67] | CCGAAAGATGTTGAATATGC   | forward   |
|      | R2450[67] | GCTTTGTTTTTAAGAGAGCGGA | reverse   |
|      | U12176[68] | TAAGGGAATCGCGAAAAATTGATCGG | forward |
|      | L3449[67] | ATCTGACTAGAAGCGCTCCA   | reverse   |
|      | U1846[67] | AGGCCGAAGTNNGAGGAAGG | forward   |
|      | L2984[67] | CTGAGCTCTGCCTTGAACCTC  | reverse   |
|      | 28SP155Ster | CTGAGAAGGGGTGTTAGACCGTGATC | forward |
|      | 28SP115Ster | TGGCGTATAGCCGGAGCTGTC | reverse   |
| H3   | H3 AF[69] | ATGCGCTCTGAACAGACAGCG  | forward   |
|      | H3 AR[69] | ATATCTCCAGCCTATRTRGAGAC | reverse   |
|      | H3FNem   | ATGGCTCTGACAAAGACAGACGC  | forward   |
|      | H3RNem   | GTCACCATCTACCCAAGAGA   | reverse   |

TimA and TimB are outer primers spanning the length of the whole fragment. S30 and S5K are internal primer for the first part and 4FB and 1806R for the second part. H3FNem and HRNem are the Colgan et al. [69] primers modified for Nemertodermatidae.

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Figure 2. Majority rule consensus tree (75%) of the LSU ML tree with collapsed terminals. The colours correspond to partitions for BP&P analyses, green indicates the Nemertinoides group, red the mainly European Sterreria subgroup and blue the extra-European Sterreria; the distant S. martindalei n.sp. and S. papuensis n.sp. are shown in orange, as they was not tested with BP&P (s. text). Bootstrap support and Bayesian posterior probabilities are projected from different ML and Bayesian analyses in the order LSU, SSU and H3 where topologies were congruent. Clades supported in at least two of the three gene trees, present as separate networks by statistical parsimony, represented by at least three specimens, and validated by multi locus Bayesian analysis (except S. martindalei n.sp. and S. papuensis n.sp., see text), are formally described and named in this paper. Clades represented by two or less specimens were considered too poorly known for formal description but represent hypothetical species shown here with abbreviations (e.g. N1, S2).

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indicates that this marker underwent neutral evolution. The H3 saturation test indicates saturation (S1).

The gene trees of the three loci estimated with RAxML (best tree with bootstrap values) and MrBayes are not resolved in the deeper topology but consistently support the same putative species (Fig. 2). In the LSU and SSU genes 20 putative species (excluding outgroups) can be identified and in the Histone 3 dataset 27 such groups are supported. The groups identified in the gene trees are identical or highly congruent with the groups identified in the pairwise distance analyses and by the haplotype networks.

In the H3 gene trees thirteen of the 101 ingroup specimens are recovered as one clade splitting from a basal trichotomy. These specimens belong to *S. lundini* n.sp., *S. papuensis* n.sp. and *S. rubra*. This grouping can be interpreted as a saturation artefact. Exclusion of these specimens from the analyses did not change the

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**Figure 3. Parsimony haplotype networks calculated with TCS.** a) LSU gene dataset, b) SSU gene dataset and c) Histone 3 gene dataset. The datasets were reduced and trimmed in order to reduce artefacts from missing data. The colours indicate the subgroups Nemertinoidea (green), mainly European *Sterreria* (red) and extra-European *Sterreria* species (blue), and the not validated *S. martindalei* n.sp. and *S. papuensis* n.sp. in orange (see Fig. 2). Some haplotypes are not connected to any other haplotype given the threshold and are represented by single boxes.

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composition of other tip groups (putative species). The recovered putative species in the gene trees, other than S. lundini n.sp. and S. patapensis n.sp., are consistent with those identified from the pairwise distances and haplotype networks.

Table 4 summarizes the support for identified groups over all methods of discovery.

Species validation

The BP&P analyses with less informative priors or an older root age supported all eleven putative species tested (Fig. 4). The analysis with an informative prior supported all putative species except Sterreria lundini n.sp., S. psammicola, S. rubra and S2 (Fig. 4).

All 35 specimens with any kind of rosy, brownish to bright red colouration recorded were found in the species Sterreria rubra and clade S2 together with four uncoloured specimens and seven specimens for which no colour data was recorded.

Seven out of the eight Sterreria species are not globally distributed, with two species being limited to Hawaii and Papua New Guinea respectively (Fig. 5). Only one species, S. variabilis n.sp., includes specimens from Hawaii, Papua New Guinea, New Caledonia, Bermuda, Portugal and the Mediterranean.

All three Nemertinoides species are distributed along the European coastline from the Mediterranean, via Portugal and the North Sea to the Swedish West Coast (Fig. 5). Within each clade however, no patterning corresponding to geographical distances could be observed. The same is true for the putative species indicated with red (Figs. 2 and 5), which also consist of European specimens and show no geographic pattern.

Clades meeting the above mentioned criteria and were supported in the BP&P analyses (except S. martindalei n.sp. and S. variabilis n.sp.) are formally described, with the exception of clade S2; the specimens of this clade lack photographs or sketches, prohibiting formal description of the species (Tab. 4).

Discussion

Our dataset multiplies the available nucleotide sequence data for Nemertodermatida more than 100 times. The analyses of nucleotide sequence data from the three genes for LSU rRNA, SSU rRNA and Histone 3 identified twelve well supported species among the collected specimens from the two nominal species Nemertinoides elongatus and Sterreria psammicola (Tab. 4). N. wolfgangii n.sp. and S. lundini n.sp. are supported in all genes and analyses but are paraphyletic in the SSU gene tree. The unified species concept defines species as a “separately evolving lineage segment” [19]. SSU has been shown to underestimate species diversity in meiofauna [37]. We therefore conclude that both species are independently evolving lineages warranting formal description as species with incomplete lineage sorting in the SSU gene.

Twelve described species, however, clearly is an underestimation of the true biodiversity in this group of Nemertodermatida (Fig. 2). Even in the material from the Mediterranean, the most densely sampled geographical area, there are several clusters of less than three specimens consistently grouping together in all analyses. These clades represent additional cryptic diversity but we refrain from formally naming such poorly sampled putative species here. This taxonomic undersampling is of course even more drastic outside the Mediterranean. A different form of undersampling emanates from the limited dataset that we have acquired: inclusion of additional molecular markers would have boosted the potential to detect additional cryptic species. Our conservative approach, based on three molecular markers, still raises the number of named species of Nemertinoides and Sterreria from two to twelve.

With the material available to us, morphological distinguishing characters could not be identified a priori for all of the herein described species while studying live specimens, but it is possible that such features will be discovered a posteriori if more material becomes available. We suspect that this is a matter of the level of detail in the morphological investigation, which could be extended from light microscopy to CLSM or electron microscopy in search of additional characters. Even if some of the new species remain diagnosable only based on nucleotide sequences, it is important to recognize such cryptic species in order to appreciate biodiversity, plan management of conservation, and understand ecosystem function [14] (with references). This is especially true if much of species diversity is constituted by cryptic taxa, as is evidently the case in Sterreria and Nemertinoides. It has been argued that species delimitations based solely on nucleotide sequence data would lead to taxonomic instability and confusion as well as taxonomic inflation [38]. However, multilocus coalescent-based methods for species delimitation are firmly grounded in evolutionary theory and population biology, and since these methods are based on explicit probabilities they can be considered more objective than traditional character based taxonomy and allow greater comparability between species [39,40]. Furthermore, if nucleotide-based species diagnoses were implemented, juvenile specimens as well as fragments of specimens, a large proportion of the nemertodermatid specimens encountered, would be available for the study of the diversity of this group.

Adams et al. 2014 [41] defined hyper-cryptic species as nominal species that actually consist of four or more valid species. Our application of molecular species discovery tools have revealed that the two nominal species Nemertinoides elongatus and Sterreria psammicola are hyper-cryptic as they are composed of at least 20 separate species-level clades. Nine of these will be formally described and named below, thereby doubling the number of nominal species of Nemertodermatida. There is no reason to believe that Nemertodermatida are unique in their extensive cryptic diversity: analyses of nucleotide sequence data have unravelled cryptic and hyper-cryptic species within many other groups of marine invertebrates. A case in point is the nominal polychaete species Eumida sanguinea (Orsted, 1843) which was studied by Nygren and Pleijel [42] who identified eight cryptic species among specimens assigned to E. sanguinea and named seven of them using nucleotide-based diagnoses. There are a number of additional cases of cryptic diversity in other polychaete taxa, e.g. the “cosmopolitan” fireworm Eurythoe complanata (Pallas 1766), which was found to consist of three species [43], and Notophyllum foliosum (Sars 1835), which was found to consist of two species [44]. There are relatively few studies at this level of taxonomic resolution in marine meiofauna, but cryptic species have been identified in the flatworm genera Pseudomonocelis [45,46] and Monocelis [47]. A noteworthy example is the “cosmopolitan” flatworm Gyraulus hermaphroditus Ehrenberg, 1831, where studies of karyotype and fine morphology revealed eight separate species in Australia [48], two separate species in the North Sea and the Mediterranean and two separate species at the French Atlantic coast [49]. Leasi et al. [50] used the coalescent-based GMYC algorithm [51] to analyse Cytochrome oxidase subunit I sequences from specimens of the rotifer Testudinella clypeata (Müller, 1786) and found seven cryptic species. Our results further corroborate the hypothesis of the oceans as a hotspot for cryptic diversity put forward by Bickford et al. [14] and exemplified above. Cryptic species occur in all animal groups and they are being identified and described at an accelerating rate.
| Species/clade          | Number of specimens sequenced per gene (LSU/SSU/H3) | Smallest interspecific vs. intraspecific pairwise distance (uncorrected) | Parsimony networks | Gene trees (ML/Bayesian) | BP&P |
|-----------------------|----------------------------------------------------|------------------------------------------------------------------------|-------------------|--------------------------|------|
|                       |                                                   | LSU | SSU | H3 | LSU | SSU | H3 | LSU | SSU | H3 | LSU | SSU | H3 | |
| N1                    | 1/1/–                                             | 2.9 | 10.6 | 0  | –   | –   | –  | yes | –   | –  | 100/1.0 | 37/0.61 | –   | –   | |
| N2                    | 2/2/–                                             | 4.8 | 3.0  | 0  | –   | yes | yes | yes | yes | yes | 100/1.0 | 100/1.0 | 100/1.0 | –   | |
| N3                    | 2/2/2                                             | 7.8 | 3.8  | 0  | 5.8 | 0.3 | yes | yes | yes | yes | 100/1.0 | 100/1.0 | 100/1.0 | –   | |
| N4                    | 2/2/2                                             | 2.9 | 1.2  | 0  | 8.7 | 0.2 | yes | yes | yes | yes | 100/1.0 | 100/1.0 | 100/1.0 | –   | |
| N. elongatus           | 21/20/15                                          | 2.9 | 2.4  | 0  | 5.8 | 0.8 | yes | yes | yes | yes* | 100/1.0 | 100/1.0 | 91/1.0 | 1/1/1 |
| N. wulfangri n.sp.     | 9/7/8                                             | 5.1 | 1.4  | 0  | 4.9 | 1.0 | yes | yes | yes | yes | 100/1.0 | –/–/– | 97/0.99 | 1/1/1 |
| N. glandulosum n.sp.   | 13/13/12                                          | 1.7 | 4.0  | 0  | 4.9 | 0.5 | yes*| yes | yes | yes | 87/0.99 | 100/1.0 | 65/0.81 | 1/1/1 |
| S. abra                | 44/45/17                                          | 1.0 | 1.2  | 0  | 7.9 | 0.9 | yes | yes | yes*| yes*| 94/1.0 | 78/0.98 | 74/0.99/1.0 | 1/0/0.93 |
| S2                    | 3/3/1                                             | 1.5 | 0.8  | 0  | 7.9 | 0.0 | yes | yes | yes | yes | 100/1.0 | 91/1.0 | 70/0.97 | 1/0/0.93 |
| S3                    | 1/1/–                                             | 1.0 | 0.8  | 0  | –   | –   | –  | yes | yes | yes | 100/1.0 | 83/1.0 | –   | –   | |
| S. lundini n.sp.       | 13/12/4                                           | 4.2 | 1.7  | 0  | –   | yes | yes | yes*| yes*| yes*| 98/0.99 | –/–/– | 99/0.99/1.0 | 1/0/0.93 |
| S. psammicola          | 6/6/2                                             | 4.2 | 1.7  | 0  | 13.4/0 | yes*| yes | yes | yes | yes | 98/1.0 | 100/1.0 | 100/1.0 | 1/0/0.93 |
| S. papuensis n.sp.     | 10/10/4                                           | 12.8| 3.4  | 0  | 10.6 | 0.3 | yes*| yes | yes*| yes*| 100/1.0 | 100/1.0 | 99/0.98/1.0 | –   |
| S7                    | 1/1/1                                             | –   | 3.4  | 0  | 10.6 | 0.0 | yes | yes | yes | yes | 100/1.0 | 100/1.0 | –   | –   | |
| S. martindalei n.sp.   | 3/3/3                                             | 13.7| 5.5  | 0  | 13.1 | 0.1 | yes | yes | yes | yes | 100/1.0 | 100/1.0 | 100/1.0 | –   | |
| S. ylva n.sp.          | 4/4/4                                             | 5.4 | 0.2  | 0.1 | 9.4 | 0.1 | yes | 2   | 10  | 99/1.0 | 93/0.58 | 97/0.94 | 1/1/1 |
| S. monolithes n.sp.    | 3/3/3                                             | 5.4 | 0.2  | 0  | 9.6 | 0.0 | yes | 2   | 10  | 100/1.0 | 89/0.97 | 100/1.0 | 1/1/1 |
| S. bounetti n.sp.      | 7/7/6                                             | 6.4 | 0.2  | 0  | 14.2 | 1.4 | yes | yes | 5   | 100/1.0 | 100/1.0 | 46/0.65 | 1/1/1 |
| P3                    | 1/1/2                                             | 14.1| 8.3  | 0  | 9.4 | 0.3 | yes | yes | yes | yes | 100/1.0 | 98/1.0 | 100/1.0 | –   | |
| S. variabilis n.sp.    | 16/16/14                                          | 6.4 | 5.2  | 0  | 14.8 | 9.9 | yes*| yes*| no  | 95/0.99 | 80/0.78 | 7/0.88 | 1/1/1 |
| NCP                   | 2/2/2                                             | –   | –    | –  | 12.5| 2.1 | –   | yes | –   | –   | 100/1.0 | 99/0.91 | –/–/– | 1/1/1 |
| NC/ PNG/H             | 4/5/4                                             | –   | –    | –  | 8.6 | 3.8 | yes*| –/–| –   | –   | 56/0.92 | –    | –   | –   | |
| Eur                   | 5/5/4                                             | –   | –    | –  | 8.6 | 0.0 | –   | yes | –/–| –   | 100/1.0 | –   | –   | –   | |
| Ber                   | 4/4/4                                             | –   | –    | –  | 13.1| 0.0 | –   | yes | –/–| –   | 100/1.0 | –   | –   | –   | |

The number of specimens is given per gene in the order LSU, SSU and H3. The uncorrected percentage of pairwise distances was calculated on the same reduced dataset as was used for the calculation of the parsimony networks; the lowest average percentage of pairwise distance between any two clades within the datasets versus the averaged intraspecific pairwise distance is shown. The parsimony networks were calculated using RAxML and MrBayes, support is given in this order; non-monophyletic clades are indicated with a hash in superscript (#). The program BP&P relies on a true guide tree, which due to the low taxon sampling we could not provide. We therefore decided to test only species with at least three specimens, thus some of the clades in the dataset were not tested and will not be named in this paper (except for S. martindalei and S. papuensis, which were well supported and described without BP&P validation). The posterior probabilities of the three different BP&P analyses are shown in the table. Due to the high variability of S. variabilis clade in all analyses, the clusters within this clade were tested for all analyses with the H3 gene. However, as this part of the tree was not resolved in either phylogenetic analysis, so no BP&B validation could be performed. A dash indicates missing data, or not tested clades.

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meiofauna is far higher than the 11%–43% estimate proposed by [14,52,53]. We hypothesize that the amount of cryptic diversity in derma westbladi they are clearly subtle. The nemertodermatid species differences in habitat specific to the identified species; if present moderate organic content. However, we did not observe any 37 m in sand that reached from coarse to very fine with low to provided in this study.

Identification of species through the metagenetic approach requires a populated database of reference sequences from specimens that were identified by a specialist, such as those [56]. It will also provide a more complete snapshot of diversity, as the morphological identification stage is eliminated. This procedure is following by PCR amplification of a selected marker, pyrosequencing [16,54,55], where DNA is extracted from sediment samples for future identification. Application of metagenetic methods e.g. groups of meiofauna, especially fragile groups such as Acoela, fragmentary knowledge of their diversity. This is true also for other live specimens, which may be part of the explanation for our nemertodermatid crown group, which includes the phylogenetic position, pertains to the clade as a whole. The vicariance hypothesis cannot explain the presence of littoral continental drift in combination with speciation. However, the poor capacity for dispersal that it implies, a first hypothesis may be to invoke vicariance, explaining the patterns by the phylogenetic position of Nemertodermatida [57,58], the group according to both of the two currently competing hypotheses on the phylogenetic position of Nemertodermatida [57,58], the group numbers refer to positions in the respective alignments and in brackets to the position in the sequences in the type specimen (genbank accession number). doi:10.1371/journal.pone.0107688.t005

Table 5. Molecular Diagnostic character of all newly described species in the three genes used in this study.

|                | LSU       | SSU       | H3        |
|----------------|-----------|-----------|-----------|
| N. glandulosum n.sp. (13185) | 2493 (1323) T, 2506 (1336) A | 175 (149) C, 696 (660) A, 725 (688) C, 727 (690) C | 43 (31) C, 70 (58) T, 79 (67) T, 103 (91) T |
| N. wolfgangi n.sp. (13453) | 2450 (1292) T, 2493 (1333) C, 2603 (1443) C | 192 (135) G | 79 G, 103 C, 262 T |
| S. boucheti n.sp. (PNG75) | 2195 (1060) G, 2205 (1070) G, 2333 (1198) G, 2487 (1288) A, 2510 (1331) A | 631 (579) G, 696 (643) A, 723 (669) A, 827 (770) A | – |
| S. lundini n.sp. (08117) | 1705–1708 (1463–1467) CTCTTC (insert) | 701 (632) T, 780 (705) C | – |
| S. martindelei n.sp. (10056) | 2037 (1611) A, 2048 (1622) C, 2193 (1757) G, 2466 (2021) A, 2523 (2061) G | 95 (91) A, 115 (110) G, 127 (122) C, 257 (235) T, 263 (241) T, 530 (506) T, 594 (570) C | 26 (11) A, 28 (13) A |
| S. monolithes n.sp. (PNG84) | 2003 (1642) T, 2326 (1953) G, 2506 (2102) T, 2518 (2113) C, 2604 (2199) A | 204 (182) C, 205 (183) A, 682 (651) A | 100 (88) A, 223 (211) T, 124 (112) G |
| S. papuensis n.sp. (PNG48) | 1825 (718) T, 1848 (727) A, 1860 (735) C, 2099 (963) A, 2337 (1192) T, 2453 (1290) T | 698 (671) T, 711 (683) G, 780 (746) T | 40 (9) G, 82 (51) C, 118 (87) A |
| S. psammicola (13508) | 1696 (1105) T, 1699 (1108) G, 1704 (1113) G, 1714 (1119) A | 701 (101) A, 710 (110) A, 775 (172) C | 61 G, 67 T, 73 A, 136 A, 142 A, 151 G |
| S. variabilis n.sp. (13452) | 2343 (1208) C, 2419 (1261) A, 2995 (1804) A | 631 (502) T, 681 (551) C, 723 (592) G, 827 (693) C | – |
| S. ylvae n.sp. (10054) | 2003 (1642) C, 2431 (2034) T, 2490 (2086) A, 2601 (2195) A | 204 (183) T, 1717 (1658) A | 124 (112) A |

Numbers refer to positions in the respective alignments and in brackets to the position in the sequences in the type specimen (genbank accession number). doi:10.1371/journal.pone.0107688.t005

[14,52,53]. We hypothesize that the amount of cryptic diversity in meiofauna is far higher than the 11%–43% estimate proposed by Appelans et al. [13]. Consequently, the approximation of total marine diversity at 0.7–1.0 million species is likely to be an underestimate.

As noted above, identification and taxonomic study of Nemertodermatida requires specialized methods and access to live specimens, which may be part of the explanation for our fragmentary knowledge of their diversity. This is true also for other groups of meiofauna, especially fragile groups such as Acoela, Platycladus and Gastrotricha that cannot be easily preserved for future identification. Application of metagenetic methods e.g. [16,54,55], where DNA is extracted from sediment samples followed by PCR amplification of a selected marker, pyrosequencing and bioinformatic analysis, has potential to change this as the morphological identification stage is eliminated. This procedure is considered cost-effective in comparison to traditional methods where a number of taxonomists would have to study each sample [56]. It will also provide a more complete snapshot of diversity, as juvenile or damaged specimens would contribute to the results. Identification of species through the metagenetic approach requires a populated database of reference sequences from specimens that were identified by a specialist, such as those provided in this study.

Habitat and Biogeography

All our specimens were found in depths between 1.5 m and 37 m in sand that reached from coarse to very fine with low to moderate organic content. However, we did not observe any differences in habitat specific to the identified species; if present they are clearly subtle. The nemertodermatid species Nemertodermus westbladi and Meunia stichopi, outgroup species in this study, occur in mud down to depths of 600 m. Sampling of nemertodermatids from deeper sediments has only been done in very few locations, mainly in the North-East Atlantic. Thus the existence of deep-water species of Sterreria and Nemertinoides cannot be ruled out.

No fossils identified as nemertodermatids are known, but according to both of the two currently competing hypotheses on the phylogenetic position of Nemertodermatida [57,58], the group is as old as the Cambrian explosion, or even predating it. When trying to explain the current distribution of such an old clade as the Nemertodermatida, and taking into account their biology with the poor capacity for dispersal that it implies, a first hypothesis may be to invoke vicariance, explaining the patterns by continental drift in combination with speciation. However, the vicariance hypothesis cannot explain the presence of littoral Nemertodermatida on younger and isolated islands such as Hawai’i and Bermuda. O’ahu island, where our Hawaiian specimens were collected, is three million years old [59]. It should also be noted that the high estimated age of Nemertodermatida, deduced from their phylogenetic position, pertains to the clade as a whole. The age of the nemertodermatid crown group, which includes the recent species of Sterreria and Nemertinoides, cannot be determined in the absence of any calibration points, but it is likely to be much younger. Clearly, dispersal is the only feasible explanation for the presence of Sterreria on O’ahu and Bermuda. The possibility remains that isolated young islands were colonized by deepwater nemertodermatid species, although currently available evidence seems to favour dispersal from distant shallow habitats, as no deep-water Sterreria specimens are known. The phylogenetic hypothesis derived from Bayesian analyses of the ribosomal datasets indicates that O’ahu was colonized at least twice as Sterreria ylvae n.sp., Sterreria martindelei n.sp. and Sterreria variabilis n.sp. are not each others closest relatives. Current
Sampling of Sterreria on O’ahu was restricted to one site. Extended sampling of nemertodermatids in the Hawaiian archipelago, where the islands are of different ages ranging from 28 Myears to 400,000 years [59], would allow an estimate of rates of colonization and speciation within Sterreria as well as indicating the relative importance of dispersal and speciation in nemertodermatid diversity. Similar studies in Macaronesia, with its volcanic islands of different ages and degrees of geographical isolation, would also shed light on the genetic distinctness of the Bermudian species and the transatlantic dispersal.

Our results show that Nemertodermatida mostly do not conform with the EiE hypothesis. The supposedly wide-ranging Sterreria psammicola and Nemertinoides elongatus both consist of complexes of cryptic species. Some of the species, e.g. Nemertinoides woflfgangi n.sp., show a distribution pattern restricted to one ocean, in this case the Mediterranean, which is, as noted above, the best represented area in this study. However, other species have more extensive distribution areas, such as Nemertinoides elongatus and Sterreria rubra, ranging from the Adriatic, through the Mediterranean via Portugal and the North Sea into the Skagerrak without exhibiting any apparent genetic structure. This indicates considerable dispersal ability in these interstitial worms. Outside Europe, the findings in the Madang lagoon, Papua New Guinea, are particularly striking: the 25 animals collected from four adjacent localities (less than 10 km distance) belong to six different species (S. papuensis n.sp., S. variabilis n.sp., S. boucheti n.sp., S. monolithes n.sp., S7 and P3). Of those, S. papuensis n.sp. and S7 are more closely related to species with European distribution, than to species from geographically closer localities. This is clearly not consistent with an isolation by distance pattern and again indicates dispersal.

Only one species in our dataset appears to be truly cosmopolitan: Sterreria variabilis n.sp. However, since the Histone 3 gene splits this nominal species into geographically structured clades, the existence of yet another unresolved species complex is possible.

**Taxonomic part**

**Family: Nemertodermatidae Steinböck, 1930.** Nemertodermatida without a female pore or bursa. Male pore supraterminal or dorsal. Sequential hermaphrodites. Sperm radially symmetric. Lithocyte in blisters. Usually with epidermal bottle glands.

**Genus: Nemertinoides Riser, 1987.** Diagnosis (emended): Nemertodermatidae with elongated body and constriction at level of statocyst. Mouth in anterior half of body, male pore variable at U40 or subterminal, testes post-oral, ovaries in posterior half of body, reaching posterior of male opening.

Remarks: The far anterior position of the male pore in *N. elongatus* with posterior ovaries were the main arguments for the naming of a new genus by Riser [11]. However, the position of the male pore in relation to body length proved to be not informative on genus level, as opposed to the relative position of the ovaries reaching posterior of the male opening.

**Three species**

- **N. elongatus Riser, 1987.** Material examined: 21 living specimens in squeeze preparation collected during summer between the years 2007 and 2013 in western Sweden (6), the North Sea (1), southern Portugal (3), the French Mediterranean (1), the Tyrrenian Sea (7), and the Adriatic (3). More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Description: Up to 6 mm long, region anterior of statocyst slimmer than rest of body. Statocyst at U4. Large bottle glands in epidermis in anterior half of body. Mouth at U25. Male opening at U40 (Fig. 6c); false seminal vesicle (fsv) directly anterior to this (Fig. 6d). Paired ovaries extend from posterior of fsv to posterior tip. Found in slightly coarser sand from the intertidal to 30 m depth.

Diagnosis (emended): Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 2.9/0.6% (LSU), 2.4/0.1% (SSU) and 5.8/0.8% (H3).

Remarks: This species conforms to the description of *N. elongatus* in Riser [11] with the position of the male opening at about U40 and the distribution of epidermal glands only in the anterior half of the body length. Species supported by all three genes in this study. The type material for *N. elongatus* was collected in the Western Atlantic (holotype at the Massachusetts coast and paratypes at the New Brunswick coast). Subsequent specimens identified as *N. elongatus* were collected in the Adriatic
by Sterrer [9]. There is some uncertainty attached to the identification of this species, as we did not have access to specimens from the type localities; it is possible that our specimens represent a species different from the original *N. elongatus*.

Distribution: Western Atlantic, Swedish West coast, North Sea, southern Portugal, Mediterranean.

**-Nemertinoides wolfgangi** n.sp. Material examined: 9 living specimens in squeeze preparation collected mostly in summer between the years 2008 and 2013 in the French Mediterranean (1), the Tyrrhenian Sea (5), Sicily (2), and the Adriatic (1). More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Typematerial: Holotype SMNH type-8632 (collection code UJ-13453). Mature specimen collected near Cherso, Croatia, by Marco Curini Galletti 22 September 2013. Photographs of the holotype specimen deposited at the Swedish Museum of Natural History, Stockholm.

Description: Reaching more than 5 mm in length. Anterior narrow with body comparatively wide, wobbly. Posterior rounded (Fig. 6f, g). Male copulatory organ (mco) at U90 (Fig. 6f). Paired ovaries in posterior half of body, also posterior of mco (Fig. 6e). Statocyst anterior of U10. Frontal glands open centrally at anterior tip, reaching to about U30. Epidermal bottle glands abundant all over body.

Diagnosis: Morphologically not clearly distinguishable from *N. glandulosum* n.sp. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 2.0/0.2% (LSU), 1.4/0.3% (SSU) and 4.9/1.0% (H3).

Remarks: Species supported by all three genes in this study, but is paraphyletic in the SSU gene tree. Distribution: Mediterranean.

**-Nemertinoides glandulosum** n.sp. Material examined: 13 living specimens in squeeze preparation collected mostly in summer between the years 2007 and 2013 in the North Sea (2), southern Portugal (2), the French Mediterranean (3), and the Tyrrhenian Sea (6). More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Typematerial: Holotype SMNH type-8631 (collection code UJ-13185), mature specimen collected near Faro, Portugal, by Inga Meyer-Wachsmuth 23 May 2013. Photographs of the holotype deposited at the Swedish Museum of Natural History, Stockholm.

Description: Up to 5 mm long. Anterior part narrow compared to wider, wobblier body. Epidermis thick and glandular (Fig. 6b). Posterior more pointy (Fig. 6b). Mouth at U35 (Fig. 6b). Statocyst anterior of U10. Frontal glands open centrally, secretions globular and connected by fine thread like pearls on a chain (Fig. 6c). Mco far in posterior (Fig. 6a).

Diagnosis: Although generally more slender and with slightly more pointy posterior, morphologically not clearly distinguishable from *N. wolfgangi* n.sp. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 2.0/0.4% (LSU), 1.4/0.5% (SSU) and 4.9/0.5% (H3).

Remarks: Species supported by all three genes in this study. Distribution: Mediterranean, southern Portugal, North Sea.

Etymology: Glandula = latin for gland. This species, in contrast to the prior described *N. elongatus*, has epidermal glands also in posterior.

**Genus: Sterreria** Lundin, 2000.

Diagnosis (emended): Elongated Nemertodermatidae without head constriction.

**Genus: Sterreria** Lundin, 2000. Diagnosis (emended): Elongated Nemertodermatidae without head constriction.
Statocyst more posterior than in *Nemertinoides*. Mouth ventral at U50. Paired testes follicular. Male pore opens dorsally at U90, fsv anterior to that. Paired ovaries anterior of mco; oocytes mature caudad.

Remarks: The species *Sterreria psammicola* has been described originally within the genus *Nemertoderma* by Sterrer in Riedl (1970). Due to differences in epidermal structure and differences in the position of the reproductive glands Lundin [60] created a new monotypic genus *Sterreria* and placed *Sterreria psammicola* in this.
The mouth has been observed only in few specimens, it is hypothesized to be a temporary feature as in Nemertoderma westbladi [61–63]. “Male maturity seems to precede female maturity” [9].

Nine species

**-Sterreria psammicola** (Sterrer, 1970). Material examined: 6 living specimens in squeeze preparation collected mostly in summer between the years 2007 and 2013 in southern Portugal (2), the Tyrrhenian Sea (2), and the Adriatic (2). More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Typematerial: Neotype SMNH type-8640 (collection code UJ-15300). Mature specimen collected by Marco Curini Galletti 28 September 2013 at Miramare near Trieste, Italy.

Description: Colourless (Fig. 7c). Frontal glands prominent, opening centre at anterior tip, secretions small, globular or oval (Fig. 7f). Epidermal glands distributed equally over body (Fig. 7c). Borders between epidermal cells not clearly visible (no “scaly” appearance). Posterior rounded, with adhesive area (Fig. 7d). Statocyst further constricted between statoliths than in other groups.

Diagnosis (emended): Morphologically not clearly distinguishable from S. lundini n.sp. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 4.2/0.8% (LSU), 1.7/0.2% (SSU) and 13.4/0.0% (H3).

Remarks: *Sterreria psammicola* has been described from Croatia as a filiform worm living in shallow sandy sediments under the name *Nemertoderma psammicola* by Sterrer in Riedl [64]. The species was described based on some specimens in which “a salmon-red longitudinal stripe is usually [italics by the present authors] present in the first third of the body length” [64]. Specimens from this clade conform to the description without colouration and two specimens were collected near the type locality of *Sterreria psammicola* in Croatia. In the formal description in 1970 no type material was deposited. Species supported by all three genes in this study.

Distribution: Mediterranean, southern Portugal.

**-Sterreria rubra** (Faubel, 1976). Material examined: 45 living specimens in squeeze preparation collected mostly in summer between 2007 and 2013 in western Sweden (1), the North Sea (6), southern Portugal (6), the Tyrrhenian Sea (29), and the Adriatic (3). More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Description: Up to 1 cm long. Opaque, dirty rose. Body surface appears “scaly” due to visible epidermal cell borders (Fig. 7j). Ovaries paired, usually two mature eggs and several small oocytes (Fig. 7h). Posterior tip wide.

Diagnosis (emended): Usually pigmented, rose, bright red or brownish; only in anterior part or all over body (Fig. 1a, b; 7 g-j). Frontal glands prominent reaching far behind statocyst; opening fanning out at the anterior tip; secretions rod shaped (Fig. 7i). Epidermal glands small, distributed densely especially in anterior third of body, but never as prominent as in *Nemertinoides*-species. Body surface appears “scaly” due to visible epidermal cell borders (Fig. 7j). Testes lateral. Male pore at U90; ov just anterior to that (Fig. 7g). Ovaries paired, usually two mature eggs and several small oocytes (Fig. 7h). Posterior tip wide.

Diagnosis: Rosy, never bright red. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 12.6/0.6% (LSU), 3.4/0.1% (SSU) and 10.6/0.3% (H3).

Remarks: This species is paraphyletic in the SSU gene tree and polyphyletic in the H3 gene tree (saturation artefact).

Distribution: Mediterranean.

ETYMOLOGY: After Kennet Lundin, the first researcher formulating a comprehensive phylogenetic hypothesis for Nemertodermatida.

**-Sterreria papuensis** n.sp. Material examined: 10 living specimens in squeeze preparation collected in November 2012 near four different islands (Siar, Tab, Panab, Wanad) in Madang Lagoon, Papua New Guinea. More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Typematerial: Holotype SMNH type-8637 (collection code UJ-08117). Male mature specimen collected near Castiglione della Pescaia, Italy, by Marco Curini Galletti 19 May 2008. Photographs of the holotype deposited at the Swedish Museum of Natural History, Stockholm.

Description: Up to 4 mm long. Colourless (Fig. 7a). Epidermal glands distributed over whole body. Frontal glands prominent with globular or oval secretions (Fig. 7b). Male pore at U90 (Fig. 7c). Posterior end rounded. Borders between epidermal cells not clearly visible.

Diagnosis: Morphologically not clearly distinguishable from *S. psammicola*. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 1.4/0.1% (LSU) and 1.7/0.1% (SSU).

Remarks: This species is paraphyletic in the SSU gene tree and polyphyletic in the H3 gene tree (saturation artefact).

Distribution: Mediterranean.

ETYMOLOGY: After Kennet Lundin, the first researcher formulating a comprehensive phylogenetic hypothesis for Nemertodermatida.
Figure 7. Diversity within the genus Sterreria. a–c: Sterreria lundini n.sp. a) Overview over whole animal. b) Anterior with statocyst with double statoliths and oval frontal gland secretions. c) Posterior with fsv and male pore (mp). d–e: Sterreria psammicola. d) Posterior with fsv male pore (mp) and adhesive area (ad). e) Overview over male mature animal with large epithelial glands distributed more or less evenly over the whole body and mco in the posterior. f) Anterior part of an animal with epithelial glands, statocyst and central frontal gland opening (fp). g–j: Sterreria nubra. g) Posterior with two mature eggs (e) and mco. h) Overview over the same male and female mature specimen with mature eggs (e) and oocytes (oo). i) Anterior with statocyst and rod-shaped frontal gland secretions. j) Detail of the epidermis in the anterior of an animal with cell borders clearly visible, giving the animal a "scaly" appearance. k: Sterreria papuensis n.sp. Anterior part of the holotype. All photographs are taken of live specimens in squeeze preparation. a, b, c, f and k are photographs of the holotype and neotype specimens respectively. The scale bars indicate 100 μm for each photograph.

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Etymology: papuensis = coming from Papua (New Guinea). This is remarkable as this species is closely related to otherwise exclusively European (putative) species.

-Sterreria bouchei n.sp. Material examined: 7 living specimens in squeeze preparation collected in November 2012 near two different islands (Panab, Wanad) in Madang Lagoon, Papua New Guinea. More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Type material: Holotype SMNH type-8633 (collection code PNG75). One immature specimen collected near Wanad Island, Papua New Guinea, by Inga Meyer-Wachsmuth in November 2012. Photographs of the holotype deposited at the Swedish Museum of Natural History, Stockholm.

Description: 1.5 mm long. Colourless. Frontal glands open centrally, secretions rod-shaped, reaching posterior until U35. Large epidermal glands all over body. Statoliths in statocyst well separated. Follicular testes interspersed with ovary. Male pore at U90; mature sperm converge to fsv anterior of that.

Diagnosis: This species is delineated molecularly by all three genes used in this study. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 4.8/0.1% (LSU), 5.2/0% (SSU) and 14.2/1.4% (H3).

Remarks: Species supported by all three genes in this study.

Distribution: Madang lagoon, Papua New Guinea.

Etymology: After Professor Philippe Bouchet, who hosted I. M-W at the 2012 Biodiversity expedition (www.ourplanetreviewed.org) and thus made safe sampling in a place as remote as Papua New Guinea possible.

-Sterreria monolithes n.sp. Material examined: 3 living specimens in squeeze preparation collected in November 2012 near two different islands (Panab, Wanad) in Madang Lagoon, Papua New Guinea. More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Type material: Holotype SMNH type-8636 (collection code PNG). Immature specimen collected near Wanad Island, Papua New Guinea, by Inga Meyer-Wachsmuth in November 2012. Photographs of the holotype deposited at the Swedish Museum of Natural History, Stockholm.

Description: Opaque silvery glossy, no pigmentation. Frontal glands open fanning out; rod-shaped criss-crossing secretions.

Diagnosis: This species cannot be morphologically delineated. It is supported by all three genes used in this study, molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 2.6/0% (LSU), 0.1/0% (SSU) and 9.6/0% (H3).

Remarks: All three specimens of species had only one statolith in the statocyst. Aberrant numbers of statoliths can be observed frequently in other species of Nemertodermatida without taxonomic implications. However, it is unusual to find only specimens with an abnormal number of statoliths. All studied specimens were immature.

Distribution: Madang lagoon, Papua New Guinea.

Etymology: monos = greek for single, lithos = greek for stone, referring to the single lithocyte in the statocyst in all three studied specimens.

-Sterreria ylvae n.sp. Examined material: 4 living specimens in squeeze preparation collected in May 2010 at Waimanolo beach, O’ahu island, Hawaii by Ulf and Ylva Jondelius. More detailed information about individuals and further photographs are accessible at http://acoela.myspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Type: Holotype SMNH type-8639 (collection code UJ-10054). One mature specimen collected at Waimanolo, Hawaii, by Ulf Jondelius 31. May 2010. Photographs of the holotype deposited at the Swedish Museum of Natural History, Stockholm.

Description: Statocyst positioned at U15, statoliths appear encapsulated into separate membrane inside statocyst (Fig. 8b). Anterior long and slender, body slightly wider, wobbly. Posterior narrow and rounded. Epidermal glands large, prominent in anterior half of body. Unicoloured, transparent, opaque only in mid-section of body (Fig. 8c).

Diagnosis: Statoliths appear to be encapsulated into separate membrane inside statocyst. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 2.6/0% (LSU), 0.2/0.1% (SSU), and 9.4/0.1% (H3).

Remarks: Delineated by all three genes used in this study.

Distribution: O’ahu Island, Hawaii, USA.

Etymology: Named after Ylva Jondelius, who collected the samples at Waimanolo beach.

-Sterreria martindalei n.sp. Examined material: 3 living specimens in squeeze preparation collected in May 2010 at Waimanolo beach, O’ahu island, Hawaii by Ulf Jondelius 31. May 2010. Photographs of the holotype deposited at the Swedish Museum of Natural History, Stockholm.

Description: Statocyst at U5. Overall larger, sturdier than Sterreria ylvae n.sp. Posterior two thirds opaque due to large and dense epidermal glands dorsally or ventrally, not laterally. Frontal gland opening central, secretions ellipsoid, reaching posterior till U35; clear border between frontal and epidermal glands. Ovaries between U50 and U75 (Fig. 8a).

Diagnosis: Statocyst more anterior than in Sterreria ylvae n.sp. and overall sturdier, however morphologically not clearly distinguishable from Sterreria ylvae n.sp. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 12.0/0.2% (LSU), 5.5/0% (SSU) and 13.0/0.1% (H3).

Remarks: Species supported by all three genes in this study.

Distribution: O’ahu Island, Hawaii, USA.
Figure 8. Diversity within the genus *Sterreria*. a: *Sterreria martindalei* n.sp. Overview over slightly damaged specimen with mature egg (e) and oocytes in the posterior. b, c: *Sterreria ylvae* n.sp. b) Anterior part with statocyst with double statoliths. c) Mid-section of the body with one mature egg (e) and few oocytes (oo). d, e: *Sterreria monolithes* n.sp. d) Overview over immature animal with only one statolith in the statocyst. e) Detail of the anterior of the same specimen with fanning out frontal gland opening and frontal gland secretions. The single statolith fills nearly the whole statocyst. f: *Sterreria boucheti* n.sp. Overview over complete, immature specimen. g, h: *Sterreria variabilis* n.sp. g) Overview over male mature specimens with club-shaped frontal gland secretions and food particles (f) in gut. h) Detail of male copulatory organ with fsv anterior of male pore (mp) and oocyte (oo). All photographs were taken of the live holotype specimens in squeeze preparation. The scale bars indicate 100 µm for each photograph.
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Etymology: After Prof. Mark Martindale, who hosted U. J. in Hawaii and made sampling possible.

*Sterreria variabilis* n.sp. Material examined: 15 living specimens in squeeze preparation collected mainly in summer between the years 2000 and 2013 in the Adriatic (2), the Tyrrenhenian Sea (2), southern Portugal (1), Bermuda (4), New Caledonia (4), Papua New Guinea (2), and Hawaii (1). More detailed information about individuals and further photographs are accessible at http://acoela.mspecies.info/, the scratchpads database for Acoela and Nemertodermatida.

Type material: Holotype SMNH type-0638 (collection code UJ-13452). One male mature specimen collected at Cherso, Croatia, collected by Marco Curini Galletti 22 September 2013. Photographs of the holotype deposited at the Swedish Museum of Natural History, Stockholm.

Description: Morphologically diverse. Bermuda and Portuguese specimens less than 1 mm long, slender, specimens from most other localities comparatively large and bulky; Portuguese specimen was immature. Opaque due to abundance of epidermal glands and oil droplets. Anterior rounded and comparatively slender; mature specimens broaden considerably at level of the statocyst; increase in width more gradual in immature specimens. Posterior tip rounded, possibly with adhesive structure. Frontal glands prominent, opening central, secretion in long bundles; in Bermudian and Portuguese specimens distal base at U35 bulbous, in Adriatic specimens club-shaped (Fig. 8h). Paired testes follicular and lateral; vasa deferentia converge to left anterior of mco (Fig. 8g). Male opening in the posterior surrounded by rosette of small glands. Paired ovaries lateral in posterior half or three quarters of body length, oocytes small.

Diagnosis: A common morphological delineation is not possible in this species. Molecular character diagnosis in table 5. Smallest interspecific pairwise distance vs. intraspecific distance 4.8/0.6% (LSU), 5.2/0.3% (SSU) and 14.8/9.9% (H3).

Remarks: This species is supported by the SSU and LSU genes but not by the higher resolving Histone 3 gene. This species is the only truly cosmopolitan species in this dataset. The Histone 3 gene splits the species in geographic clades. This makes the existence of another, not yet resolved species complex, possible. Population genetic studies of this species with increased geographic sampling and sample size might give valuable insights into the dispersal capabilities of interstitial meiofauna.

Distribution: Mediterranean, southern Portugal, Bermuda, New Caledonia, Papua New Guinea, Hawaii.

Etymology: *variabilis* = latin for variable, as this species is morphologically and genetically changeable throughout its known range.

Supporting Information

**Figure S1** Saturation plot for the H3 gene across the whole dataset. Plotted are the uncorrected p-distances versus the phylogenetic distances between pairs of sequences. The level distribution of the points indicates saturation.

(TIFF)

**Figure S2** Best ML tree calculated with RAxML of the LSU rRNA dataset with bootstrap support plotted on the branches. Putative species with binomial names are formally described in the present study, those with abbreviations represent candidate species. The branch colours correspond to partitions for BP&P analyses, green indicates the Nemertinoides group, red the mainly European Sterreria subgroup and blue the extra-European Sterreria species; orange species have not been validated with BP&P.

(TIFF)

**Figure S3** Majority rule consensus tree estimated with MrBayes of the LSU rRNA dataset with Bayesian posterior probabilities plotted on the nodes. Putative species with binomial names are formally described in the present study, those with abbreviations represent candidate species. The branch colours correspond to partitions for BP&P analyses, green indicates the Nemertinoides group, red the mainly European Sterreria subgroup and blue the extra-European Sterreria species; orange species have not been validated with BP&P.

(TIFF)

**Figure S4** Best ML tree calculated with RAxML of the SSU rRNA dataset with bootstrap support plotted on the nodes. Putative species with binomial names are formally described in the present study, those with abbreviations represent candidate species. The branch colours correspond to partitions for BP&P analyses, green indicates the Nemertinoides group, red the mainly European Sterreria subgroup and blue the extra-European Sterreria species; orange species have not been validated with BP&P.

(TIFF)

**Figure S5** Majority rule consensus tree estimated with MrBayes of the SSU rRNA dataset with Bayesian posterior probabilities plotted on the nodes. Putative species with binomial names are formally described in the present study, those with abbreviations represent candidate species. The branch colours correspond to partitions for BP&P analyses, green indicates the Nemertinoides group, red the mainly European Sterreria subgroup and blue the extra-European Sterreria species; orange species have not been validated with BP&P.

(TIFF)

**Figure S6** Best ML tree calculated with RAxML of the Histone 3 dataset with bootstrap support plotted on the nodes. Putative species with binomial names are formally described in the present study, those with abbreviations represent candidate species. The branch colours correspond to partitions for BP&P analyses, green indicates the Nemertinoides group, red the mainly European Sterreria subgroup and blue the extra-European Sterreria species; orange species have not been validated with BP&P.

(TIFF)

**Figure S7** Majority rule consensus tree estimated with MrBayes of the Histone 3 dataset with Bayesian posterior probabilities plotted on the nodes. Putative species with binomial names are formally described in the present study, those with abbreviations represent candidate species. The branch colours correspond to partitions for BP&P analyses, green indicates the Nemertinoides group, red the mainly European Sterreria subgroup and blue the extra-European Sterreria species; orange species have not been validated with BP&P.

(TIFF)

**Table S1** Reduced datasets. Table showing, which sequences per dataset had been excluded for pairwise distance calculations and TCS analyses.

(XLSX)

**Table S2** Averaged pairwise distances across the LSU rRNA dataset.

(XLSX)

**Table S3** Averaged pairwise distances across the SSU rRNA dataset.

(XLSX)
Table S4  Averaged pairwise distances across the Histone 3 gene dataset. For Sterreria variabilis n.s.p. pairwise distances are also given for the four subdivisions within the species. (XLSX)

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