



The First Phylogenetic Analysis of Palpigradi (Arachnida)—The Most Enigmatic Arthropod Order

Citation

Gonzalo, Gilbert, McIntyre Erin, Christian Erhard, Espinasa Luis, Ferreira Rodrigo L., Francke Óscar F., Harvey Mark S., Isaia Marco, Kováč Ĺubomír, McCutchen Lynn, Souza Maysa F. V. R., and Zagmajster Maja. 2014. "The First Phylogenetic Analysis of Palpigradi (Arachnida) – The Most Enigmatic Arthropod Order." Invertebrate Systematics 28: 350–360. doi: 10.1071/IS13057

Published Version

doi:10.1071/IS13057

Permanent link

http://nrs.harvard.edu/urn-3:HUL.InstRepos:12313557

Terms of Use

This article was downloaded from Harvard University's DASH repository, and is made available under the terms and conditions applicable to Open Access Policy Articles, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#OAP

Share Your Story

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

Accessibility

1 The first phylogenetic analysis of Palpigradi (Arachnida)—the most

- 2 enigmatic arthropod order
- 3 Gonzalo Giribet^{A,K}, Erin McIntyre^A, Erhard Christian^B, Luis Espinasa^C, Rodrigo L. Ferreira^D, Óscar F.
- 4 Francke^E, Mark S. Harvey^F, Marco Isaia^G, Ľubomīr Κονάč^H, Lynn McCutchen^I, Maysa F. V. R.
- 5 Souza^D and Maja Zagmajster^J

6

- 7 AMuseum of Comparative Zoology, Department of Organismic and Evolutionary Biology, Harvard
- 8 University, 26 Oxford Street, Cambridge, MA 02138, USA.
- 9 BInstitut für Zoologie, Universität für Bodenkultur, Gregor-Mendel-Straße 33, 1180 Wien, Austria.
- 10 ^cSchool of Science, Marist College, 3399 North Road, Poughkeepsie, New York, USA.
- 11 De Centro de Estudos em Biologia Subterrânea, Departamento de Biologia, Universidade Federal
- de Lavras, Lavras, MG. CEP 37200-000, Brazil.
- 13 ^EColección Nacional de Arácnidos, Instituto de Biologia, UNAM, Apartado Postal 70-153, C. P.
- 14 04510, Mexico, D. F., Mexico.
- 15 FDepartment of Terrestrial Zoology, Western Australian Museum, Locked Bag 49, Welshpool DC,
- 16 WA 6986, Australia.
- 17 ^GDipartimento di Scienze della Vita e Biologia dei Sistemi, Università di Torino, Via Accademia
- 18 Albertina 13, 10123 Torino, Italy.
- ¹⁹ ^HDepartment of Zoology, Institute of Biology and Ecology, Faculty of Science, P. J. Šafárik
- 20 University, Moyzesova 11, 040 01 Košice, Slovakia.
- ¹Department of Biology, Kilgore College, 1100 Broadway, Kilgore, TX 75662, USA.
- ¹SubBioLab, Department of Biology, Biotechnical Faculty, University of Ljubljana, Večna pot 111,
- 23 SI-1000 Ljubljana, Slovenia.
- 24 ^KCorresponding author. Email: ggiribet@g.harvard.edu

Abstract. Palpigradi are a poorly understood group of delicate arachnids, often found in caves or other subterranean habitats. Concomitantly, they have been neglected from a phylogenetic point of view. Here we present the first molecular phylogeny of palpigrades based on specimens collected in different subterranean habitats, both endogean (soil) and hypogean (caves), from Australia, Africa, Europe, South America and North America. Analyses of two nuclear ribosomal genes and COI under an array of methods and homology schemes found monophyly of Palpigradi, Eukoeneniidae, and a division of Eukoeneniidae into four main clades, three of which include samples from multiple continents. This supports either ancient vicariance or long-range dispersal, two alternatives we cannot distinguish with the data at hand. In addition, we show that our results are robust to homology scheme and analytical method, encouraging further use of the markers employed in this study to continue drawing a broader picture of palpigrade relationships.

- 39 Additional keywords: Arachnida, micro-whip scorpions, palpigrades, speleobiology,
- 40 biogeography.

Introduction

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

The arachnid order Palpigradi (micro-whip scorpions or palpigrades) is one of the smallest, rarest and most neglected groups of terrestrial arthropods, and one of the last arachnid orders to be discovered—it was first reported only in 1885 (Grassi and Calandruccio 1885). The first photographs of living palpigrades did not appear published until the first decade of the 21st century (Kováč et al. 2002; Beccaloni 2009). Additionally, only a handful of DNA sequence data are available in GenBank; with only 64 sequences, 56 are for Prokoenenia wheeleri (Rucker, 1901), a species that was part of a multi-gene phylogeny of arthropods (Regier et al. 2010), while the remaining eight sequences are unidentified specimens from three studies on chelicerate phylogenetics (Giribet et al. 2002; Pepato et al. 2010; Arabi et al. 2012). Contrary to this, one can find more DNA sequences for other small arachnid orders in GenBank: 105 for Uropygi, 200 for Schizomida, 200 for Ricinulei, 251 for Amblypygi, and 502 for Pseudoscorpiones, [checked on October 25th, 2013]. In addition, there are only 2 sequences available in the Barcode of Life website (http://www.barcodinglife.org). Palpigrades are delicate animals that walk sensing the substrate with what seems a nervous behaviour of the first pair of walking legs, and use their unmodified palps for walking, unlike all other arachnids (Fig. 1). While moving, most palpigrades keep the flagellum upward, moving it laterally. Accordingly, it is possible that the uplifted flagellum is associated with perception of the environment (Ferreira and Souza 2012). These small, depigmented and highly translucent arachnids range in size from 0.65 mm in Eukoenenia grassii (Hansen, 1901) to 2.4 mm in the "giant" E. draco (Peyerimhoff, 1906) from caves on the island of Majorca (Mayoral and Barranco 2013). Eukoenenia spelaea (Peyerimhoff, 1902) from Slovakia has recently been reported to feed on heterotrophic Cyanobacteria (Smrž et al. 2013). The mode of sperm transfer in these arachnids remains unknown. The living members of the order are currently divided in two families, Eukoeneniidae Petrunkevitch, 1955, with 4 genera and 85 named species, and Prokoeneniidae Condé, 1996, with 2 genera and 7 named species (Harvey 2002; Prendini 2011; Souza and Ferreira 2013). Eukoeneniidae includes the genera Allokoenenia Silvestri, 1913 (1 sp. from West Africa), Eukoenenia Börner, 1901 (71 spp., on all continents under tropical and subtropical climate; in temperate regions predominantly in caves), Koeneniodes Silvestri, 1913 (8 Palaeotropical spp.) and Leptokoenenia Condé, 1965 (5 spp. in the Afrotropical, Neotropical and Palearctic regions).

72 Prokoeneniidae includes the genera *Prokoenenia* Börner, 1901 (6 spp. in the Nearctic,

73 Neotropical and Oriental regions) and *Triadokoenenia* Condé, 1991 (1 sp. from Madagascar).

74 Further unnamed new species are known to us from various parts of the world.

The position of Palpigradi among the arachnid orders remains highly debated. The largest set of data analysed to date places them as the sister group to Acariformes mites in a basal position within arachnids, although without support (Regier *et al.* 2010). The most recent morphological cladistic analysis of arachnid relationships leaves them mostly unresolved among the clades Stomothecata, Haplocnemata, Pantetrapulmonata, and Acaromorpha (Shultz 2007). Earlier studies combining morphology and a small set of molecular data placed Palpigradi as the sister group of Ricinulei + Tetrapulmonata or as sister to Pycnogonida when fossils were considered, although again, without significant clade support (Giribet *et al.* 2002); as sister to a clade including Acari and Solifugae, based on the same two markers used in earlier studies (Pepato *et al.* 2010); or in an unresolved position within arachnids (Arabi *et al.* 2012). Even less is known about the internal relationships of the group, since no published study—molecular or morphological—has yet incorporated information for more than one palpigrade species, and only one unpublished masters thesis has explored palpigrade relationships cladistically, using morphology (Montaño Moreno 2008).

To bridge this important gap in the knowledge of this arachnid order, although acknowledging the difficulties in sampling and identification of these elusive animals, we obtained samples for as many species of palpigrades as possible and from as many localities as possible with the aim to obtain molecular DNA sequence data to generate a first hypothesis of internal palpigrade relationships.

Materials and Methods

Taxon sampling

Palpigrades are difficult to obtain and identify, and success of field sampling differed among regions included in the study. In Western Australia, many samples were collected indirectly in caves and bore holes. In Brazil and Europe, they can be abundant in caves, where fresh specimens have recently become available for inclusion in molecular studies. Additional samples

were from soil samples in Australia, Italy and the USA. In addition to fresh material collected for this study, older specimens were used, especially from the diverse cave systems in Brazil, where several new species have been recently described (Souza and Ferreira 2010; Ferreira et al. 2011; Souza and Ferreira 2011a; Souza and Ferreira 2011b; Souza and Ferreira 2012a; Souza and Ferreira 2012b). While a recently collected specimen of Eukoenenia ferratilis Souza & Ferreira, 2011 amplified well for some of the studied markers, none of the six specimens of Allokoenenia spp. and the two specimens of Leptokoenenia sp. collected from the caves yielded workable DNA. We also obtained a relatively large collection of specimens from the Western Australian bore holes from Barrow Island and the Pilbara, but these were collected from litter traps and many specimens did not amplify or only yielded some amplicons. Some of these specimens are probably related to the Western Australian endemic E. guzikae Barranco & Harvey, 2008, but unrelated to the more widespread species E. mirabilis (Grassi & Calandruccio, 1885), also found in Western Australia (Harvey et al. 2006; Barranco and Harvey 2008). A single specimen of Prokoenenia wheeleri was obtained from the Austin area (Texas, USA), but amplified well for all fragments attempted. In addition, we obtained samples of Eukoenenia mirabilis from Italy (Christian et al. 2010) and Australia (Harvey et al. 2006), E. spelaea (Peyerimhoff, 1902) from multiple localities in Slovenia and Slovakia (Kováč et al. 2002; Zagmajster and Kováč 2006; Král et al. 2008). Italian samples also include E. bonadonai Condé, 1979 and E. strinatii Condé, 1977, collected in caves. We also included specimens from multiple localities from the hansenichilanga group of Eukoenenia from Mexico and the USA (Montaño-Moreno 2012). Additional specimens come from Mexican caves and South Africa. Details on collecting localities are available in Table 1 and in MCZBASE (http://mczbase.mcz.harvard.edu/SpecimenSearch.cfm). Vouchers or additional specimens are deposited in the Museum of Comparative Zoology, Harvard University (MCZ), and in the Western Australian Museum (WAM).

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

We included three species available in GenBank, one from South Africa sequenced by Giribet *et al.* (2002), one from Brazil from Pepato *et al.* (2010), and one of unknown origin published by Arabi *et al.* (2012). Here we added sequences from an additional South African specimen from the same collection of that from Giribet *et al.* (2002), and a specimen of *E. ferratilis* from Brazil, which was identical to the specimen reported by Pepato *et al.* (2010) as *Eukoenenia* sp., and to which we refer to as *E. cf. ferratilis* in the present study. Outgroup taxa were selected from GenBank (Table 2), mostly from previous studies on arthropod or arachnid phylogeny using nuclear ribosomal genes (Giribet *et al.* 2002; Mallatt and Giribet 2006).

Molecular methods

Although we attempted to amplify and sequence five molecular markers typically used in other analyses of arachnid systematics (e.g., Dimitrov *et al.* 2012; Giribet *et al.* 2012), the mitochondrial 16S rRNA gene only amplified for *Prokoenenia wheeleri* and the nuclear proteinencoding gene histone H3, although amplified for several samples, did not produce clean reads. We thus restricted our study to the two broadly available nuclear ribosomal genes, the complete 18S rRNA and ca. 2.2 Kb of 28S rRNA, and the mitochondrial protein-encoding cytochrome *c* oxidase subunit I (COI hereafter) (as in Murienne *et al.* 2008), although the latter gene only amplified for about a third of the specimens (Table 1). For two of the bore-hole Western Australian specimens, poorly preserved, only the middle amplicon of 28S rRNA worked.

Total DNA was extracted from whole specimens or from the opisthosomal region using Qiagen's DNEasy® tissue kit (Valencia, CA, USA). Although we were aiming to preserve the digested carcass as a morphological voucher, it was completely digested and not recoverable. Purified genomic DNA was used as a template for Polymerase chain reactions (PCR) amplification. PCR, visualization by agarose gel electrophoresis, and direct sequencing were conducted for most specimens as described in earlier work, e.g., Edgecombe and Giribet (2009). Chromatograms obtained from the automatic sequencer were read and sequences assembled using the sequence editing software Sequencher™ (Gene Codes Corporation, Ann Arbor, MI, USA). Sequence data were edited in MacGDE (Linton 2005). The three genes were analysed as follows:

18S rRNA: This marker was amplified in three amplicons (*a*, *b*, *c*), as in previous studies (Edgecombe and Giribet 2009; Giribet *et al*. 2010; Giribet *et al*. 2012). In the present study we include 27 palpigrade specimens plus 8 outgroups, for a total of 1760-1771 bp per complete sequence (up to 1805 bp for one of the outgroups). From the 27 palpigrade sequences all but three were complete; *E. spelaea* is missing fragment *a* and the sample of *Eukoenenia* from South Africa (DNA100456.2) is missing fragment *b*. For the direct optimization analyses the three amplicons were treated as a single input file, containing 23 sequences, and divided into six fragments. The three amplicons were concatenated for the static alignment analyses.

COI: This widely used mitochondrial marker amplified for ten palpigrade terminals in a single amplicon using primers LCO—HCO, showing no length variation (654 bp analysed), plus one was available in GenBank. COI did not amplify for many individuals, perhaps due to major changes in this marker, as evidenced by the deletion of one amino acid with respect to the outgroups. Five outgroup sequences were obtained from GenBank, but these were 3 bp longer in all cases except for the pseudoscorpion. It was analysed as a single fragment; not pre-aligned due to the length difference with some outgroups.

Phylogenetic analyses

Parsimony analyses were based on a direct optimization (DO) approach (Wheeler 1996) using POY v. 5.0 (Varón *et al.* 2012). Tree searches were performed using the timed search function in POY, i.e., multiple cycles of (a) building Wagner trees, (b) subtree pruning and regrafting (SPR), and (c) tree bisection and reconnection (TBR), (d) ratcheting (Nixon 1999), and (e) tree-fusing (Goloboff 1999, 2002) [command: search (max_time:00:01:00, min_time:00:00:10, hits:20, memory:gb:2)]. For the individual partitions, timed searches of 1 hour were run on 4 processors under six parameter sets, as in Giribet *et al.* (2012) (see Table 3). For the combined analysis of the three markers we started with the same search strategy, giving the 28S rRNA trees as input—as these contained all the taxa in the combined data set—, and the resulting trees were given as input for a second round of analyses (sensitivity analysis tree fusing; SATF),

as described by Giribet (2007), and continued until the tree lengths stabilised (Giribet *et al.* 2012). The optimal parameter set was estimated using the modified wILD metrics (Wheeler 1995; Sharma *et al.* 2011), as a proxy for the parameter set that minimizes overall incongruence among data partitions (Table 4). Nodal support for the optimal parameter set was estimated via jackknifing (250 replicates) with a probability of deletion of e⁻¹ (Farris *et al.* 1996) using auto sequence partition, as discussed in earlier work (Giribet *et al.* 2012).

Maximum likelihood (ML) analyses were conducted on static multiple sequence alignments (MSA) inferred in MUSCLE v. 3.6 (Edgar 2004) through the EMBL-EBI server (http://www.ebi.ac.uk/Tools/msa/muscle/). We also used an implied alignment (IA) generated in POY (Wheeler 2003; Giribet 2005) for subsequent analyses based on static alignments, as recently explored by Giribet and Edgecombe (2013b) for a centipede data set. The MUSCLE alignments were conducted for each gene independently. The IA and MSA therefore were based on the same data (see length for each gene in Table 5). In order to evaluate the impact of the hypervariable regions in the data set, MSAs and IAs were subsequently trimmed with Gblocks v. 0.91b (Castresana 2000; Talavera and Castresana 2007) to cull positions of ambiguous homology (see length for each trimmed gene in Table 5). In the case of 28S, fragments a and bc were Gblocked separately, due to the larger proportion of missing data in the a fragment, which otherwise would be deleted from the final 28S alignment. These data sets are thus based on different data from their original sources and from each other, but the remaining data use the same homology scheme as the source. Data sets were concatenated with SequenceMatrix (Vaidya et al. 2011).

Maximum likelihood analyses were conducted using RAxML ver. 7.2.7 (Stamatakis et~al. 2008b) in the CIPRES server (Miller et~al. 2010). For the searches, a unique General Time Reversible (GTR) model of sequence evolution with corrections for a discrete gamma distribution (GTR + Γ) was specified for each data partition, and 100 independent searches were conducted. Nodal support was estimated via the rapid bootstrap algorithm (1000 replicates) using the GTR-CAT model (Stamatakis et~al. 2008a). Bootstrap resampling frequencies were thereafter mapped onto the optimal tree from the independent searches.

In total we analysed five data sets accounting for different optimality criteria, homology schemes, and/or amount of data, as follows:

- Analysis 1. Direct optimization/dynamic homology under parsimony (full sensitivity analysis of 6 parameter sets) analysed in POY
 - Analysis 2. Static homology from the implied alignment for the optimal parameter set under ML (analysed in RAxML)
 - Analysis 3. Static homology from the implied alignment for the optimal parameter set trimmed with Gblocks under ML (analysed in RAxML)
 - Analysis 4. Static homology based on MUSCLE multiple sequence alignment (analysed in RAxML)
 - Analysis 5. Static homology based on MUSCLE/Gblocks (analysed in RAxML)

231

232

233

234

235

236

237

238

239

240

241

242

243

244

245

246

247

248

249

250

251

222

223

224

225

226

227

228

229

230

Results and Discussion

All phylogenetic analyses yielded very similar results with respect to the ingroup relationships, while the outgroup relationships were incongruent from analysis to analysis and unsupported for the most part (Figs. 2 and 3). The latter was expected given the small amount of data and outgroup taxa and the poor resolution in deep arachnid relationships in other studies (e.g., Wheeler and Hayashi 1998; Giribet et al. 2002; Pepato et al. 2010; Regier et al. 2010). The optimal parameter set under parsimony direct optimization was 3211 (where indel opening costs 3, indel extension 1, transversions cost 2 and transitions cost 1; wILD = 0.00913), with a cost of 10,408 weighted steps (Fig. 2). Nearly all examined parameter sets concurred on the topology of the optimal parameter set, with the exception of Eukoenenia spelaea IZ-19346 from Slovenia, and the resolution of one of the Eukoenenia clades (see below). Likewise, the analyses of the four data sets analysed under maximum likelihood were nearly identical, except for some of the shallowest relationships. One of these trees, the one for the multiple sequence alignment trimmed with Gblocks—the one that could be potentially the most different from the POY analysis—is presented in Fig. 3, and it is virtually identical to the direct optimization tree. From the 10 nodes depicted in Fig. 2 summarizing the six direct optimization and the four maximum likelihood analyses, 5 were recovered in all analyses. Support values for these five nodes is high for most analyses (jackknife values are lower by definition), with the exception of clades III and IV in the DO analysis. Basically, nearly all analyses concur on the overall topology of the palpigrade tree.

All analyses show a basal dichotomy between *Prokoenenia wheeleri* (the only Prokoeneniidae represented in our analyses) and the remaining samples, which we consider as Eukoenenia for further discussion—even if some samples from GenBank or from the Australian boreholes were not identified. Eukoenenia is divided into four main clades, indicated in Figures 2 and 3. Clade I includes E. florenciae from Slovakia, Brazil, and unidentified specimens probably belonging to the same species from the USA and Mexico, and another species from a cave in Guerrero, Mexico (IZ-128499). Clade II includes E. spelaea and E. s. hauseri Condé, 1974 from Slovenia and Slovakia, and several additional samples from Slovenia and Italy, including E. strinatii, E. bonadonai and E. austriaca (Hansen, 1926); E. spelaea IZ-19346 from Slovenia clusters with these species in some analyses, but not all (Fig. 2). Clade III includes E. ferratilis from Brazil, the specimens from the Australian bore holes, and an undescribed species from Brazil (IZ-19345). Clade IV includes E. mirabilis from Australia and Italy, and unidentified specimens from South Africa, plus a specimen from a cave in Chiapas, Mexico (IZ-136274) and a GenBank specimen (JA-2011) of unknown origin. Clades I and II are supported in all analyses; Clade III is supported in all analyses except for the DO analysis under parameter set 211; Clade IV is unsupported in the ML analysis of the trimmed MSA. Eukoenenia spelaea IZ-19346 appears as the sister group to Clade II under 4 analytical parameter sets in DO and in the untrimmed ML analyses, both for the IA and for the MSA. The E. florenciae clade (Clade I) always forms the sister group of the E. spelaea clade (Clade II), although E. spelaea IZ-19346 sometimes forms the sister group of the E. florenciae clade. While the E. ferratilis clade (Clade III) often forms the sister group to the E. mirabilis clade (Clade IV) (Figs. 2, 3), and is well supported in the probabilistic analyses (97 to 100% bootstrap support, depending on the analysis), under some parameter sets Clade III is sister to the E. spelaea—E. florenciae clade (parameter sets 111, 211, 221, 3221).

252

253

254

255

256

257

258

259

260

261

262

263

264

265

266

267

268

269

270

271

272

273

274

275

276

277

278

279

280

281

282

283

alternative methods (parsimony and maximum likelihood) based on identical raw data with different homology schemes (implied alignments versus multiple sequence alignments), or different data sets (trimmed implied alignments and trimmed multiple sequence alignments). There are very few cases with such consistency across weighting schemes, homology schemes, and methodologies, but a recent case was documented for scutigeromorph centipedes (Giribet and Edgecombe 2013b). In that case, the fossil record and denser sampling allowed for accurate molecular dating and analyses of diversification of lineages through time, and it was suggested

that the congruence across analyses was due to constant rates of diversification through more than 400 million years of evolution in the group. We can only guess this for palpigrades, as the fossil record for this group is rare, and a single Pliocene specimen is known (Rowland and Sissom 1980; Delclòs *et al.* 2008; Dunlop 2010), although the group must be much older in origin (see for example Giribet and Edgecombe 2013a).

Phylogenetic analysis of the three molecular markers combined and for all analyses performed resolves into Prokoeneniidae (although represented by a single species) and Eukoeneniidae, supporting the monophyly of Eukoeneniidae—palpigrades without sternal opisthosomal vesicles (Condé 1996). We were, however, unable to obtain samples of *Triadokoenenia* or of additional *Prokoenenia* species, thus not being able to test the taxon Prokoeneniidae. Within Eukoeneniidae, the four main clades discussed above are supported in nearly all analyses. But species identifications in palpigrades do not seem straightforward. Within Clade I, the specimens of *Eukoenenia* from Texas (USA), the Mexican state of Yucatán, *E. cf. florenciae* from Brazil and *E. florenciae* from Slovakia show nearly identical COI sequences and identical nuclear ribosomal RNA sequences, suggesting that they may be conspecific (see Edgecombe and Giribet 2008; Vélez *et al.* 2012). In contrast, Clade II includes three lineages of the morphospecies *E. spelaea*. From these, two samples identified as *E. spelaea* and *E. spelaea hauseri* from Slovenia appear identical for the nuclear ribosomal genes (but did not amplify for COI).

Clade III includes the Western Australian samples and *Eukoenenia ferratilis* from the Iron caves of Minas Gerais (Brazil). Difficulties in amplifying the Australian samples and the lack of COI information for any of the members of the clade precludes us from understanding genetic variability within this clade of geographically distant species (both between the continents, but also among the Western Australian localities), although most analyses consistently resolve this clade of six individuals with reciprocal monophyly of the two geographic regions.

Clade IV, although with less support than the other three clades, includes the sample of unknown provenance sequenced by Arabi *et al.* (2012), a specimen from caves in Chiapas, and the cosmopolitan *E. mirabilis*, including two specimens from Italy (identical for all markers) and two putative members of this species from South Africa plus a sample of *E. mirabilis* from Australia. While *E. mirabilis* has been suggested to be a synanthropic species originating in the Mediterranean region with recent introductions to South Africa, Australia, Chile and

Madagascar (Harvey *et al.* 2006), our limited data suggest a close relationship between one of the South African samples and the Australian specimen, even in the absence of COI data, and therefore suggesting changes in the nuclear ribosomal genes with respect to the Italian sample. Further study of Gondwanan *E. mirabilis* and addition of circum-Mediterranean samples should be undertaken to bring this matter to conclusion.

Given the sampling of this study it is still early to make any firm conclusions about palpigrade relationships. We were not able to test for the monophyly of Prokoeneniidae, and monophyly of *Eukoenenia* is not thoroughly tested either. Attempts to sequence *Allokoenenia* and *Leptokoenenia* were unsuccessful, and we were unable to obtain specimens of the Palaeotropical *Koeneniodes* and *Triadokoenenia*. Few studies have looked at variation among palpigrade species, but Král *et al.* (2008) investigated the karyotypes of *E. spelaea* from Slovakia and *E. mirabilis*, which appear in different clades in our study (Clades II and IV, respectively). However, the karyotypes of both species showed no variation, both consisting of a low number of tiny chromosomes that decrease gradually in size and a lack of morphologically differentiated sex chromosomes, suggesting that molecular data may be more informative than karyotypic data for separating species.

Morphologically, the characters used to differentiate *Eukoenenia* species are mostly restricted to the number of lobules in the lateral organs or the number of setae in different body regions, but the significance of these characters has not been tested phylogenetically—for example, *E. mirabilis* and *E. ferratilis* are very similar morphologically with many somatic traits, considered important for taxonomy, virtually identical (Souza and Ferreira 2011a). However, these two species belong to different clades, reflecting that their differences in genital morphology and chaetotaxy may be better systematic characters than the ones outlined above. Our study thus provides a new framework for adding new sequences and testing the significance of these characters. Additional samples and especially more genera must however be added before we can attempt a taxonomic revision of the higher taxa in Palpigradi.

Conclusions

Palpigrades are a poorly understood group of tiny soil arthropods, often found exclusively in caves, and have received little attention from a phylogenetic point of view. Here we were able

to amass specimens from different environments (caves and soil) from Australia, Africa, Europe, South America and North America with the aim of generating a molecular phylogenetic hypothesis for the group. The difficulty in obtaining well-preserved material for molecular work is reflected in the large number of specimens that did not yield DNA of enough quality for sequencing, but we were able to propose the first phylogenetic hypothesis of the group based on molecular data to find monophyly of Eukoeneniidae and its division into four main clades, three of these including samples from multiple continents. Given the absence of denser sampling and proper clock calibrations, our data cannot discern whether palpigrades are a very old group that diversified prior to the breakup of Pangaea, or a group of animals that disperses across large geographic distances, as suggested by some widespread species. Long-range dispersal is however difficult to reconcile with the narrow ecological conditions and the facility with which these animals desiccate once removed from their environments.

Acknowledgements

Julián Bueno-Villegas and Jesús A. Cruz-López helped with fieldwork in Yucatán; T. Delić and S. Prevorčnik in Slovenia; P. Ľuptáčik and A. Mock in Slovakia; L. Galli and M. Zinni provided samples from Italy; J. Van der Schyff from South Africa; J. Christophoryová from Slovakia; staff from the environmental companies Biota Environmental Sciences and Subterranean Ecology provided samples from Western Australia. Gustavo Hormiga and Nikolaj Scharff kindly provided comments that helped improve this article. This work has been supported by internal funds from the Museum of Comparative Zoology and by NSF grant #1144417 to G. G. and G. Hormiga (Collaborative Research: ARTS: Taxonomy and systematics of selected Neotropical clades of arachnids).

368	References
369 370	Arabi, J., Judson, M. L., Deharveng, L., Lourenço, W. R., Cruaud, C., and Hassanin, A. (2012).
371	Nucleotide composition of CO1 sequences in Chelicerata (Arthropoda): detecting new
372	mitogenomic rearrangements. <i>Journal of Molecular Evolution</i> 74 , 81-95.
373	doi:10.1007/s00239-012-9490-7
374	Barranco, P., and Harvey, M. S. (2008). The first indigenous palpigrade from Australia: a new
375	species of <i>Eukoenenia</i> (Palpigradi : Eukoeneniidae). <i>Invertebrate Systematics</i> 22 , 227-
376	233.
377	Beccaloni, J. (2009). 'Arachnids.' (The Natural History Museum: London.)
378	Castresana, J. (2000). Selection of conserved blocks from multiple alignments for their use in
379	phylogenetic analysis. Molecular Biology and Evolution 17, 540-552.
380	Christian, E., Capurro, M., and Galli, L. (2010). Phenology of two syntopic <i>Eukoenenia</i> species in a
381	northern Italian forest soil (Arachnida: Palpigradi). Revue suisse de Zoologie 117, 829-
382	834.
383	Condé, B. (1996). Les palpigrades, 1885-1995: acquisitions et lacunes. Revue suisse de Zoologie
384	hors série, 87-106.
385	Delclòs, X., Nei, A., Azar, D., Bechly, G., Dunlop, J. A., Engel, M. S., and Heads, S. W. (2008). The
386	enigmatic Mesozoic insect taxon Chresmodidae (Polyneoptera): New palaeobiological
387	and phylogenetic data, with the description of a new species from the Lower Cretaceous
388	of Brazil. Neues Jahrbuch für Geologie und Palaontologie-Abhandlungen 247 , 353-381.
389	Dimitrov, D., Lopardo, L., Giribet, G., Arnedo, M. A., Álvarez-Padilla, F., and Hormiga, G. (2012).
390	Tangled in a sparse spider web: single origin of orb weavers and their spinning work
391	unravelled by denser taxonomic sampling. Proceedings of the Royal Society B: Biological
392	Sciences 279 , 1341-1350. doi:10.1098/rspb.2011.2011
393	Dunlop, J. A. (2010). Geological history and phylogeny of Chelicerata. Arthropod Structure &
394	Development 39, 124-142. doi:10.1016/j.asd.2010.01.003

395	Edgar, R. C. (2004). MUSCLE: multiple sequence alignment with high accuracy and high
396	throughput. Nucleic Acids Research 32, 1792-1797. doi:10.1093/nar/gkh340
397	Edgecombe, G. D., and Giribet, G. (2008). A New Zealand species of the trans-Tasman centipede
398	order Craterostigmomorpha (Arthropoda : Chilopoda) corroborated by molecular
399	evidence. Invertebrate Systematics 22, 1-15. doi:10.1071/Is07036
100	Edgecombe, G. D., and Giribet, G. (2009). Phylogenetics of scutigeromorph centipedes
101	(Myriapoda: Chilopoda) with implications for species delimitation and historical
102	biogeography of the Australian and New Caledonian faunas. Cladistics 25, 406-427.
103	doi:10.1111/j.1096-0031.2009.00253.x
104	Farris, J. S., Albert, V. A., Källersjö, M., Lipscomb, D., and Kluge, A. G. (1996). Parsimony
105	jackknifing outperforms neighbor-joining. Cladistics 12, 99-124.
106	Ferreira, R. L., and Souza, M. F. V. R. (2012). Notes on the behavior of the advanced troglobite
107	Eukoenenia maquinensis Souza & Ferreira 2010 (Palpigradi: Eukoeneniidae) and its
108	conservation status. Speleobiology Notes 4, 17-23.
109	Ferreira, R. L., Souza, M. F. V. R., Machado, E. O., and Brescovit, A. D. (2011). Description of a
110	new Eukoenenia (Palpigradi: Eukoeneniidae) and Metagonia (Araneae: Pholcidae) from
111	Brazilian caves, with notes on their ecological interactions. The Journal of Arachnology
112	39 , 409-419. doi:10.1636/Ha11-03.1
113	Giribet, G. (2005). Generating implied alignments under direct optimization using POY. <i>Cladistics</i>
114	21 , 396-402.
115	Giribet, G. (2007). Efficient tree searches with available algorithms. <i>Evolutionary Bioinformatics</i>
116	Online 3 , 341-356.
117	Giribet, G., and Edgecombe, G. D. (2013a). The Arthropoda: a phylogenetic framework. In
118	'Arthropod Biology and Evolution – Molecules, Development, Morphology'. (Eds A.
119	Minelli, G. Boxshall and G. Fusco) pp. 17-40. (Springer: Berlin.)
120	Giribet, G., and Edgecombe, G. D. (2013b). Stable phylogenetic patterns in scutigeromorph
121	centipedes (Myriapoda: Chilopoda: Scutigeromorpha): dating the diversification of an

423	doi:10.1071/IS13019
424 425 426	Giribet, G., Edgecombe, G. D., Wheeler, W. C., and Babbitt, C. (2002). Phylogeny and systematic position of Opiliones: a combined analysis of chelicerate relationships using morphological and molecular data. <i>Cladistics</i> 18 , 5-70.
120	morphological and morecular data. Charistics 10, 5 70.
427	Giribet, G., Sharma, P. P., Benavides, L. R., Boyer, S. L., Clouse, R. M., de Bivort, B. L., Dimitrov, D.
428	Kawauchi, G. Y., Murienne, J. Y., and Schwendinger, P. J. (2012). Evolutionary and
429	biogeographical history of an ancient and global group of arachnids (Arachnida:
430	Opiliones: Cyphophthalmi) with a new taxonomic arrangement. Biological Journal of the
431	Linnean Society 105 , 92-130. doi:10.1111/J.1095-8312.2011.01774.X
432	Giribet, G., and Shear, W. A. (2010). The genus Siro Latreille, 1796 (Opiliones, Cyphophthalmi,
433	Sironidae), in North America with a phylogenetic analysis based on molecular data and
434	the description of four new species. Bulletin of the Museum of Comparative Zoology 160
435	1-33.
436	Giribet, G., Vogt, L., Pérez González, A., Sharma, P., and Kury, A. B. (2010). A multilocus approach
437	to harvestman (Arachnida: Opiliones) phylogeny with emphasis on biogeography and
438	the systematics of Laniatores. Cladistics 26, 408-437. doi:10.1111/j.1096-
439	0031.2009.00296.x
440	Goloboff, P. A. (1999). Analyzing large data sets in reasonable times: solutions for composite
441	optima. <i>Cladistics</i> 15 , 415-428.
442	Goloboff, P. A. (2002). Techniques for analyzing large data sets. In 'Techniques in Molecular
443	Systematics and Evolution'. (Eds R. DeSalle, G. Giribet and W. Wheeler) pp. 70-79.
444	(Brikhäuser Verlag: Basel.)
445	Grassi, B., and Calandruccio, S. (1885). Intorno ad un nuovo aracnide artrogastro (Koenenia
446	mirabilis) che crediamo rappresentante d'un nuovo ordine (Microteliphonida).
447	Naturalista Siciliano 4 , 127-133, 162-169.
448	Harvey, M. S. (2002). The neglected cousins: What do we know about the smaller arachnid
449	orders? The Journal of Arachnology 30 , 357-372.

450	Harvey, M. S., Staniavsky, F., and Theron, P. D. (2006). The distribution of Eukoenenia mirabilis
451	(Palpigradi: Eukoeneniidae): a widespread tramp. Records of the Western Australian
452	Museum 23 , 199-203.
453	Kováč, L., Mock, A., Ľuptáčik, P., and Palacios-Vargas, J. G. (2002). Distribution of <i>Eukoenenia</i>
454	spelaea (Peyerimhoff, 1902) (Arachnida, Palpigradida) in the Western Carpathians with
455	remarks on its biology and behaviour. In 'Studies on Soil Fauna in Central Europe'. (Eds K
456	Tajovský, V. Balík and V. Pižl) pp. 93-99: České Budějovice.)
457	Král, J., Kováč, L., Šťahlavský, F., Lonský, P., and L'uptácik, P. (2008). The first karyotype study in
458	palpigrades, a primitive order of arachnids (Arachnida: Palpigradi). <i>Genetica</i> 134 , 79-87.
459	Linton, E. W. (2005). MacGDE: Genetic Data Environment for MacOS X. (Software available at
460	http://www.msu.edu/~lintone/macgde/)
461	Mallatt, J., and Giribet, G. (2006). Further use of nearly complete 28S and 18S rRNA genes to
462	classify Ecdysozoa: 37 more arthropods and a kinorhynch. Molecular Phylogenetics and
463	Evolution 40, 772-794. doi:10.1016/J.Ympev.2006.04.021
464	Mayoral, J. G., and Barranco, P. (2013). Rediscovery of the troglobious palpigrade Eukoenenia
465	draco (Peyerimhoff 1906) (Palpigradi: Eukoeneniidae), with notes on the adaptations to
466	a cave-dwelling life. Zootaxa 3635 , 174-184. doi:10.11646/zootaxa.3635.2.5
467	Miller, M. A., Pfeiffer, W., and Schwartz, T. (2010). Creating the CIPRES Science Gateway for
468	Inference of Large Phylogenetic Trees. In 'Proceedings of the Gateway Computing
469	Environments Workshop (GCE)'. New Orleans pp. 1-8
470	Montaño Moreno, H. (2008). Revisión taxonómica de los palpígrados (Arachnida: Palpigradi) de
471	México. Masters thesis, Universidad Nacional Autónoma de México.
472	Montaño-Moreno, H. (2012). Redescripción de <i>Eukoenenia hanseni</i> (Arachnida: Palpigradi) y
473	descripción de una nueva especie de palpígrado de México. Revista Ibérica de
474	Aracnología 20 , 1-15.
475	Murienne, J., Harvey, M. S., and Giribet, G. (2008). First molecular phylogeny of the major clades
476	of Pseudoscorpiones (Arthropoda: Chelicerata). Molecular Phylogenetics and Evolution
177	49 170-184 doi:10.1016/j.vmney.2008.06.002

478	Nixon, K. C. (1999). The Parsimony Ratchet, a new method for rapid parsimony analysis.
479	Cladistics 15 , 407-414.
480	Pepato, A. R., da Rocha, C. E., and Dunlop, J. A. (2010). Phylogenetic position of the acariform
481	mites: sensitivity to homology assessment under total evidence. BMC Evolutionary
482	Biology 10 , 235. doi:10.1186/1471-2148-10-235
483	Prendini, L. (2011). Order Palpigradi Thorell, 1888 (In: Animal biodiversity: An outline of higher-
484	level classification and survey of taxonomic richness). Zootaxa 3148, 121.
485	Regier, J. C., Shultz, J. W., Zwick, A., Hussey, A., Ball, B., Wetzer, R., Martin, J. W., and
486	Cunningham, C. W. (2010). Arthropod relationships revealed by phylogenomic analysis
487	of nuclear protein-coding sequences. <i>Nature</i> 463 , 1079-1083. doi:10.1038/nature08742
488	Rowland, J. M., and Sissom, W. (1980). Report on a fossil palpigrade from the Tertiary of Arizona,
489	and a review of the morphology and systematics of the order (Arachnida, Palpigradida).
490	The Journal of Arachnology 8 , 69-86.
491	Sharma, P. P., Vahtera, V., Kawauchi, G. Y., and Giribet, G. (2011). Running wILD: The case for
492	exploring mixed parameter sets in sensitivity analysis. Cladistics 27, 538-549.
493	doi:10.1111/j.1096-0031.2010.00345.x
494	Shultz, J. W. (2007). A phylogenetic analysis of the arachnid orders based on morphological
495	characters. Zoological Journal of the Linnean Society 150, 221-265.
496	Smrž, J., Kováč, Ĺ., Mikeš, J., and Lukešová, A. (2013). Microwhip scorpions (Palpigradi) feed on
497	heterotrophic Cyanobacteria in Slovak caves – A curiosity among Arachnida. PLoS ONE 8,
498	e75989. doi:10.1371/journal.pone.0075989
499	Souza, M. F. V. R., and Ferreira, R. L. (2010). Eukoenenia (Palpigradi: Eukoeneniidae) in Brazilian
500	caves with the first troglobiotic palpigrade from South America. The Journal of
501	Arachnology 38 , 415-424.
502	Souza, M. F. V. R., and Ferreira, R. L. (2011a). A new species of <i>Eukoenenia</i> (Palpigradi:
503	Eukoeneniidae) from Brazilian iron caves. Zootaxa 2886, 31-38.

505	Eukoeneniidae) from Brazil. <i>Journal of Arachnology</i> 39 , 185-188. doi:10.1636/Ha10-43.1
506 507	Souza, M. F. V. R., and Ferreira, R. L. (2012a). <i>Eukoenenia virgemdalapa</i> (Palpigradi: Eukoeneniidae): a new troglobitic palpigrade from Brazil. <i>Zootaxa</i> 3295 , 59-64.
508	Souza, M. F. V. R., and Ferreira, R. L. (2012b). A new highly troglomorphic species of <i>Eukoenenia</i>
509	(Palpigradi: Eukoeneniidae) from tropical Brazil. <i>The Journal of Arachnology</i> 40 , 151-158.
510	Souza, M. F. V. R., and Ferreira, R. L. (2013). Two new species of the enigmatic Leptokoenenia
511	(Eukoeneniidae: Palpigradi) from Brazil: First record of the genus outside intertidal
512	environments. PLoS ONE 8, e77840. doi:10.1371/journal.pone.0077840
513	Stamatakis, A., Hoover, P., and Rougemont, J. (2008a). A rapid bootstrap algorithm for the
514	RAxML Web servers. Systematic Biology 57 , 758-771. doi:10.1080/10635150802429642
515	Stamatakis, A. P., Meier, H., and Ludwig, T. (2008b). RAxML: A parallel program for phylogenetic
516	tree inference.
517	Talavera, G., and Castresana, J. (2007). Improvement of phylogenies after removing divergent
518	and ambiguously aligned blocks from protein sequence alignments. Systematic Biology
519	56 , 564-577. doi:10.1080/10635150701472164
520	Vaidya, G., Lohman, D. J., and Meier, R. (2011). SequenceMatrix: concatenation software for the
521	fast assembly of multi-gene datasets with character set and codon information.
522	Cladistics 27, 171-180. doi:10.1111/j.1096-0031.2010.00329.x
523	Varón, A., Lucaroni, N., Hong, L., and Wheeler, W. C. (2012). POY 5.0.0. (American Museum of
524	Natural History. http://research.amnh.org/scicomp: New York)
525	Vélez, S., Mesibov, R., and Giribet, G. (2012). Biogeography in a continental island: population
526	structure of the relict endemic centipede Craterostigmus tasmanianus (Chilopoda,
527	Craterostigmomorpha) in Tasmania using 16S rRNA and COI. Journal of Heredity 103, 80-
528	91. doi:10.1093/jhered/esr110
529	Wheeler, W. C. (1995). Sequence alignment, parameter sensitivity, and the phylogenetic analysis
530	of molecular data. Systematic Biology 44, 321-331.

531 532	Wheeler, W. (1996). Optimization alignment: the end of multiple sequence alignment in phylogenetics? <i>Cladistics</i> 12 , 1-9.
533 534	Wheeler, W. C. (2003). Implied alignment: a synapomorphy-based multiple-sequence alignment method and its use in cladogram search. <i>Cladistics</i> 19 , 261-268.
535 536	Wheeler, W. C., and Hayashi, C. Y. (1998). The phylogeny of the extant chelicerate orders. Cladistics 14, 173-192.
537538539	Whiting, M. F., Carpenter, J. M., Wheeler, Q. D., and Wheeler, W. C. (1997). The Strepsiptera problem: phylogeny of the holometabolous insect orders inferred from 18S and 28S ribosomal DNA sequences and morphology. <i>Systematic Biology</i> 46 , 1-68.
540	Zagmajster, M., and Kováč, Ĺ. (2006). Distribution of palpigrades (Arachnida, Palpigradi) in
541 542	Slovenia with a new record of <i>Eukoenenia austriaca</i> (Hansen, 1926). <i>Natura Sloveniae</i> 8 , 23-31.
543	

544 Fig. 1. Photographs of (A) Eukoenenia spelaea, Ardovská Cave (Slovak Karst, Slovakia), 545 photographed by Ľ. Kováč & V. Kóňa; (B) Prokoenenia wheeleri, Austin (Texas, USA), 546 photographed by L. McCutchen; (C) Eukoenenia mirabilis, flagellum, segments 1 to 10; (D) 547 Eukoenenia bonadonai, male genital lobes; (E) E. bonadonai, female genital lobes; (F) E. 548 bonadonai, mouth cone and chelicerae (C-D photographed by E. Christian). 549 550 Fig. 2. Left: Optimal tree at 10,408 weighted steps obtained from the direct optimization 551 analysis under parameter set 3211 of the combined analysis of the three genes. Numbers on 552 branches indicate jackknife support values. Navajo rugs are shown in selected nodes; Black 553 square indicates monophyly, white square non-monophyly. Specific parameter sets or analyses 554 indicated in the figure. Numerals indicate parameter set under parsimony direct optimization; IA 555 (ML analysis using implied alignment under parameter set 3211); IAg (Idem, Gblocked); MSA 556 (ML analysis of the MUSCLE multiple sequence alignment); MSAg (Idem, Gblocked). Clades I to 557 IV are indicated. 558 559 Fig. 3. Optimal maximum likelihood tree (-LnL = -24955.690470) of the combined data set using

560

561

bootstrap support values.

the MUSCLE multiple sequence alignment trimmed with Gblocks. Numbers on nodes indicate

Table 1. Palpigrade specimens, accession numbers, collecting information and amplified loci with GenBank accession numbers

IZ: Department of Invertebrate Zoology, Museum of Comparative Zoology, Cambridge; DNA: MCZ DNA collection; WAM: Western Australian Museum, Perth; MNHN: Muséum national d'histoire Naturelle, Paris. A dash (-) indicates a missing amplicon. New sequences are KF823823 to KF823883

					18S rRNA			28S rRNA		COI
	MCZ No.		Country	а	b	С	а	b	с	
Prokoenenia wheeleri	IZ-134477	DNA107078	Texas, USA	KF823823	KF823823	KF823823	KF823848	KF823848	KF823848	KF823874
Eukoenenia austriaca	IZ-19349	-	Slovenia	KF823824	KF823824	KF823824	KF823849	KF823849	KF823849	-
Eukoenenia bonadonai	IZ-19340	-	Italy	KF823825	KF823825	KF823825	KF823850	KF823850	KF823850	-
Eukoenenia ferratilis	IZ-127609	-	Brazil	KF823826	KF823826	KF823826	KF823851	KF823851	KF823851	-
Eukoenenia cf. ferratilis	-	GenBank		HM070336	HM070336	HM070336	HM070299	HM070299	HM070299	-
Eukoenenia florenciae	IZ-19351	-	Slovakia	KF823827	KF823827	KF823827	KF823852	KF823852	KF823852	KF823875
Eukoenenia cf. florenciae	IZ-19343	-	Brazil	KF823828	KF823828	KF823828	KF823853	KF823853	KF823853	-
Eukoenenia mirabilis	IZ-127901	-	Italy	KF823829	KF823829	KF823829	KF823854	KF823854	KF823854	KF823876
Eukoenenia mirabilis	IZ-127902	-	Italy	KF823830	KF823830	KF823830	KF823855	KF823855	KF823855	KF823877
Eukoenenia mirabilis	IZ-16117	-	Australia	KF823831	KF823831	KF823831	KF823856	KF823856	KF823856	-
Eukoenenia spelaea	IZ-135126	DNA106786	Slovakia	-	KF823832	KF823832	KF823857	KF823857	KF823857	-
Eukoenenia spelaea	IZ-19346	-	Slovenia	KF823833	KF823833	KF823833	KF823858	KF823858	KF823858	KF823878
Eukoenenia spelaea	IZ-19347	-	Slovenia	KF823834	KF823834	KF823834	KF823859	KF823859	KF823859	-
Eukoenenia spelaea hauseri	IZ-19348	-	Slovenia	KF823835	KF823835	KF823835	KF823860	KF823860	KF823860	-
Eukoenenia strinatii	IZ-19341	-	Italy	KF823836	KF823836	KF823836	KF823861	KF823861	KF823861	-
Eukoenenia sp.	IZ-19350	-	Slovenia	KF823837	KF823837	KF823837	KF823862	KF823862	KF823862	KF823879

Eukoenenia sp.	-	DNA100456.1	South Africa	AF207648	AF207648	AF207648	-	AF207653	-	-
Eukoenenia sp.	-	DNA100456.2	South Africa	KF823838	-	KF823839	-	KF823863	-	-
Eukoenenia sp.	IZ-134549	DNA107079	USA	KF823840	KF823840	KF823840	KF823864	KF823864	KF823864	KF823880
Eukoenenia sp.	IZ-127598.1	-	Mexico	KF823841	KF823841	KF823841	KF823865	KF823865	KF823865	KF823881
Eukoenenia sp.	IZ-127598.2	-	Mexico	KF823842	KF823842	KF823842	KF823866	KF823866	KF823866	KF823882
Eukoenenia sp.	IZ-128499	-	Mexico	KF823843	KF823843	KF823843	KF823867	KF823867	KF823867	KF823883
Eukoenenia sp.	IZ-136274	-	Mexico	KF823844	-	KF823844	KF823868	KF823868	KF823868	-
Eukoenenia sp.	IZ-127636	WAM T81111	Australia		-	-	-	KF823869	-	-
Eukoenenia sp.	IZ-127639	WAM T116012	Australia	KF823845	KF823845	KF823845	-	KF823870	KF823870	-
Eukoenenia sp.	IZ-127640	WAM T111422	Australia	-	-	-	-	KF823871	-	-
Eukoenenia sp.	IZ-127643	-	Australia	KF823846	KF823846	KF823846	-	KF823872	KF823872	-
Eukoenenia sp.n.	IZ-19345	-	Brazil	KF823847	KF823847	KF823847	-	KF823873	KF823873	-
Palpigradi sp.	-	MNHN-JAA76		JN018286.1	JN018286.1	JN018286.1	JN018383.1	JN018383.1	JN018383.1	JN018169.1

Table 2. Outgroup sampling with GenBank accession numbers

		18S rRNA	28S rRNA	COI
Anoplodactylus portus	Pycnogonida	AY859551	AY859550	GQ912859
Limulus polyphemus	Xiphosura	U91490	AF212167	AF216203
Pandinus imperator	Scorpiones	AY210831	AY210830	AY156582
Metasiro americanus	Opiliones	DQ825542	DQ825595	DQ825645
Calocheiridius termitophilus	Pseudoscorpiones	AY859559	AY859558	EU559544
Dermacentor sp.	Acari	Z74480	AY859582	-
Eremobates sp.	Solifugae	AY859573	AY859572	-
Mastigoproctus giganteus	Uropygi	AF005446	AY859587	JN018215

Table 3. Result of the POY timed searches (search) and improvement after each round of SATF for the six explored parameter sets

	1	SATF2	SATF3
111	6520	6520	6520
121	10076	10076	10076
211	7543	7543	7543
221	11851	11851	11851
3211	10408	10408	10408
3221	13526	13526	13526

Table 4. Number of weighted steps for each data partition, the combination of them (MOL) ${\rm and} \ _{\rm W} {\rm ILD} \ {\rm value}$

The optimal parameter set is indicated in italics

	188	28\$	COI	MOL	wILD
111	1125	3967	1354	6520	0.01135
121	1655	6272	2051	10076	0.00973
211	1246	4840	1381	7543	0.01008
221	1867	7780	2080	11851	0.01046
3211	1704	6535	2074	10408	0.00913
3221	2314	8305	2777	13526	0.00961

Table 5. Length of each data partition (28S rRNA is divided into three amplicons) and total length of alignment

IA (121) is for implied alignment under parameter set 121; IA+Gb is for implied alignment trimmed with Gblocks; Muscle is for MUSCLE multiple sequence alignment; Muscle+Gb is for multiple sequence alignment trimmed with Gblocks

	185	28Sa	28Sbc	COI	TOTAL
Unaligned	1760-1805	832-873	1265-1347	654-657	
IA (3211)	1860	1323	1555	669	5407
IA+Gb	1676	378	1162	626	3842
Muscle	1818	1046	1409	663	4936
Muscle+Gb	1695	609	1212	636	4152