

MOLECULAR PHYLOGENY OF NORTH SEA SEPIOLINAE (CEPHALOPODA: SEPIOLIDAE) REVEALS AN OVERLOOKED *SEPIOLA* SPECIES

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ABSTRACT

Recent bottom trawl surveys in the North Sea have broadened our understanding of an obscure group of bobtail squids (Sepioidae: Sepiolinae). There are at least two *Sepietta* species in this region, viz. *Sepietta oweniana* and *Sepietta neglecta*, and three *Sepiolo* species, one of which is undescribed, viz. *Sepiolo atlantica*, *Sepiolo pfefferi* and *Sepiolo* sp. nov. *Sepiolo pfefferi* is distinct from *Sepiolo aurantiaca* because of differences in the hectocotylus; the type localities of both species are far apart and data in GenBank indicate a substantial genetic difference between them. It is unclear how to distinguish *Sepiolo* sp. nov. morphologically from its sister species *S. atlantica*, but molecular phylogenetic analyses and distributional data readily set them apart. The occurrence of *Sepiolo rondeleti* in the northeastern Atlantic could not be confirmed. Several obviously incorrect or dubious identifications regarding GenBank data are listed.

INTRODUCTION

The Sepiolinae are small, poorly known, so-called bobtail squids, with a mantle length of up to 5 cm for the largest species used in this study. The subfamily encompasses 5 of the 15 known genera of Sepioidae, viz. *Euprymna*, *Inioteuthis*, *Rondeletiolo*, *Sepietta* and *Sepiolo*. Of these, only *Sepiolo* and *Sepietta* are known from the North Sea, where they are represented by seven species (Reid & Jereb, 2005). These are *Sepiolo atlantica* d'Orbigny, 1839–1842 (it is unknown in what year the plate with legends which validate this name were published; see Tillier & Boucher-Rodoni, 1993), *Sepiolo intermedia* Naef, 1912, *Sepiolo rondeleti* Leach, 1817, *Sepiolo aurantiaca* Jatta, 1896, *Sepiolo pfefferi* Grimpe, 1921, *Sepietta neglecta* Naef, 1916 and *Sepietta oweniana* (d'Orbigny, 1839–1841) (the plate with legends, which could validate this name, was published in 1839–1842, whereas the text was published in 1841).

Although Reid & Jereb (2005) include the northeastern Atlantic in the geographical distribution of *S. intermedia*, there is a query about its distribution west of the Strait of Gibraltar (Reid & Jereb, 2005: 163, fig. 236). The occurrence of *S. rondeleti* in the North Sea is doubtful as well, since the recent records (Reid & Jereb, 2005: 168, fig. 244) all refer directly or indirectly to relatively old data in the literature, i.e. Joubin (1895, 1902) and Naef (1912). Apart from that, Reid & Jereb (2005) did not cite a critical paper by Grimpe (1925), in which he attributed the records by Joubin and Naef to *S. oweniana*. Later, Grieg (1933) came to the same conclusion for records of so-called *S. rondeleti* from the Scandinavian part of the North Sea. *Sepiolo rondeleti* is also missing from comprehensive overviews of the German, Dutch and British areas of the North Sea (Jaekel, 1958; Janssen, 1975; Hayward & Ryland, 1990).

Sepiolo aurantiaca was described from the Tyrrhenian Sea by Jatta (1896). At the beginning of the twentieth century, Russell collected twenty specimens of sepiolids from the North

Sea, off the northeast coast of Scotland (*c.* 56°N, 3°W). Despite some details of the hectocotylus, i.e. the lack of the peculiar lobe at the base of the first arm right (or right dorsal arm) (Fig. 1; illustration in Russell, 1922), Russell (1909) attributed these bobtail squids to *S. aurantiaca*. Grimpe (1921) introduced *S. pfefferi* as a new species for sepiolids from the east coast of England (*c.* 53°N, 1°E). *Sepiolo pfefferi* could allegedly be discerned from *S. aurantiaca* by a deep V-shaped (instead of U-shaped) mantle outline (Fig. 2) and a basal lobe only on the left dorsal arm of the hectocotylus. According to Naef (1923) however, the differences between *S. pfefferi* and *S. aurantiaca* are so minimal that he considered *S. pfefferi* to be a form of *S. aurantiaca*.

Clearly, the Sepiolinae represent a controversial group in which species are often difficult to identify morphologically (especially juvenile and female specimens). Sibling species are perhaps most easily distinguished genetically. In such cases a database like GenBank will be most valuable. However, the utility of such databases depends on how well the taxonomic group under study is represented and whether reliably identified voucher specimens are available for more detailed morphological studies. While trying to characterize the North Sea Sepiolinae by DNA analyses ('barcoding') and to unravel their phylogenetic relationships, our initial views, largely based on data in the literature, had to be modified.

MATERIAL AND METHODS

Sampling

Sepiolinae in general are small, obscure, bottom-dwelling organisms, that are most commonly known from the by-catch of fisheries. Except for SCUBA diving, which is not a feasible means to survey larger areas, there does not seem to be an easy way specifically to catch these small cuttlefish. We therefore made use of trawl surveys in this study. For a number of years, IBTS (International Bottom Trawl Survey) and BTS (Beam Trawl Survey) surveys have been conducted in the North Sea

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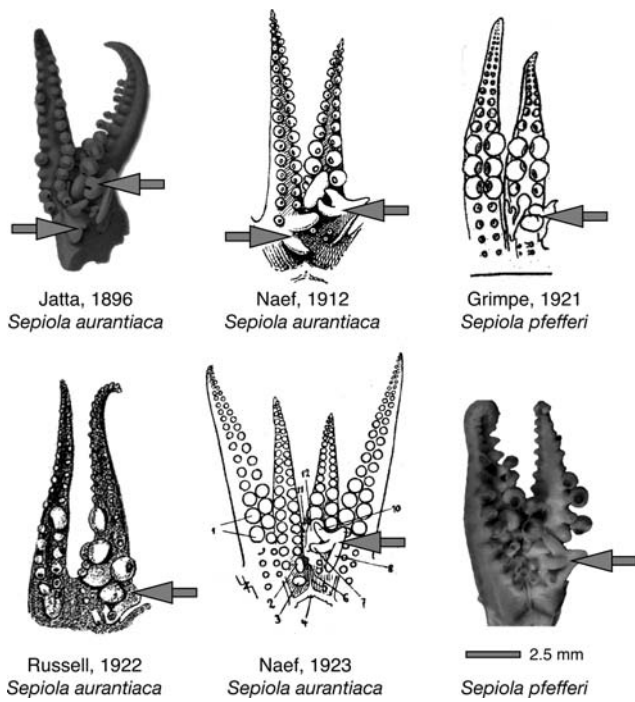


Figure 1. Ventral view of dorsal arms of *Sepiola aurantiaca* and *Sepiola pfefferi*. Arrows indicate basal lobes on the hectocotylus. The scale refers only to the photo in the lower right corner of a hectocotylus of *S. pfefferi*, collected in 1931 from Cove Bay, off Aberdeen (57°06'N 2°03'W).

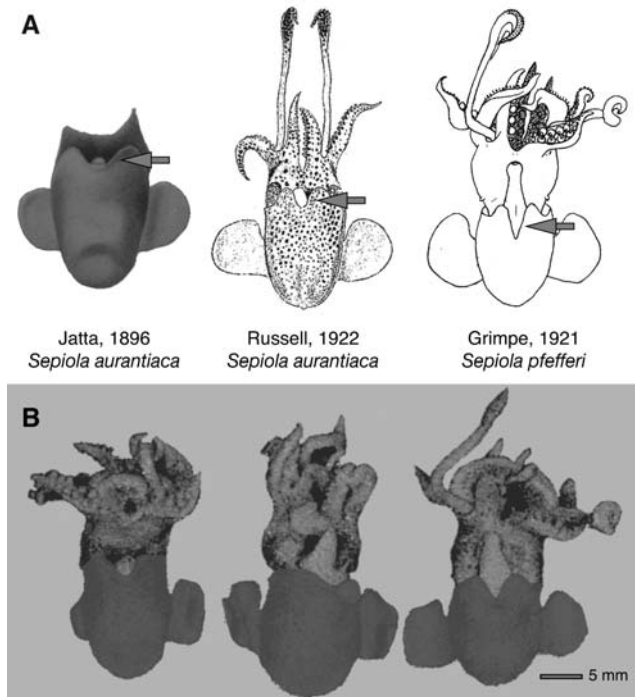


Figure 2. **A.** U- and V-shaped mantle outline (arrows) of *Sepiola aurantiaca* and *Sepiola pfefferi*. **B.** Illustration after Grimpe (1921) showing the type specimens of *S. pfefferi* with their mantles highlighted. The scale only refers to the photographed part of this figure.

by IMARES (the Netherlands Institute for Fisheries Research) under auspices of ICES (the International Council for the Exploration of the Sea). The sepiolids from these surveys were

stored in 96% ethanol and added to the collection of the National Museum of Natural History *Naturalis* (Leiden, The Netherlands). Our identifications were primarily based on morphological data published by Grimpe (1925) and Reid & Jereb (2005), i.e. size, characters of the hectocotylus, the number and arrangement of suckers, and the presence and position of light organs. All species that were indisputably known to occur in the North Sea were represented in our material: *Sepiola atlantica* (21 specimens), *Sepiola aurantiaca*/*Sepiola pfefferi* (5), *Sepietta oweniana* (4) and *Sepietta neglecta* (8). Hence we assume that the IMARES surveys yielded a representative sample of this subfamily for this area. Specimens and associated data (depth, gender) are listed in Table 1 and sampling sites are illustrated in Figure 3.

Type specimens and vouchers

The syntypes of neither *S. aurantiaca* nor *S. pfefferi* could be located in the institutes where they reportedly had been deposited, i.e. the Stazione Zoologica di Napoli, Naples (A. Travaglini, personal communication) and the Museum für Naturkunde der Humboldt-Universität, Berlin (M. Glaubrecht, personal communication), respectively. We are not aware of any recently collected material of undisputable *S. aurantiaca* and a voucher specimen for the sequence in GenBank (AF035708) was not available (M.E. Nishiguchi, personal communication).

DNA isolation, PCR and sequencing

Total genomic DNA was extracted from up to 15 mg of muscle tissue, using a DNeasy® Tissue Kit (Qiagen), following the manufacturer's protocol. Specimen vouchers were deposited at the National Museum of Natural History *Naturalis*, Leiden, The Netherlands. A 658-bp fragment of cytochrome oxidase subunit I (COI), that was shown to be informative in other cephalopod studies (Nishiguchi, Ruby & McFall-Ngai, 1998; Lindgren, Giribet & Nishiguchi, 2004; Nishiguchi, Lopez & Boletzky, 2004), was amplified with primers L1490 5'-GGTCA ACAATCATAAAGATATTGG-3' and H2198 5'-TAAACTT CAGGGTGACCAAAAATCA-3' (Folmer *et al.*, 1994). PCR conditions were 1.5 mM MgCl₂, 0.2 mM dNTPs, 0.4 μM of each primer and 5 U of *Taq* DNA polymerase (Qiagen) per reaction. Reactions were always carried out in a volume of 25 μl using a thermocycle profile of 3 min at 94°C, followed by 40 cycles of 15 s at 94°C, 30 s at 50°C and 40 s at 72°C, and a final extension of 5 min at 72°C. PCR products were purified with Nucleospin® Extract II columns (Macherey-Nagel) and both strands were directly sequenced using the same primers. Sequencing was done on an ABI 3730 automated sequencer (Applied Biosystems) at Macrogen Corp. (Korea). Forward and reverse sequences were assembled and checked using Sequencher version 4.2 (Gene Codes Corp.). Sequences were aligned manually using MacClade 4.08 (Maddison & Maddison, 2003).

Phenetic analysis

A neighbour-joining phylogram (optimality criterion = distance, total character difference, distance measure = uncorrected *p*) was constructed with PAUP 4.0b10 (Swofford, 2003), including sequences from this study, all Sepiolineae sequences that had been previously deposited in GenBank and four out-group sequences (representing Rossinae, Heteroteuthinae and Sepiadariidae), also from GenBank. Genetic distances (uncorrected *p*-distances; absolute number of differences, Table 2) were calculated with PAUP 4.0b10 as well. Although corrected distances [such as General Time Reversible (GTR) model, see

Table 1. Specimens, sampling information and GenBank accession numbers (GB ACCN).

| Species | Lat (°N) | Long (°E) | Depth (m) | Sex | Date | GB ACCN | RMNH_MOL |
|--|-----------|-----------|-----------|-----|--------------|----------|----------|
| <i>Sepietta neglecta</i> Naef, 1916 | 56.33653 | 4.5567 | 62 | F | 2.ii.2005 | FJ231301 | 105647 |
| <i>Sepietta neglecta</i> Naef, 1916 | 56.66683 | -1.61233 | 58 | F | 8.ii.2008. | FJ231312 | 110308 |
| <i>Sepietta neglecta</i> Naef, 1916 | 56.66683 | -1.61233 | 58 | F | 8.ii.2008. | FJ231324 | 110308.1 |
| <i>Sepietta neglecta</i> Naef, 1916 | 56.66683 | -1.61233 | 58 | F | 8.ii.2008. | FJ231325 | 110308.2 |
| <i>Sepietta neglecta</i> Naef, 1916 | 56.66683 | -1.61233 | 58 | F | 8.ii.2008. | FJ231326 | 110308.3 |
| <i>Sepietta neglecta</i> Naef, 1916 | 56.66683 | -1.61233 | 58 | M | 8.ii.2008. | FJ231327 | 110309.1 |
| <i>Sepietta neglecta</i> Naef, 1916 | 56.66683 | -1.61233 | 58 | M | 8.ii.2008. | FJ231328 | 110309.2 |
| <i>Sepietta neglecta</i> Naef, 1916 | 56.66683 | -1.61233 | 58 | M | 8.ii.2008. | FJ231329 | 110309.3 |
| <i>Sepietta oweniana</i> (d'Orbigny, 1839–1841) | 55.73613 | -0.27547 | 73 | M | 9.ix.2005 | FJ231297 | 107426 |
| <i>Sepietta oweniana</i> (d'Orbigny, 1839–1841) | 57.82507 | -2.42627 | 89 | M | 26.viii.2005 | FJ231298 | 108685 |
| <i>Sepietta oweniana</i> (d'Orbigny, 1839–1841) | 55.68268 | 6.40243 | 43 | - | 23.viii.2007 | FJ231299 | 110057 |
| <i>Sepietta oweniana</i> (d'Orbigny, 1839–1841) | 54.64303 | 0.52227 | 70 | F | 12.ix.2005 | FJ231300 | 108941 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 53.4611 | 0.9181 | 20 | F | 15.ii.2006 | FJ231302 | 102844 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 53.106798 | 1.7576 | 38 | M | 26.i.2005 | FJ231303 | 105641 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 53.106798 | 1.7576 | 20 | M | 26.i.2005 | FJ231304 | 105658 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 51.57346 | 2.77096 | 32 | M | 25.i.2005 | FJ231308 | 105638 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 53.62875 | 2.46205 | 25 | M | 11.ix.2007 | FJ231309 | 110059 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 52.2099 | 2.99283 | 38 | F | 23.8.2004 | FJ231310 | 99500 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 54.67276 | 2.57207 | 20 | F | 9.ii.2005 | FJ231311 | 105655 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 54.70469 | 0.37152 | 68 | M | z2006 | FJ231314 | 105637 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 52.68033 | 4.04217 | 24 | F | 28.i.2008 | FJ231316 | 110291 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 54.28333 | 8.03333 | 19 | M | 29.i.2008 | FJ231317 | 110293 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 54.12033 | 4.90917 | 44 | M | 30.i.2008 | FJ231318 | 110297 |
| <i>Sepioloidea atlantica</i> d'Orbigny, 1839–1842 | 53.8275 | 4.21467 | 29 | F | 13.ii.2008 | FJ231322 | 110316 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 56.78483 | -0.23631 | 79 | F | 7.ii.2005 | FJ231305 | 105650 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 57.82509 | -2.42626 | 63 | M | 26.viii.2005 | FJ231306 | 108684 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 55.68268 | 6.40243 | 43 | M | 21.viii.2007 | FJ231307 | 110058 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 54.77194 | -0.25279 | 66 | M | z2006 | FJ231313 | 105636 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 55.21893 | -0.50832 | 75 | F | z2005 | FJ231315 | 108938 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 55.83367 | 3.35133 | 64 | M | 5.ii.2008 | FJ231319 | 110298 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 56.27517 | 4.45883 | 65 | F | 6.ii.2008 | FJ231320 | 110299 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 56.09333 | 1.19967 | 83 | M | 11.ii.2008 | FJ231321 | 110311 |
| <i>Sepioloidea</i> sp. nov. (Goud & de Heij, in prep.) | 54.30417 | -1.27367 | 94 | M | 7.ii.2008 | FJ231323 | 110301 |
| <i>Sepioloidea pfefferi</i> Grimpe, 1921 | 57.74243 | -1.3547 | 90 | M | 26.viii.2005 | FJ231292 | 108846.1 |
| <i>Sepioloidea pfefferi</i> Grimpe, 1921 | 57.74243 | -1.3547 | 90 | - | 26.viii.2005 | FJ231293 | 108846.2 |
| <i>Sepioloidea pfefferi</i> Grimpe, 1921 | 53.44067 | 0.96167 | 20 | F | 26.i.2005 | FJ231294 | 105642 |
| <i>Sepioloidea pfefferi</i> Grimpe, 1921 | 53.46110 | 0.9181 | 38 | F | 15.ii.2006 | FJ231295 | 110056 |
| <i>Sepioloidea pfefferi</i> Grimpe, 1921 | 53.11671 | 1.7827 | 38 | M | 26.i.2005 | FJ231296 | 108845 |

RMNH_MOL refers to the voucher reference number (Molluscan collection) of the National Museum of Natural History, Naturalis (Leiden, The Netherlands). Sex determination: M, male; F, female; -, juvenile (or undetermined).

the paragraph below] will be more realistic (e.g. they compensate for multiple substitutions at the same position), we choose to use uncorrected distances, because these at least show the minimum amount of change.

Phylogenetic analysis

To reduce computational time and focus on the genera present in the area of study, we retained one representative of the sequences from GenBank for *Euprymna tasmanica* (AY293713), *Euprymna scolopes* (AF035701) and *Euprymna hyllebergi* (AY293714) for Bayesian analysis. The sequence in GenBank for *Rondeletiola minor* (AF035714) was also excluded, as we suspect that it represents a specimen of *E. scolopes* (see Discussion). For the Bayesian analysis, the best-fit model of sequence evolution was selected with MRMODELTEST version 2.2 (Nylander, 2004). Both hLRT and AIC selected a GTR model ($N_{st} = 6$) with gamma distribution rate variation across

sites and a proportion of invariable sites (rates = invgamma; GTR + I + G). A flat Dirichlet distribution was selected as prior on state frequencies (statefreqpr = dirichlet (1,1,1,1)). Bayesian inference was done with MRBAYES 3.1.2. (Ronquist & Huelsenbeck, 2003), using two independent runs of four Markov chains with 10,000,000 generations. Sampling was done every 100 generations and the first 5,000,000 generations were discarded (burnin = 50,000; standard deviation of split frequencies was 0.0176 and dropped below 0.01 after 8,775,000 generations). *Sepioloidea lineolata* (AF000064) was specified as outgroup.

RESULTS

Both phenetic and cladistic analyses (Figs 4, 5, respectively) distinguished *Sepietta neglecta* (group F) and *Sepietta oweniana* (group G). No intraspecific variation was observed in either species according to our own data, whereas the interspecific divergence between these species was 7.6% (Table 2).

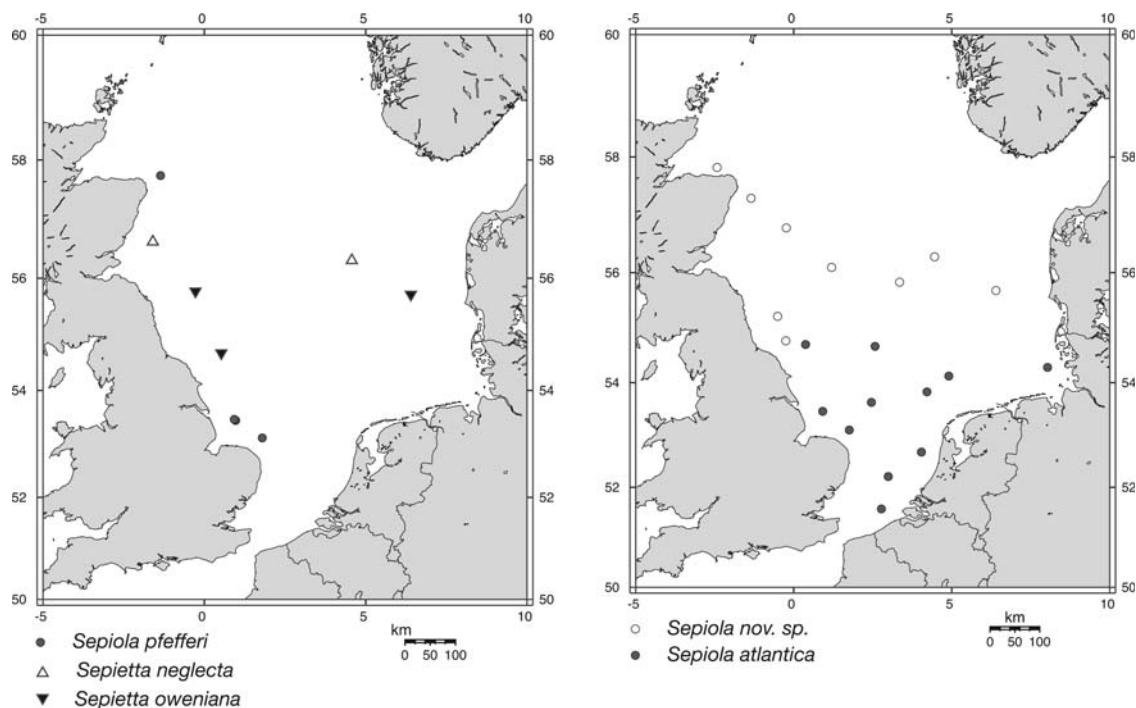


Figure 3. Sampling localities of *Sepioloidea* and *Sepietta* in the North Sea.

Sequences of specimens provisionally named *Sepiola pfefferi* (group E) differed substantially (14.5% sequence divergence) from the only available alleged *Sepiola aurantiaca* sequence (AF035708, GenBank), while intraspecific variation within *S. pfefferi* was low (0.15% or less, Table 2). Specimens initially identified as *Sepiola atlantica*, turned out to represent two monophyletic groups: *S. atlantica* and *Sepiola* sp. nov. (groups A and B respectively; Figs 4, 5) with a minimum sequence divergence of 3.50% (Table 2). There was no intraspecific sequence divergence within *Sepiola* sp. nov. and our data show low intraspecific sequence divergence (0.15%) within *S. atlantica* (Table 2). Numerous Sepioloidea species were represented by more than one COI sequence in GenBank, but rarely did allegedly conspecific individuals form monophyletic groups (Fig. 4, Table 3).

DISCUSSION

Five species of Sepioloidea occur in the North Sea, representing two genera, viz. *Sepietta oweniana* (group G) and *Sepietta neglecta* (group F), and *Sepiola atlantica* (group B), *Sepiola pfefferi* (group E) and *Sepiola* sp. nov. (group A), with different distributional patterns (Fig. 3). For the moment, *S. atlantica* and *Sepiola* sp. nov. are considered sibling species. Based on DNA analyses of COI, a provisional phylogeny reconstruction (Fig. 5) is given, including sepiolid taxa from elsewhere using data in GenBank. Our data lead us to conclude that *Sepiola* is paraphyletic, whereas *Sepietta* is monophyletic. This study indicates that a high percentage of the species, genera and, in one case, subfamilies of Sepioloidea in GenBank are most probably misidentified. This hampered our conclusions, in particular since voucher specimens were not available for inspection.

There is a striking contrast between sequences obtained with this study, that clearly show distinct species that are well separated from each other (small intraspecific sequence divergences and large interspecific divergences) and sequences deposited in GenBank, which showed two other extremes: either sequences of the same so-called species that showed large interspecific

divergences (Table 3) or sequences with limited intraspecific divergences of which the vouchers were identified as different species (group D, Figs 4, 5).

Both this incongruence between our data and the data from GenBank, as well as the high number of likely misidentifications (Fig. 4, Table 3), limited our ability to evaluate whether the sequence of *Sepiola aurantiaca* is actually distinct from group D (Fig. 4, 5). According to Nishiguchi *et al.* (1998, 2004) group D represents seven species. Among these all, except *S. atlantica* (AF035707), were collected in the Mediterranean Sea; most came from the same locality (Banyuls-sur-Mer; Nishiguchi *et al.*, 1998, 2004). Comparing the topology of Figures 4, 5, and divergence among sequences (Table 2) referred to as *S. atlantica* d'Orbigny, 1839–1842, *Sepiola intermedia* Naef, 1912, *Sepiola affinis* Naef, 1912 and *Sepiola ligulata* Naef, 1912, led us to conclude that GenBank entries AF035707, AY293718, AY293716 and AF035710, respectively (group D), were probably misidentified and likely all pertain to the species *Sepiola robusta*.

Considering that *Heteroteuthis dispar* (Rüppell, 1844) (AF035713, group D) belongs to the subfamily Heteroteuthinae instead of Sepioloidea, and does not group with either its congener *Heteroteuthis hawaiiensis* (Berry, 1909) (AY293728) or with other representatives of the Heteroteuthinae, like *Stoloteuthis leucoptera* (Verrill, 1878) (AF000044), this specimen may also have been misidentified. One *S. robusta* Naef, 1912 sequence (AF035711, group D) is problematic in several ways: it has an insertion that causes a frame shift, coding region information is not in GenBank and it has four amino acid substitutions that are otherwise unique among sepiolid COI sequences in GenBank.

The sequence assigned to *S. aurantiaca* in GenBank (AF035708) differs considerably from other members of group D (7.85% sequence divergence between AF035708 and AY293718; Table 2). However, AF035708 is of unusual length and bears four insertions and two deletions, which are unique among sepiolid COI sequences (GenBank). AF035708 also contains stopcodons regardless of the chosen reading frame and even using an optimized alignment to minimize the number of

Table 2. Sequence divergence percentages (calculated as the uncorrected p -distance times one hundred) between sequences used in this study.

| Group | Taxon | Group | Taxon | Seq. divergence (%) |
|-------|---------------------------------------|-------|---------------------------------------|---------------------|
| A | <i>Sepiolo sp. nov.</i> FJ231306 | A | <i>Sepiolo sp. nov.</i> FJ231305 | 0.00 |
| B | <i>Sepiolo atlantica</i> FJ231317 | B | <i>Sepiolo atlantica</i> FJ231302 | 0.15 |
| B | <i>Sepiolo atlantica</i> AY293721 | B | <i>Sepiolo atlantica</i> FJ231317 | 0.77 |
| A | <i>Sepiolo sp. nov.</i> FJ231305 | B | <i>Sepiolo atlantica</i> FJ231302 | 3.50 |
| B | <i>Sepiolo atlantica</i> FJ231317 | A | <i>Sepiolo sp. nov.</i> FJ231305 | 3.65 |
| C | <i>Sepiolo rondeleti</i> AY293720 | C | <i>Sepiolo affinis</i> AF035706 | 0.16 |
| C | <i>Sepiolo rondeleti</i> AY293720 | C | <i>Sepiolo intermedia</i> AF035709 | 0.99 |
| C | <i>Sepiolo intermedia</i> AF035709 | C | <i>Sepiolo affinis</i> AF035706 | 0.83 |
| C | <i>Sepiolo rondeleti</i> AY293720 | B | <i>Sepiolo atlantica</i> FJ231317 | 1.67 |
| C | <i>Sepiolo rondeleti</i> AY293720 | A | <i>Sepiolo sp. nov.</i> FJ231305 | 3.96 |
| C | <i>Sepiolo affinis</i> AF035706 | B | <i>Sepiolo atlantica</i> FJ231317 | 1.81 |
| C | <i>Sepiolo affinis</i> AF035706 | A | <i>Sepiolo sp. nov.</i> FJ231305 | 3.96 |
| C | <i>Sepiolo intermedia</i> AF035709 | B | <i>Sepiolo atlantica</i> FJ231317 | 2.65 |
| C | <i>Sepiolo intermedia</i> AF035709 | A | <i>Sepiolo sp. nov.</i> FJ231305 | 4.78 |
| – | <i>Sepiolo rondeleti</i> AF035712 | C | <i>Sepiolo rondeleti</i> AY293720 | 4.00 |
| – | <i>Sepiolo rondeleti</i> AF035712 | C | <i>Sepiolo affinis</i> AF035706 | 3.84 |
| – | <i>Sepiolo rondeleti</i> AF035712 | C | <i>Sepiolo intermedia</i> AF035709 | 3.67 |
| – | <i>Sepiolo rondeleti</i> AF035712 | B | <i>Sepiolo atlantica</i> FJ231318 | 5.50 |
| – | <i>Sepiolo rondeleti</i> AF035712 | A | <i>Sepiolo sp. nov.</i> FJ231305 | 7.50 |
| D | <i>Sepiolo robusta</i> AY293719 | B | <i>Sepiolo atlantica</i> AF035707 | 0.00 |
| D | <i>Sepiolo robusta</i> AY293719 | D | <i>Sepiolo intermedia</i> AY293718 | 0.46 |
| D | <i>Sepiolo robusta</i> AY293719 | D | <i>Heteroteuthis dispar</i> AF035713 | 0.00 |
| D | <i>Sepiolo robusta</i> AF035711 | D | <i>Sepiolo ligulata</i> AF035710 | 2.66 |
| D | <i>Sepiolo ligulata</i> AF035710 | D | <i>Sepiolo intermedia</i> AY293718 | 1.06 |
| D | <i>Sepiolo ligulata</i> AF035710 | D | <i>Sepiolo affinis</i> AY293716 | 1.77 |
| D | <i>Sepiolo robusta</i> AY293719 | D | <i>Sepiolo aurantiaca</i> AF035708 | 7.85 |
| E | <i>Sepiolo pfefferi</i> FJ231296 | E | <i>Sepiolo pfefferi</i> FJ231295 | 0.00 |
| E | <i>Sepiolo pfefferi</i> FJ231296 | E | <i>Sepiolo pfefferi</i> FJ231294 | 0.15 |
| F | <i>Sepietta neglecta</i> FJ231301 | E | <i>Sepiolo pfefferi</i> FJ231294 | 10.03 |
| F | <i>Sepietta neglecta</i> FJ231329 | F | <i>Sepietta neglecta</i> FJ231301 | 0.00 |
| – | <i>Rondeletiola minor</i> AY293725 | F | <i>Sepietta neglecta</i> FJ231301 | 11.57 |
| – | <i>Sepiolo ligulata</i> AY293717 | E | <i>Sepiolo pfefferi</i> FJ231294 | 9.29 |
| D | <i>Sepiolo aurantiaca</i> AF035708 | E | <i>Sepiolo pfefferi</i> FJ231294 | 14.48 |
| G | <i>Sepietta oweniana</i> FJ231300 | G | <i>Sepietta oweniana</i> FJ231297 | 0.00 |
| F | <i>Sepietta neglecta</i> AY293722 | G | <i>Sepietta oweniana</i> FJ231297 | 0.31 |
| F | <i>Sepietta neglecta</i> AY293722 | – | <i>Euprymna stenodactyla</i> AF035704 | 3.22 |
| – | <i>Euprymna stenodactyla</i> AF035704 | G | <i>Sepietta oweniana</i> FJ231297 | 2.92 |
| G | <i>Sepietta obscura</i> AF036912 | F | <i>Sepietta neglecta</i> AY293722 | 0.46 |
| F | <i>Sepietta neglecta</i> FJ231329 | G | <i>Sepietta oweniana</i> FJ231297 | 7.60 |
| D | <i>Sepiolo robusta</i> AF035711 | D | <i>Sepiolo aurantiaca</i> AF035708 | 7.13 |
| D | <i>Sepiolo intermedia</i> AY293718 | D | <i>Sepiolo aurantiaca</i> AF035708 | 7.85 |
| – | <i>Sepietta obscura</i> AY293723 | F | <i>Sepietta neglecta</i> FJ231301 | 9.30 |
| – | <i>Sepietta obscura</i> AY293723 | G | <i>Sepietta oweniana</i> FJ231300 | 9.45 |
| B | <i>Euprymna tasmanica</i> DQ646730 | B | <i>Sepiolo atlantica</i> FJ1231302 | 1.52 |
| B | <i>Euprymna tasmanica</i> DQ646730 | – | <i>Euprymna tasmanica</i> AY293713 | 12.94 |

amino acid substitutions, it has 14 unique amino acids at positions that do not differ through sequences in the family. A BLAST search showed that this sequence is cephalopod in origin. In our opinion the large number of point mutations, indels and stopcodons could indicate a nuclear mitochondrial pseudogene (NUMT), though no bias in nucleotide composition was observed. NUMTs can also be detected by unexpected phylogenetic placement of the taxon under study (Bensasson *et al.*, 2001), but the unclear phylogenetic relationships within the Sepiolinae, the availability of only a single sequence (AY035708) and the absence of a voucher (to check the morphological identification of the specimen in question) preclude use of these methods. Since syntypes of *S. aurantiaca*

are missing, whereas according to Jereb, Mazzola & Di Stefano (1997) the records of the species are limited to those published by Jatta (1896) and Naef (1912), we consider *S. aurantiaca* a nomen dubium, referring to a species described from the Tyrrhenian Sea, most probably not occurring in the North Sea.

After exclusion of the dubious sequences AY035708 and AF035711, the maximum sequence divergence within group D is 1.77% (between AF035710 and AY293716, Table 2). Accepting a threshold of 3% sequence divergence for species status, as suggested by Hebert *et al.* (2003), this supports our assumption that group D represents only a single species, instead of seven according to Nishiguchi *et al.* (1998, 2004).

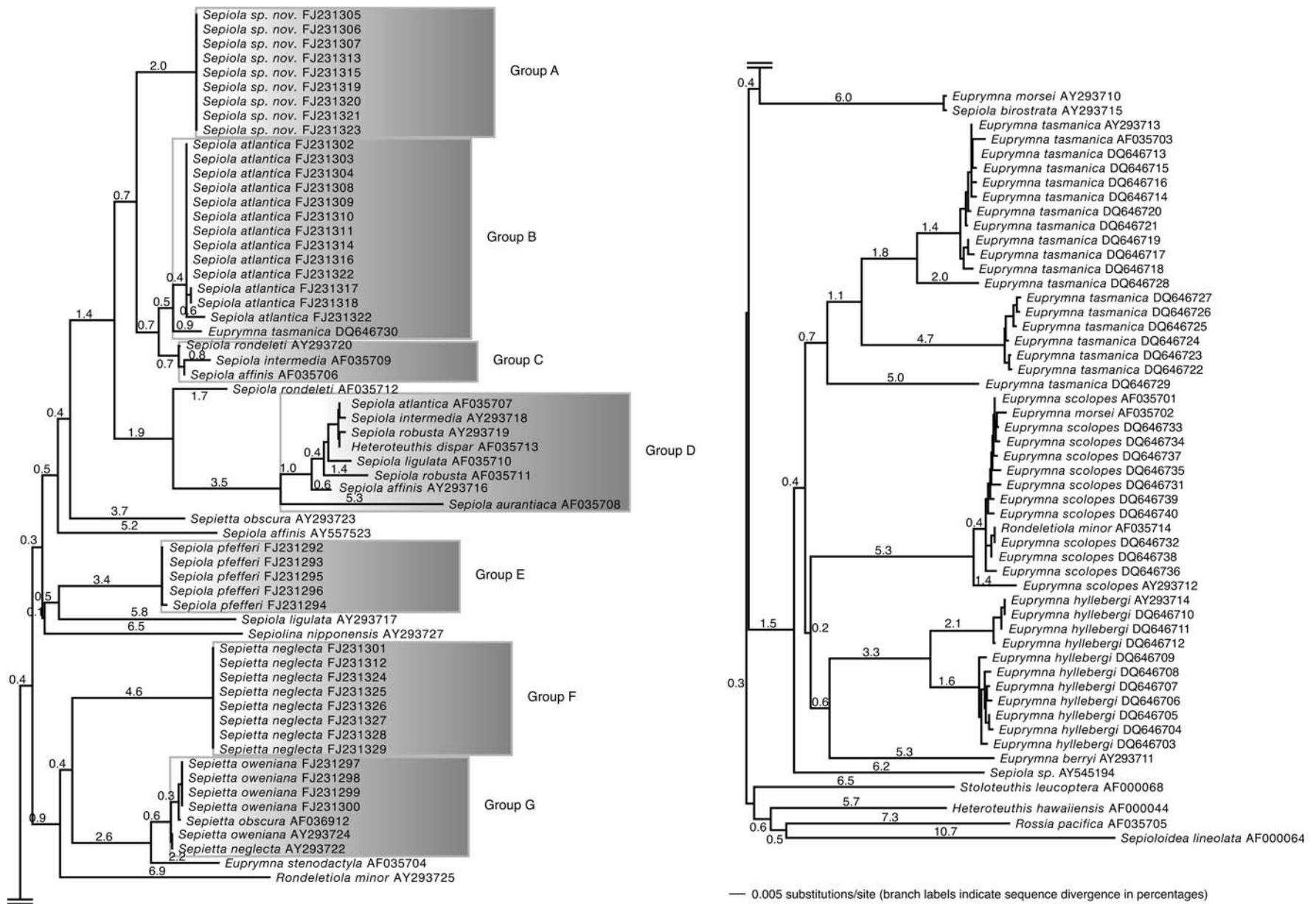


Figure 4. Neighbour-joining phylogram (branch lengths indicate sequence divergence percentages). Grey shading indicates groups of sequences of which the voucher specimens are considered conspecific in this study.

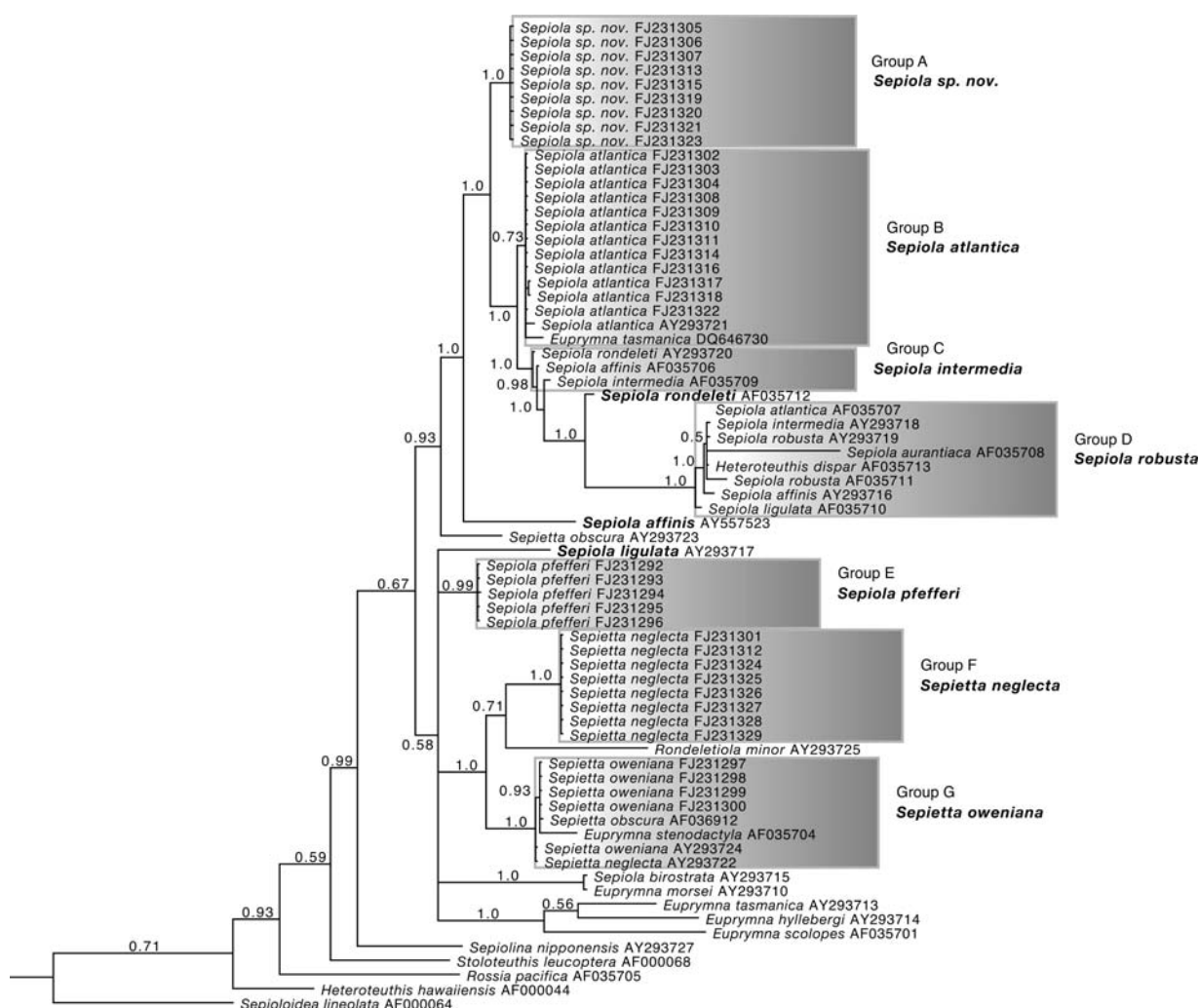


Figure 5. Bayesian phylogram with posterior probabilities. Grey shading indicates groups of sequences of which the voucher specimens are considered conspecific in this study.

Table 3. Accession numbers of conspecific specimens (according to GenBank registration) that do not form monophyletic groups.

| | |
|---------------------------|--|
| <i>Sepiola atlantica</i> | AY293721 and AF035707 |
| <i>Sepiola rondeleti</i> | AY293720 and AF035712 |
| <i>Sepiola affinis</i> | AF035706, AY557523 and AY293716 |
| <i>Sepiola intermedia</i> | AF035709 and AY293718 |
| <i>Sepiola ligulata</i> | AF035710 and AY293717 |
| <i>Sepietta obscura</i> | AY293723 and AF036912 |
| <i>Euprymna tasmanica</i> | DQ646730 and all other conspecifics (Fig. 4) |
| <i>Euprymna morsei</i> | AY293710 and AF035702 |
| <i>Rondeletiola minor</i> | AY293725 and AF035714 |

Assuming that incorrect names were assigned to GenBank entries AF035707, AY293718, AY293716, AF035710 and AF035713, group D only encompasses *S. robusta* (AY293719), which is one of the most frequently captured Mediterranean *Sepiola* species (Jereb *et al.*, 1997; Reid & Jereb, 2005) and is suggested to be the sister species (Fig. 5) of *Sepiola rondeleti* according to Bello (1998).

In our view, Grimpe's (1921) description of *S. pfefferi* was inaccurate. The photograph of the type specimen (Fig. 2B; Grimpe, 1925) clearly showed a V-shaped mantle outline

(specimen on the right), but the two syntypes on the left of it, seemed to have that structure more U- than V-shaped (as indicated for *S. aurantiaca*). Our specimens did not show the typical V-shaped mantle outline cited by Grimpe (1921). The drawing that was added (illustration on the right of Fig. 2A) is further misleading in that the size of the V-shaped incision extends up to one-third of the mantle length, whereas on the photograph the length is about one-fifth (Grimpe, 1925: specimen on the right of Fig. 2B). Even though the shape of the mantle outline is a dubious character, a clear-cut difference exists in the hectocotylus between *S. pfefferi* and *S. aurantiaca*. There are no lobes on the base of the right dorsal arm of the hectocotylus in *S. pfefferi* (Grimpe, 1921), while both Jatta (1896) and Naef (1912) depict lobes on the right dorsal arm of *S. aurantiaca* (Fig. 1). The geographical distance between the type localities of *S. aurantiaca* (Tyrrhenean Sea) and *S. pfefferi* (North Sea, East coast of England) is noteworthy as well. Despite our doubts on the correctness of the alleged *S. aurantiaca* sequence (AF035708), we have included it in our table of sequence divergences (Table 2). The *S. pfefferi* sequences (group E) differ from the sequences of all other *Sepiola* species included in this study, in particular from the alleged *S. aurantiaca* sequence. With 9.29% minimum sequence divergence (Table 2) with its sister species (Fig. 4) *S. ligulata* (AY293717), *S. pfefferi* warrants species status if the same

threshold of 3% sequence divergence (Hebert *et al.*, 2003) is imposed.

Only two *Sepietta* species, *S. neglecta* and *S. oweniana*, are known from the North Sea. Reid & Jereb (2005) indicated that these species can be difficult to distinguish morphologically; both are easily separated by their COI sequences (group F and G, Figs 4, 5, Table 2), however. Therefore, we accept *S. neglecta* and *S. oweniana* as separate taxa (Bayesian posterior probability of 1.0, Fig. 5). Our data indicated no intraspecific variation in the small samples of *S. neglecta* ($n = 8$) and *S. oweniana* ($n = 4$, Table 1, Fig. 3). Addition of the GenBank *Sepietta* sequences (AY293722, AY293724 and AF036912) once more brought misidentifications to light. These sequences form a monophyletic group with our sequences of *S. oweniana* (group G), only slightly increasing the sequence divergence within that clade to 0.31% (Table 2). If we again impose a divergence threshold of 3%, group G represents a single species. This implies that the specimens belonging to GenBank sequences AF036912 (Nishiguchi *et al.*, 1998) and AY293722 (Nishiguchi *et al.*, 2004) identified as *Sepietta obscura* and *S. neglecta*, most likely are *S. oweniana*.

A second sequence (AY293723) is referred to as *S. obscura* in GenBank. This is the only *Sepietta* sequence that indicates that this genus is not monophyletic. Except for *Rondeletiola minor* (AY293725) and *Euprymna stenodactyla* (AF035704) AY293723, the other 15 *Sepietta* sequences used in this study, form a monophyletic group (Figs 4, 5). Consequently, we are sceptical about the identification of AY293723 as *S. obscura* by Nishiguchi *et al.* (2004); we suggest it might belong to a *Sepiola* species. More sequences of reliably identified *S. obscura* are required to clarify this point.

The phylogenetic analysis (Fig. 5) and genetic distances (Table 2) show that individuals which we initially identified as *S. atlantica* represent two groups, viz. *S. atlantica* (group B) and a cryptic sister species, *Sepiola* sp. nov. (group A). Although we cannot (yet) distinguish the latter species from *S. atlantica* on morphological characters such as size, the number and arrangement of suckers and comparison of the hectocotyls, we consider it justified to report it as a separate taxon. Even within the relatively small area of the North Sea, there seems to be a sharp geographic delineation (Fig. 3) between *S. atlantica* (group B, the southern species) and *Sepiola* sp. nov. (group A, the northern species). Further investigation has to show to what extent the sharp delineation between these groups should be attributed to either different geographical distributions of both taxa indeed (southern and northern North Sea, respectively; Fig. 3) or to a preference for different depths (on average 31 vs 70 m, respectively; Table 1). The separation between groups A and B is supported by high Bayesian posterior probabilities (Fig. 5). Both analyses (Figs 4, 5) show that not group A, but group C is the sister taxon of group B. Due to the inclusion of groups C and D (Fig. 5), group B is paraphyletic. According to our data ($n = 21$) no intermediates were detected between groups A and B. The minimum sequence divergence between these clades (3.5%, Table 2) exceeds the species threshold of 3% used here, while intraspecific divergences were low (0.15% and 0% within *S. atlantica* and *Sepiola* sp. nov., respectively). Due to a transversion (T to A), all individuals of *Sepiola* sp. nov. have a methionine residue at amino acid position 160, which apparently is typical for this species (all other sepiolid COI sequences in GenBank show a leucine residue at that position).

In both analyses two GenBank sequences (AY293721 and DQ646730) are placed in group B. The specimen belonging to sequence AY293721 was collected near Vigo, Spain, and (unlike the voucher for AF035707, Table 3) seems to be correctly identified as *S. atlantica*. Inclusion of AY293721 slightly increases the maximum divergence within group B to 0.77%

(Table 2). According to Jones *et al.* (2006), the voucher for sequence DQ646730 was collected near Adelaide, Australia and represents *Euprymna tasmanica* (Pfeffer, 1884). This is surprising, since the other nineteen *E. tasmanica* sequences in GenBank form a monophyletic, if heterogeneous, group (Fig. 4). Possibly an invading specimen of *S. atlantica* (e.g. traveling in the ballast water of a ship from the North Sea) was caught and not recognized as such, this far outside its normal geographic distribution. Including DQ646730 in group B increases its maximum divergence to 1.52%. For the moment, we conclude that the sequence DQ646730 is actually from *S. atlantica* (Table 3).

The position of group C (Figs 4, 5) is unclear. The maximum sequence divergence between groups B and C (2.65%, Table 2) is inflated as both AF035709 and AF035706 have an insertion that causes a frame shift and coding region information is not provided to GenBank. As groups B and C cluster on the neighbour-joining (NJ) tree (Fig. 4), it is tempting to regard them as a single species, *S. atlantica*. The Bayesian analysis (Fig. 5), on the other hand, shows that groups C and D are monophyletic. According to the species identifications now registered in GenBank, there is an allegedly conspecific specimen elsewhere in the tree (Table 3) for each so-called species in group C. Among the three so-called *S. affinis* sequences in GenBank, AY557523 is basal to the groups A, B, C and D and AF035706 and AY293716 belong to groups C and D, respectively. The minimum sequence divergence between groups C and D (between AY293720 and AY293716) is 9.74% (Table 2), which (with a 3% threshold for species status) suggests that these groups are not conspecific. As shown (see the paragraph on group D), sequence AY293716 most likely belongs to *S. robusta*. Sequence AY557523 appears to be the sole representative of a separate group (sequence divergence >3%), maybe *S. affinis*, indicating that sequence AF035706 probably does not belong to this species.

Sepiola rondeleti and *S. intermedia* are each represented by two sequences, viz. AY293720 and AF035712, and AY293718 and AF035709. Of the former, AY293720 belongs to group C, whereas sequence AF035712 probably represents a separate group (minimum distance to group C, *S. rondeleti* AY293720 is 3.84% and to group D, *S. affinis* AY293716 is 7.97%, Table 2), maybe *S. rondeleti* indeed. Of the latter species, sequence AY293718 likely belongs to *S. robusta* (see the paragraph on group D). Consequently, the only sequence of group C that does not refer to a conspecific, more reliably identified specimen elsewhere in the tree (AF035709) was identified as *S. intermedia* by Nishiguchi *et al.* (1998). Hence we tentatively label group C as *S. intermedia*.

There is a discrepancy, which cannot yet be resolved, between the relationships shown in the NJ (Fig. 4) and the Bayesian (Fig. 5) figures, in particular regarding the position of group D relative to *S. atlantica*. We propose this can be attributed to a difference between phenetic and cladistic methods. The topology of a minimum evolution analysis (results not shown) is identical to Figure 4, whereas the topology of a maximum parsimony analysis (results not shown) is similar to that obtained by Bayesian inference (Fig. 5). As to the sequences identified as *R. minor* in GenBank, we can only conclude that they are not conspecific (Table 3) and that AF035714 is from *E. tasmanica* (Fig. 4). Given the consistency between the molecular phylogeny and our morphological determinations, and the observation that the intraspecific sequence divergences of these specimens are generally small (Table 2), we are convinced that other explanations for the observed discrepancies between the molecular phylogeny and the species identifications registered in GenBank, like incomplete lineage sorting or mitochondrial introgression, are all far less probable than our suggestion of misidentification.

In addition to contributing to our knowledge of the systematics and phylogeny of the Sepiolidae, in particular of the genera *Sepiola* and *Sepietta*, this study should serve as a warning against indiscriminate use of GenBank data, as well as being a plea for the deposition of voucher specimens in easily accessible institutional collections. DNA barcoding can be advantageous when dealing with cryptic species or for an initial assessment of biodiversity. However, to avoid confusion and error, it should be preceded or combined with a critical taxonomic analysis.

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