Species limits and phylogeography of Newportia (Scolopendromorpha) and implications for widespread morphospecies

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters

| Citation         | Edgecombe, Gregory D., Varpu Vahtera, Gonzalo Giribet, and Pipsa Kaunisto. 2015. "Species limits and phylogeography of Newportia (Scolopendromorpha) and implications for widespread morphospecies." ZooKeys (510): 65-77. doi:10.3897/zookeys.510.8573. http://dx.doi.org/10.3897/zookeys.510.8573. |
|------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Published Version| doi:10.3897/zookeys.510.8573                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| Citable link     | http://nrs.harvard.edu/urn-3:HUL.InstRepos:21461193                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| Terms of Use     | This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA                                                                                                                                                                                                                                                                                                                                                           |
Species limits and phylogeography of *Newportia* (Scolopendromorpha) and implications for widespread morphospecies

Gregory D. Edgecombe¹, Varpu Vahtera², Gonzalo Giribet³, Pipsa Kaunisto²

¹ Department of Earth Sciences, The Natural History Museum, Cromwell Road, London SW7 5BD, UK
² Zoological Museum, Department of Biology, University of Turku, Turku FI-20014, Finland
³ Museum of Comparative Zoology, Department of Organismic and Evolutionary Biology, Harvard University, 26 Oxford Street, Cambridge, MA 02138, USA

Corresponding author: Gregory D. Edgecombe (g.edgecombe@nhm.ac.uk)

Academic editor: Ivan H. Tuf | Received 12 September 2014 | Accepted 15 May 2015 | Published 30 June 2015

http://zoobank.org/C2A99E47-11F6-4F1B-B7BE-2088DCA23573

Citation: Edgecombe GD, Vahtera V, Giribet G, Kaunisto P (2015) Species limits and phylogeography of *Newportia* (Scolopendromorpha) and implications for widespread morphospecies. In: Tuf IH, Tajovský K (Eds) Proceedings of the 16th International Congress of Myriapodology, Olomouc, Czech Republic. ZooKeys 510: 65–77. doi: 10.3897/zookeys.510.8573

Abstract

The genus *Newportia* Gervais, 1847, includes some 60 nominal species distributed in the Caribbean islands and from Mexico to central South America. Modern keys to species and subspecies are available, greatly facilitating identification, but some species are based on few specimens and have incomplete documentation of taxonomically-informative characters. In order to explore genetic variability and evolutionary relationships within geographically-widespread morphospecies, specimens of *N. (N.) stolli* (Pocock, 1896) and *N. (N.) divergens* Chamberlin, 1922, two nominal species distinguished principally by differences in suture patterns on T1, were sequenced for mitochondrial 16S rRNA and cytochrome c oxidase subunit I (COI) genes from populations in southern Mexico, Guatemala, Honduras and Brazil. *N. (N.) stolli* is paraphyletic with respect to *N. (N.) divergens* Chamberlin, 1922, two nominal species distinguished principally by differences in suture patterns on T1, were sequenced for mitochondrial 16S rRNA and cytochrome c oxidase subunit I (COI) genes from populations in southern Mexico, Guatemala, Honduras and Brazil. *N. (N.) stolli* is paraphyletic with respect to *N. (N.) divergens* within a clade from Guatemala, Honduras, and Chiapas (Mexico), most trees being consistent with a single loss of a connection between the anterior transverse suture on T1, whereas specimens of "*N. (N.) stolli*" from Brazil are not closely allied to those from the Mesomerican type area. The widespread morphospecies *N. (N.) monticola* Pocock, 1890, was sequenced for the same loci from populations in Costa Rica, Colombia and Brazil, finding that specimens from these areas do not unite as a monophyletic group. Samples of *N. (N.) oreina* Chamberlin, 1915, from different regions of Mexico form geographic clusters that resolve as each other’s closest relatives. These results suggest that some widespread species of *Newportia* may...
be taxa of convenience more so than natural groupings. In several cases geographic proximity fits the phylogeny better than taxonomy, suggesting that non-monophyletic species do not result from use of inappropriate molecular markers. Molecular identification is possible for specimens missing taxonomically informative morphological characters, notably damaged specimens that lack the ultimate leg pair, a protocol that may also apply to other taxonomically difficult genera that are prone to damage (such as Cryptops).

**Keywords**
Scolopocryptopidae, Newportiinae, Neotropics, phylogeny

**Introduction**

*Newportia* Gervais, 1847 is a species-rich Neotropical genus that belongs to the family Scolopocryptopidae, encompassing blind Scolopendromorpha with 23 leg-bearing segments, pectinate second maxillary claws, and kinked and pineapple-shaped processes in the gizzard (Shelley and Mercurio 2005; Koch et al. 2009, 2010). *Newportia* has until recently been classified as one of two genera in the subfamily Newportiinae, distinguished from *Tidops* Chamberlin, 1915, by different forcipular structures (Chagas-Júnior 2011). Phylogenetic analyses based on multi-locus molecular sequence data have, however, indicated that *Tidops* nests within *Newportia* rather than being the sister group, as does another clade that had been assigned to a separate subfamily, the Mesoamerican Ectonocryptopinae (Vahtera et al. 2013).

The geographic distribution of *Newportia* (including *Tidops*, *Ectonocryptops* Crabill, 1977, and *Ectonocryptoides* Shelley & Mercurio, 2005 as subgenera: Vahtera et al. 2013) extends from northern Mexico throughout Central America and the Caribbean islands to Paraguay. Most species of *Newportia* have tarsus 2 of the ultimate leg divided into five to nearly 40 tarsomeres, or with indistinct separation of tarsi 1 and 2. Currently some 60 nominal species or subspecies are recognised (Minelli et al. 2006 and onwards; Schileyko 2013). In many species, diagnostic features involve the spinose processes on the ultimate prefemora and femora and the number of tarsomeres, all inconvenient characters because individuals frequently lose these legs when collected.

We propose a solution to the taxonomic impediment of missing ultimate legs by using mitochondrial sequence data to supplement identifications. We also explore phylogeographic patterns within and between select species of *Newportia* from Mexico and Central America using parsimony and maximum likelihood methods. The resultant phylogenies allow the taxonomic value of purportedly diagnostic morphological characters to be evaluated and for the limits of morphospecies to be tested.

**Methods**

Thirty-four specimens of *Newportia* from Mexico, Guatemala, Honduras, and Costa Rica were sorted mostly from collections made by the LLAMA (Leaf Litter Survey of
Species limits and phylogeography of Newportia (Scolopendromorpha)...

Mesoamerica) project, deposited in the Museum of Comparative Zoology (MCZ), Harvard University, Cambridge Massachusetts, USA and accessible through the dedicated data base MCZbase (http://mczbase.mcz.harvard.edu). All tissues were fixed in absolute ethanol and thus were amendable to DNA sequencing.

Identifications were made using the most recent key for *N. (Newportia)* (Schileyko, 2013), supplemented with taxonomic descriptions in modern literature (Schileyko and Minelli 1998; Chagas-Júnior and Shelley 2003), standard monographs (Attems 1930), original descriptions, and examination of type material designated by R. I. Pocock in The Natural History Museum (London) and or by R. V. Chamberlin in the MCZ.

LLAMA specimens keyed to either *N. (N.) monticola* Pocock, 1890, *N. (N.) stolli* (Pocock, 1896), *N. (N.) oreina* Chamberlin, 1915, or *N. (N.) divergens* Chamberlin, 1922. All LLAMA specimens were sequenced for two mitochondrial loci: 16S rRNA and cytochrome *c* oxidase subunit I (COI). These loci were selected because they vary both within and between species, and even between individuals from geographically close populations. The 34 LLAMA samples were supplemented with *N. (Newportia)* and *N. (Ectonocryptoides)* sequences from our previous work (Vahtera et al. 2013), nine new Newportia specimens from five localities in Amazonas and Roraima, Brazil, and novel sequences for an individual of *N. (N.) pusilla* Pocock, 1893, from Ecuador (see Table 1 for morphospecies determinations and locality data).

Total DNA was extracted from the legs utilizing the NucleoSpin®Tissue kit (Macherey-Nagel). Samples were incubated overnight. PCR amplifications were performed with illustra TM PuReTaq TM Ready-To-GoTM PCR Beads (GE Healthcare). The COI fragments were amplified using primer pair HCO1490 (Folmer et al. 1994) and HCOout (Carpenter and Wheeler 1999) and the 16S rRNA fragments using primer pair 16Sa/16Sb (Xiong and Kocher 1991; Edgecombe et al. 2002). The normal amplification cycle for COI consisted of an initial denaturation step (2 min at 95 °C), followed by 35 cycles of denaturation (1 min at 95 °C), annealing (1 min at 43 °C) and extension (1.5 min at 72 °C), followed by a final extension step (4 min at 72 °C). For the 16S rRNA fragment the cycle consisted of an initial denaturation step (2 min at 94 °C), followed by 35 cycles of denaturation (30 s at 94 °C), annealing (30 s min at 43 °C) and extension (1 min at 72 °C), followed by a final extension step (7 min at 72 °C). Visualization of the PCR products was done by 1 % agarose electrophoresis using Midori Green Advanced DNA Stain and FastGene® GelPic LED Box (Nippon Genetics, GmbH).

Samples were purified using ExoSAP-IT (Affymetrix) and sent to FIMM (Institute for Molecular Medicine Finland) for sequencing. Chromatograms were visualized and assembled using Sequencer 5.0.1 (Gene Codes Corp., Ann Arbor, Michigan, USA). Sequence alignment editor Se-Al (Rambaut 1996) was used to visualize the sequences simultaneously. GenBank registrations for new sequences are listed in Table 1.

Parsimony analysis was conducted with POY ver. 5.1.1 (Wheeler et al. 2014) run in 16 nodes in the high-performance supercluster Taito at CSC (IT-Center of Science), Finland. A timed search of three hours was first performed on the unaligned data set.
| Species                        | Voucher ID number | Lab code | Country (State) | Country (State) | 16S Accession | COI Accession | Lat. (degrees) | Long. (degrees) |
|-------------------------------|------------------|---------|-----------------|-----------------|---------------|---------------|----------------|----------------|
| Newportia adisi               | 130770           | -       | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | -              | -              |
| Newportia collis              | 18827            | -       | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 2.93355S       | 59.96611W      |
| Newportia quadrimeropus       | 99154            | 82      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | -              | -              |
| Newportia divergens           | 98078            | 81      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 0.99185N       | 62.15915W      |
| Newportia divergens           | 99129            | 75      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 14.6137724N    | 89.0697228W    |
| Newportia divergens           | 88919            | 85      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 14.18139N      | 87.53225W      |
| Newportia ernstei             | 94265            | 57      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 23.0344N       | 99.18697W      |
| Newportia monticola           | 98065            | 40      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 8.40607N       | 82.59333W      |
| Newportia oreina              | 94726            | 49      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 8.78658N       | 82.59333W      |
| Newportia pusilla             | 88758            | 40      | MCZ, Brazil (Amazonas) | -               | KP099504      | HQ405349      | 0.6083333N     | 77.8825W       |
| Species          | Voucher ID number | Lab code | Country (State) | COI   | 16S       | Lat. (degrees) | Long. (degrees) |
|------------------|-------------------|----------|-----------------|-------|-----------|----------------|-----------------|
| N. sp.           | 81282             | 54       | MCZ             | Costa Rica | KP099544 | 8.94997N       | 82.83757W       |
| N. sp.           | 81635             | 22       | MCZ             | Brazil (Roraima) | KP099502 | 0.99595S       | 61.59094W       |
| N. sp.           | 106516            | 37       | MCZ             | Brazil (Roraima) | KP099510 | 10.18518N      | 62.0822W        |
| N. sp.           | 81560             | 44       | MCZ             | Brazil (Roraima) | KP099467 | 14.19185N      | 91.14958N       |
| N. sp.           | 79982             | 47       | MCZ             | Brazil (Roraima) | KP099463 | 16.1353333N    | 90.90146667W    |
| N. sp.           | 80143             | 48       | MCZ             | Mexico (Chiapas) | KP099469 | 16.93583N      | 89.676667W      |
| N. sp.           | 8175              | 78       | MCZ             | Mexico (Chiapas) | KP099472 | 1.2173667N     | 93.14999W       |
| N. sp.           | 80175             | 48       | MCZ             | Mexico (Chiapas) | KP099464 | 16.97316667N   | 91.58991667W    |
| N. sp.           | 81563             | 50       | MCZ             | Mexico (Chiapas) | KP099465 | 15.08405N      | 89.94991W       |
| N. sp.           | 82008             | 71       | MCZ             | Mexico (Chiapas) | KP099471 | 16.75181N      | 92.1062W        |
| N. sp.           | 99225             | 71       | MCZ             | Mexico (Chiapas) | KP099513 | 15.09708N      | 90.21480799W    |
| N. sp.           | 99279             | 78       | MCZ             | Guatemala       | KP099473 | 15.02124133N   | 89.5458833N     |
| N. sp.           | 18826             | 88a      | MCZ             | Brazil (Roraima) | KP099477 | 16.444446667N  | 89.5487322W     |
| N. sp.           | 18830             | 93a      | MCZ             | Brazil (Roraima) | KP099478 | 17.24935734N   | 89.60290417W    |
| N. stolli        | 18827             | 95a      | MCZ             | Brazil (Roraima) | KP099514 | 16.58991667W   | 91.58991667W    |
| N. stolli        | 89566             | 99       | MCZ             | Mexico (Chiapas) | KP099512 | 15.02124133N   | 89.5458833N     |
| N. stolli        | 89321             | 100      | MCZ             | Guatemala       | KP099479 | 15.21318939N   | 90.21213699W    |
| N. stolli        | 89396             | 102      | MCZ             | Guatemala       | KP099480 | 16.6444466667N | 89.5458833N     |
| N. stolli        | 89355             | 103      | MCZ             | Guatemala       | KP099515 | 15.21318939N   | 90.21213699W    |
| N. stolli        | 89306             | 104      | MCZ             | Guatemala       | KP099475 | 16.444446667N  | 89.5458833N     |
| N. stolli        | 13057             | -        | MCZ             | Italy           | KP099547 | 13.36111N      | 92.63490655W    |

*Cryptops punicus* 130604 - MCZ - 61.62111N 8.54909E

*Scolopocryptops mexicanus* 105626 - MCZ - 1.336111N 77.263095W
The resulting tree was used as the starting tree for the next round in which an additional timed search of six hours was performed. Parameter set 111 (indel/transversion and transversion/transition costs all equal) was used throughout the searches and branch lengths were reported using the newly implemented command “report ("file_name.tre", trees:(total, branches:true))”. Nodal support was calculated using parsimony jackknifing (Farris et al. 1996).

Additional analyses used a probabilistic approach with the maximum likelihood program RAxML ver. 8.0.22 (Stamatakis 2014). For these, multiple sequence alignments (MSA) were first estimated with MUSCLE ver. 3.6 (Edgar 2004) and then trimmed using Gblocks ver. 0.91b (Castresana 2000; Talavera and Castresana 2007) to remove areas of ambiguous alignment. Since COI sequences showed no length variation, they were not trimmed in Gblocks. The amount of 16S rRNA data that remained after trimming was 59% of the original 585 positions. The two data sets were concatenated using SequenceMatrix (Vaidya et al. 2011) and the concatenated data were analyzed with RAxML in the CIPRES Science Gateway (Miller et al. 2010). A unique general time reversible (GTR) model was specified for each partition independently. Nodal support was estimated using the rapid bootstrap algorithm (applying the Majority Rule Criterion) using the GTR-CAT model (Stamatakis et al. 2008).

**Results**

The combined analysis of both COI and 16S fragments using parsimony as the optimality criterion resulted in two most parsimonious (MP) trees of length 4625 steps.

![Map of Mesoamerica, the Caribbean and northern South America showing geographic distribution of Newportia specimens analyzed herein (see Table 1 for coordinates of samples).](image-url)
Species limits and phylogeography of Newportia (Scolopendromorpha)...

Figure 2. Strict consensus of two optimal cladograms for Newportia under parameter set 111 for parsimony (POY) analysis. Abbreviations: BRA, Brazil; COL, Colombia; CR, Costa Rica; DR, Dominican Republic; ECU, Ecuador; FRG, French Guiana; GUA, Guatemala; HON, Honduras; MEX, Mexico.

The strict consensus tree (Fig. 2) shows these two trees are almost identical, differing only in the placement of two Brazilian specimens of *N. (N.) stolli* in relation to each other. Comparing strongly supported clades, the maximum likelihood tree (lnL -14054.372302: Fig. 3) shows the same major geographic and taxonomic groupings as the parsimony tree. This congruence is noteworthy because the data sets analyzed under these two optimality criteria were different (unaligned in POY and analyzed using the concept of dynamic versus static homologies with some regions removed in RAxML), as are the resampling methods (jackknifing and bootstrapping, respectively). Parts of the trees that are incongruent between the two analyses involve nodes that received low resampling supports in both analyses (e.g., the positions of *N. (N.) adisi* and Brazilian specimen 89b relative to other species). Both analyses depict substantial branch lengths both within and between species, with only a few instances of no (or minimal) variation between specimens from the same or geographically close populations.

As in previous analyses based on sparser sampling for Newportia (Vahtera et al. 2013), *Tidops (T. collaris)* and *Ectonocryptoides (E. quadrimeropus)* nest within Newportia in all
analyses. Specifically, they unite with *Newportia* (*Newportia*) spp. that inhabit the same geographic region i.e., *N. (T.) collaris* from the Brazilian Amazon groups within a clade composed of species of *Newportia* (*Newportia*) from there, whereas *N. (E.) quadrimeropus* from Jalisco, Mexico, groups with the Mexican *N. (N.) oreina*. These results reinforce proposals to classify *Tidops*, *Ectonocryptoides* and presumably allied *Ectonocryptops* within *Newportia* and to regard *Ectonocryptopinae* as subordinate to *Newportiinae* (Vahtera et al. 2013). The traditional classification of *N. (Tidops)* and *N. (Ectonocryptoides)* as separate genera because of their obvious phenotypic differences from *N. (Newportia)* might have predicted that they would be markedly different from *N. (Newportia)* genetically. However, neither *N. (Tidops) collaris* nor *N. (Ectonocryptoides) quadrimeropus* depict long branch divergences from their closest relatives with respect to the studied loci, indeed being shorter than some population-level branches within species.

*Newportia oreina* consists of two geographical clades and this division is found in both parsimony and likelihood analyses; one clade consists of all specimens from Tamaulipas (JK, BS 100) and the other of ones from Oaxaca (JK 100, BS 98). Interestingly, *N. (Ectonocryptoides) quadrimeropus* forms a well-supported (JK 99, BS 73) clade with the *N. (N.) oreina* populations from Oaxaca, rendering *N. (N.) oreina* para-

---

**Figure 3.** Maximum likelihood tree (lnL = -14054.372302). Abbreviations for countries as in Fig. 2.
phyletic with respect to *Ectonocryptoides* (and presumably *Ectonocryptops*). A previous scolopendromorph phylogeny (Vahtera et al. 2013) had also indicated affinity between *N.* (*N.*) *oreina* and *N.* (*Ectonocryptoides*) *quadrimeropus*; analyses based on combined molecular and morphological data resolved them as sister-groups, although only one individual of each was then available. We note that *N. oreina* possesses a shorter tarsus than most congeners. The phylogeny interprets the ancestral condition of the ultimate leg tarsi of *Newportia* as being elongate and divided into tarsomeres, with the relatively short tarsus 2 of *N.* (*N.*) *oreina* being a possible precursor to the stout tarsi of the submerged taxon, “Ectonocryptopinae”. This transformation series increases the plausibility of the subclavate “ectonocryptopine” ultimate legs being derived from an ancestor with flagelliform tarsi, a result that was already strongly signaled by molecular phylogenies (Vahtera et al. 2013) and is reinforced by the current trees.

A Mesoamerican clade uniting *N.* (*N.*) *stolli* and *N.* (*N.*) *divergens* from Mexico (Chiapas), Guatemala and Honduras is recovered in both parsimony and likelihood analyses (Figs 2, 3), though resampling methods did not strongly support it (JK <50, BS 57). *N.* (*N.*) *divergens* is resolved as monophyletic in the POY analyses but is nested within a paraphyletic *N.* (*N.*) *stolli*, implying a single loss of the median part of the anterior transverse suture on T1 (Fig. 2). However, there is no jackknife support for the *divergens* clade. In contrast, the likelihood analysis did not support monophyly of *N.* (*N.*) *divergens*; six individuals from Guatemala and Honduras resolve as a well-supported clade (BS 98), but two others from Honduras (81, 82) are grouped with two Mexican *N.* (*N.*) *stolli* specimens, albeit with weak nodal support.

Specimens identified as *N.* (*N.*) *stolli* from the Brazilian Amazon do not unite with supposed congeners from Mesoamerica but are instead most closely related to other taxa from the same region, i.e., a specimen identified as *N.* (*N.*) *monticola* (91) and *N.* (*Tidops*) *collaris*. This result implies that *N.* (*N.*) *stolli* is polyphyletic and an indistinct segmentation of ultimate tarsus 2 has multiple (convergent) origins. This character had once served as the basis for recognising a subgenus *N.* (*Scolopendrides*), e.g., in the classification of Bücherl (1974), but this taxon is not used in current classifications (Schileyko and Minelli 1998). We re-examined the *N.* (*N.*) *stolli* specimens again in light of the signal for non-monophyly in the phylogenetic analysis, attempting to recognize any morphological character(s) that would separate the specimens from Brazil from those in Mesoamerica. However, we found no distinctive characters between the samples; the specimens appear to be morphologically indistinguishable and using the existing keys they would all be identified as *N.* (*N.*) *stolli* with confidence.

Costa Rican specimens of *N.* (*N.*) *monticola* unite as a monophyletic group (JK 100, BS 99) in both analyses. In the maximum likelihood tree (Fig. 3) a Colombian specimen of *N.* (*N.*) *monticola* (103974) is resolved as a sister taxon to the Costa Rican clade but this relationship is not found in the parsimony tree (Fig. 2). In neither analysis did a Brazilian specimen identified as *N.* (*N.*) *monticola* unite with the other supposed conspecifics.

The two included specimens of *N.* (*N.*) *pusilla*, one from Ecuador (specimen 86) and the other from Brazilian Amazonas (specimen 90), likewise do not form a clade
but instead are situated in different parts of the tree. The Brazilian specimen conforms to “Amazonian type *pusilla*” of Schileyko and Minelli (1998), characterized by rudimentary paramedian sutures on T1 (in contrast to their complete absence in other populations). Both analyses group this Brazilian specimen together with *N. (N.) longitarsis stechowi* but since there is no strong resampling support in either analysis (JK <50, BS 69), the question about its identity and closest relative remains unclear.

We also included a few *Newportia* specimens that could not be identified morphologically since they lacked ultimate legs, were juveniles, or did not key out to any known species. A specimen (54) from Costa Rica has a unique character combination and is apparently a distinct species but lacks its ultimate legs. In the POY analysis it groups together, although with weak support, with the Costa Rican *N. (N.) monticola* clade. A very distinctive Brazilian specimen (89b) with all tarsi bipartite and tarsus 2 of the ultimate leg undivided groups at the base of the Mexican *N. (N.) oreinal/N. (E.) quadrimeropus* clade in the parsimony analysis. However, there is poor resampling support for this grouping and it is instead allied to species with indistinctly segmented ultimate tarsus 2 and the Brazilian clade in the likelihood tree. The poor support values and topological instability under different analytical conditions render the affinities of this undescribed species uncertain.

**Discussion**

Some of the specimens used in this study were either of small size because of the collection methods employed (and thus may not have been appropriate for keying using traditional criteria formulated for mature specimens) or were missing their taxonomically-informative ultimate legs. Nonetheless, several such specimens could be identified with a high degree of accuracy because their sequence data placed them within clades whose nomenclature could be established based on standard external morphological characters. An example is provided by a juvenile from Brazil (92) that is in poor condition and cannot be identified to species. However, the analysis shows it to be a juvenile of a Brazilian clade assigned to *N. (N.) stolli*. This approach is likely to be valuable in other groups of taxonomically-difficult centipedes that rely heavily on characters of the ultimate leg pair but often lack those legs in fixed specimens, such as *Cryptops*, where the numbers of tibial and tarsal saw teeth are fundamental taxonomic characters. The identification of developmental stages or adults without key taxonomic characters is becoming standard for many groups of animals, including other arthropod groups, such as insects (Monaghan et al. 2009; Gattolliat and Monaghan 2010) and arachnids (Fernández et al. 2014).

Some morphologically delimited species were found to be monophyletic groups, like *N. (N.) divergens* in the parsimony analysis, but others were paraphyletic or polyphyletic. This could be interpreted as a failure of the taxonomic characters traditionally used to delimit species or a failure in reconstructing an accurate tree by the markers selected. The second option is unlikely for the reasons outlined below, especially the biogeographical patterns exhibited in many clades where “distinct” species from the same regions tend
to cluster together and not with their supposed conspecifics from other geographical regions. In particular \textit{N. (N.) stolli} formed a series of geographic groupings that in part were paraphyletic with respect to sympatric species (specifically, to \textit{N. (N.) divergens} in Mesoamerica) or in other cases were found to be distantly related (Brazilian “\textit{N. (N.) stolli}”). The first pattern is consistent with \textit{N. (N.) stolli} being a grade united by a plesiomorphy (a continuous anterior transverse suture on T1), some parts of which are most closely related to a species defined by an apomorphic state (i.e., loss of the median extent of the anterior transverse suture). The tree topology, however, suggests that the Brazilian specimens identified as \textit{N. (N.) stolli} are misidentified. \textit{Newportia (N.) monticola} is likewise a questionable taxon, the monophyletic Costa Rican group never uniting with a specimen of the same putative species from Brazil and only variably so with one from Colombia. Brazilian \textit{N. (N.) monticola} and \textit{N. (N.) stolli} unite in a well-supported clade (JF and BS 100), indicating that, in this instance, geography is a better predictor of relationships than taxonomy. It is noteworthy that \textit{N. (N.) stolli} and \textit{N. (N.) monticola} are among the most geographically widespread “species” of \textit{Newportia}, but our results suggest that the wide distribution is partly an artifact of morphologically-based identifications. The same evidently applies to \textit{N. (N.) pusilla}, a morphospecies that is regarded as ranging from St. Vincent through Colombia to the Brazilian Amazon (Schileyko and Minelli 1998; Chagas-Júnior et al. 2014). Polyphyly of this species in the molecular trees suggests that its diagnostic characters (absent or rudimentary paramedian sutures on T1 and a lack of ventral spinose processes on the ultimate leg femora) evolved convergently in different regions.

Centipede systematics, still strongly influenced by mid 20th Century conceptualisations of species (see Edgecombe 2007), primarily assumes polymorphic and geographically widespread entities. The existing concepts that \textit{N. (N.) monticola} and \textit{N. (N.) stolli} are widespread throughout much of Central and South America exemplify where morphospecies do not appear to correspond to clades but rather are classes defined by combinations of characters. In these instances, molecular tools may prove to be invaluable for species delimitations, and novel morphological characters will need to be identified to rediagnose polyphyletic species.

\textbf{Acknowledgements}

Most samples were collected from the LLAMA (Leaf Litter Survey of Mesoamerica) survey, a collecting program supported by NSF grant DEB-0640015 to John Longino. Laboratory expenses were covered by a grant from the Finnish Entomological Society. Funding for GG’s fieldwork in Brazil was provided by the National Geographic Society to the Amazon (2012), and by US National Science Foundation grant #1144417 (Collaborative Research: ARTS: Taxonomy and systematics of selected Neotropical clades of arachnids) to GG and G. Hormiga. We thank the University of Turku for supporting GDE’s visit in January 2014, and CSC – IT Center for Science Ltd. for the allocation of computational resources. Emil Vahtera and Kari Kaunisto assisted with figures. Input from Arkady Schileyko and Rowland Shelley as referees improved the manuscript.
References

Attems C (1930) Myriapoda 2. Scolopendromorpha. In: Schulze FE, Kükenthal W (Eds) Das Tierreich, 54. Walter de Gruyter, Berlin, 1–308.

Bücherl W (1974) Die Scolopendromorpha der Neotropischen Region. In: Blower JG (Ed.) Myriapoda. Symposia of the Zoological Society of London 32. Academic Press, 99–133.

Carpenter JM, Wheeler WC (1999) Towards simultaneous analysis of morphological and molecular data in Hymenoptera. Zoologica Scripta 28: 251–260. doi: 10.1046/j.1463-6409.1999.00009.x

Castresana J (2000) Selection of conserved blocks from multiple alignments for their use in phylogenetic analysis. Molecular Biology and Evolution 17: 540–552. doi: 10.1093/oxfordjournals.molbev.a026334

Chagas-Júnior A (2011) A review of the centipede genus *Tidops* Chamberlin (Scolopendromorpha, Scolopocryptopidae, Newportiinae). International Journal of Myriapodology 5: 63–82. doi: 10.3897/ijm.5.1649

Chagas-Júnior A, Chaparro E, Galvis Jiménez S, Triana Triana HD, Flórez E, Sícoli Seoane JC (2014) The centipedes (Arthropoda, Myriapoda, Chilopoda) from Colombia: Part 1. Scutigeromorpha and Scolopendromorpha. Zootaxa 3779: 133–156. doi: 10.11646/zootaxa.3779.2.2

Chagas-Júnior A, Shelley RM (2003) The centipede genus *Newportia* Gevais, 1847, in Mexico: description of a new troglomorphic species; redescription of *N. sabina* Chamberlin, 1942; revival of *N. azteca* Humbert & Saussure, 1869; and a summary of the fauna (Scolopendromorpha: Scolopocryptopidae: Newportiinae). Zootaxa 379: 1–20.

Crabill RE Jr. (1977) A new cryptopid genus, with key to the genera known to occur in North America including Mexico (Chilopoda: Scolopendromorpha: Cryptopidae). Proceedings of the Entomological Society of Washington 79: 346–349.

Edgar RC (2004) MUSCLE: multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res. 32(5): 1792–1797. doi: 10.1093/nar/gkh340

Edgecombe GD (2007) Centipede systematics: progress and problems. Zootaxa 1668: 327–341.

Edgecombe GD, Giribet G, Wheeler WC (2002) Phylogeny of Henicopidae (Chilopoda: Lithobiomorpha): a combined analysis of morphology and five molecular loci. Systematic Entomology 27: 31–64. doi: 10.1046/j.0307-6970.2001.00163.x

Farris JS, Albert VA, Källersjö M, Lipscomb D, Kluge AG (1996) Parsimony jackknifing outperforms neighbor-joining. Cladistics 12: 99–124. doi: 10.1111/j.1096-0031.1996.tb00196.x

Fernández R, Vélez S, Giribet G (2014) Linking genetic diversity and morphological disparity: biodiversity assessment of a highly unexplored family of harvestmen (Arachnida : Opiliones : Neopilionidae) in New Zealand. Invertebrate Systematics 28: 590–604. doi: 10.1071/IS14029

Gattolliat JL, Monaghan MT (2010) DNA-based association of adults and larvae in Baetidae (Ephemeroptera) with the description of a new genus *Adnoptilum* in Madagascar. Journal of the North American Benthological Society 29: 1042–1057. doi: 10.1899/09-119.1

Koch M, Edgecombe GD, Shelley RM (2010) Anatomy of *Ectonocryptoides* (Scolopocryptopidae: Ectonocryptopinae) and the phylogeny of blind Scolopendromorpha (Chilopoda). International Journal of Myriapodology 3:51–81. doi: 10.1163/187525410X12578602960344
Species limits and phylogeography of Newportia (Scolopendromorpha)... 77

Koch M, Pärschke S, Edgecombe GD (2009) Phylogenetic implications of gizzard morphology in scolopendromorph centipedes (Chilopoda). Zoologica Scripta 38: 269–288. doi: 10.1111/j.1463-6409.2008.00372.x

Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for inference of large phylogenetic trees. In: Proceedings of the Gateway Computing Environments Workshop (GCE), 14 Nov. 2010, New Orleans, LA, 1–8. doi: 10.1109/GCE.2010.5676129

Minelli A, Bonato L, Dioguardi R, Chagas-Júnior A, Edgecombe GD, Lewis JGE, Pereira LA, Shelley RM, Stoev P, Uliana M, Zapparoli M (2006 and onwards) CHILOBASE. A web resource for Chilopoda taxonomy. http://chilobase.bio.unipd.it

Monaghan MT, Wild R, Elliot M, Fujisawa T, Balke M, Inward DJ, Lees DC, Ranivosolo R, Eggleton P, Barraclough TG, Vogler AP (2009) Accelerated species inventory on Madagascar using coalescent-based models of species delineation. Systematic Biology 58: 298–311. doi: 10.1093/sysbio/syp027

Schileyko AA (2013) A new species of Newportia Gervais, 1847 from Puerto Rico, with a revised key to the species of the genus (Chilopoda, Scolopendromorpha, Scolopocryptopidae). ZooKeys 276: 39–54. doi: 10.3897/zookeys.276.4876

Schileyko AA, Minelli A (1998) On the genus Newportia Gervais, 1847 (Chilopoda: Scolopendromorpha: Newportiidae). Arthropoda Selecta 7: 265–299.

Shelley RM, Mercurio R (2005) Ectonocryptoides quadrimeropus, a new centipede genus and species from Jalisco, Mexico; proposal of Ectonocryptopinae, analysis of subfamilial relationships, and a key to subfamilies and genera of the Scolopocryptopidae (Scolopendromorpha). Zootaxa 1094: 25–40.

Stamatakis A, Hoover P, Rougemont J (2008) A rapid bootstrap algorithm for the RAxML web servers. Systematic Biology 57: 758–771. doi: 10.1080/10635150802429642

Stamatakis A (2014) RAxML Version 8: A tool for Phylogenetic Analysis and Post-Analysis of Large Phylogenies. Bioinformatics (2014) 30(9): 1312–1313. doi: 10.1093/bioinformatics/btu033

Talavera G, Castresana J (2007) Improvement of phylogenies after removing divergent and ambiguously aligned blocks from protein sequence alignments. Systematic Biology 56: 564–577. doi: 10.1080/10635150701472164

Vahtera V, Edgecombe GD, Giribet G (2013) Phylogeny of scolopendromorph centipedes: Can denser taxon sampling improve an artificial classification? Invertebrate Systematics 27: 578–602. doi: 10.1071/IS13035

Vaidya G, Lohman DJ, Meier R (2011) SequenceMatrix: concatenation software for the fast assembly of multi-gene datasets with character set and codon information. Cladistics 27: 171–180. doi: 10.1111/j.1096-0031.2010.00329.x

Wheeler WC, Lucaroni N, Hong L, Crowley LM, Varón A (2014) POY version 5: phylogenetic analysis using dynamic homologies under multiple optimality criteria. Cladistics 31: 189–196. doi: 10.1111/cla.12083

Xiong B, Kocher TD (1991) Comparison of mitochondrial DNA sequences of seven morphospecies of black flies (Diptera: Simuliidae). Genome 34: 306–311. doi: 10.1139/g91-050