Review of Orthochiroides Kovařík, 1998 with description of a new species (Scorpiones: Buthidae)

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May 2022 — No. 349
Euscorpius
Occasional Publications in Scorpiology

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Publication date: 14 May 2022

http://zoobank.org/urn:lsid:zoobank.org:pub:2C279DDB-CF64-480C-8267-38F950B5E785
**Summary**

The genus *Orthochiroiides* Kovařík, 1998 is reanalyzed. Revised diagnoses and new illustrations for the genus and all four of its species are presented. A new species, *O. somalilandus* sp. n. from Somaliland is described and illustrated. Phylogenetic relationships of the genus with several other similar genera of small buthids are inferred from a parsimony analysis of 43 discrete morphological characters. The recent synonymy of *Orthochiroiides* with *Orthochirus* is refuted and the genus is revalidated.

**Introduction**

In 1899, Pocock described *Butheolus insularis*, a small, dark scorpion from Socotra Island (Yemen) with a trapezoidal carapace, short pedipalps and incrassate, heavily sclerotized posterior metasomal segments IV–V. The lateral and ventral surfaces of those segments were smooth, and were marked with numerous small, shallow depressions or punctae. The punctate metasoma differed from that of the type species of the genus, *Butheolus thalassinus* Simon, 1882, in which the posterior metasomal segments are granulate and non-punctate. However, it was similar to that of *B. aristidus* Simon, 1992, originally also described under the same genus. Subsequently, Simon (1910) transferred *B. aristidus* to *Orthochirus* Karsch, 1891, a genus characterized by smooth, punctate metasomal segments. Birula (1917) listed under *Orthochirus* several other species with punctate metasomal segments that were previously assigned to *Butheolus*, including *B. insularis*. Levy & Amitai (1980), in a checklist, downgraded *Orthochirus insularis* to a subspecies of *Orthochirus bicolor* (Pocock, 1897), without explanation.

Kovařík (1998) described the monotypic genus *Orthochiroiides*, with type species *Orthochiroiides vachoni* from Somalia. The species was named after Max Vachon, who in 1976 examined all 38 type specimens (VA No. 1405). Vachon sorted the material into males, females and juveniles, and attached to it the label “*Orthochiroiides* gen. nov.”, a name that he never published. *Orthochiroiides vachoni* is a small, dark scorpion, superficially resembling *Orthochirus* in habitus, but differing in at least two characters: the shape of the telson vesicle is bulbous in *O. vachoni*, vs. slender and pyriform in *Orthochirus*; and the morphosculture of the lateral and ventral surfaces of metasoma IV–V is rugose-reticulate with large depressions in *O. vachoni*, vs. smooth with small punctae in *Orthochirus*. Subsequently, Kovařík (2004) revised *Orthochiroiides*, expanding it to include *Butheolus insularis*, and described a second species from Socotra Island, *O. socotrensis*. The two Socotra species both bear smooth, punctate metasomal segments, a surface morphosculpture similar to that of *Orthochirus*. This left the shape of the telson as the main diagnostic character for separating the two genera.

Recently, the genus *Orthochiroiides* was rejected by Lourenço & Ythier (2021). After studying a paratype male of *Orthochiroiides vachoni* deposited in the Paris museum (MNHN), these authors concluded that telson shape was not a valid generic character, that no other differences from *Orthochirus* justified generic status, and hence that *Orthochiroiides* Kovařík, 1998, was a junior synonym of *Orthochirus* Karsch, 1891. Here, we revisit the question about the validity of the genus *Orthochiroiides*. We analyze in detail the morphology of the species previously assigned to *Orthochiroiides*. We compare them to species described under *Orthochirus*, and several other genera of small buthids that share a number of characters with *Orthochirus*. Our results support the reinstatement of *Orthochiroiides*. We also describe a new species of *Orthochiroiides*.

**Methods, Material & Abbreviations**

Nomenclature and measurements generally follow Stahnke (1971), Sissom et al. (1990), Kovařík (2009), Kovařík & Ojajüren Affilastro (2013) and Lowe et al. (2014). Nomenclature of trichobothria largely follows Vachon (1974, 1975), of hemispermatophores Kovařík et al. (2018), and of pedipalp chela carination, Acosta et al. (2008). External morphology was examined under a dissecting microscope, viewing reflected white light or fluorescence emission under UV LED illumination. Hemispermatophore capsules were
imaged with a Mitutoyo M Plan Apo 10X objective. Focus stacking was implemented in Zerene Stacker 1.04 (Zerene Systems, LLC). Biometrics were measured with an ocular reticule or by digital image analysis with ImageJ 1.52a (Rasband, 2018). Cladistic analyses were conducted in TNT 1.5 (Goloboff & Catalano, 2016). Heuristic searches were performed by generating 1,000 random addition sequences with tree-bisection-reconnection (TBR) branch swapping, holding 50 trees per replicate. Trees were collapsed during searches with minimum length zero. Searches were performed under equal weights (EW), and under implied weights (IW) (Goloboff, 1993) testing a range of concavity constants. Unambiguous synapomorphies were mapped in TNT to the most parsimonious trees (MPTs) retrieved. Consistency indices (CI) and retention indices (RI) of trees and characters were calculated using scripts “stats.run” and “wstats.run”. Node supports and average tree supports were estimated by jackknife resampling (4,000 pseudoreplicates, probability 36%) expressed as group present/ contradicted (GC) frequency differences (Goloboff et al., 2003). Absolute or relative Bremer supports were also estimated, using up to 40,000 suboptimal trees generated by successive TBR branch swapping of MPTs and increasingly suboptimal trees. Specimen depositories: BMNH (The Natural History Museum, London, United Kingdom); FKCP (František Kovařík, private collection, Prague, Czech Republic); will in future be merged with the collections of the National Museum of Natural History, Prague, Czech Republic); GLPC (Graeme Lowe, private collection, Auckland, New Zealand); MNHN (Muséum National d’Histoire Naturelle, Paris, France); MNHW (Museum für Naturkunde der Humboldt-Universität, Berlin, Germany); MZUF (Museo Zoologico de "La Specola", Firenze, Italy); NHMB (Naturhistorisches Museum, Basel, Switzerland); NMPC (National Museum of Natural History, Prague, Czech Republic); NMK (National Museums of Kenya, Nairobi, Kenya); ONHM (Oman Natural History Museum, Muscat, Oman); USNM (United States National Museum of Natural History (Smithsonian Institution), Washington, DC, USA); ZMHB (Museum für Naturkunde der Humboldt-Universität, Berlin, Germany); WDS (W. David Sissom, private collection, Canyon, Texas, USA); and ZMUH (Centrum für Naturkunde (CeNak), Center of Natural History Universität Hamburg, Zoological Museum, Hamburg, Germany). Abbreviations: morphometrics: D, depth; L, length; W, width; movable finger dentition: ID, inner denticles; OD, outer denticles.

### Systematics

**Family Buthidae C. L. Koch, 1837**


(Figures 1–121, 130–133, 140–146, 166–167, 205–211; Tables 1–5)


**Type species.** *Orthochiroioides vachoni* Kovařík, 1998.

**Diagnosis (adults).** Total length of adults up to 24 mm (♂), or 34 mm (♀). Carapace strongly trapezoidal, posterior width/ anterior width > 2.2 (♂), > 2.3 (♀); surface densely, finely granulate, carinae indistinct; median eyes and ocular tubercle located in posterior 2/3 of carapace; preocular area in lateral view inclined downward from median eyes to anterior margin of carapace; 5 pairs of lateral eyes present, of which 2 pairs are either smaller or indistinct (Type 5 pattern; Loria & Prendini, 2014). Pectines hirsute, with fulcra, without enlarged basal teeth or enlarged basal middle lamellae. Pectinal tooth count range: 16–20 (♂), 14–18 (♀). Tergites I–VI either weakly mono- or tricarinata, or with carinae indistinct. Stermites IV–V, and maybe III, with prominent posterior median smooth patches; sternite VI antero-medial surface matte; posterior margins of sternites III–VI with fringes of regular, non-contiguous, enlarged, blunt denticles; spiracles hemi-elliptic or broadly slit-like. Metasomal segments relatively short, stout, either almost uniform in width or slightly increasate posteriorly (metasoma V W/ metasoma I W < 1.1); carination moderately or weakly developed on metasoma I–II, weak or mostly masked by heavy cuticular morphosculpature on metasoma III–V; ventromedian surface of metasoma I granulate; lateral and ventral surfaces of metasoma IV–V either smooth and punctate, or rugose-recticulate. Antero-lateral corners of metasomal segments II–IV flush, not extended anteriorly and outwardly in wedge-shaped processes; metasomal segments II–III widening gradually in front of posterior margins. Posterior margins of tergite VII and metasomal segments I–III bearing fine fringes of microsetae. Telson with vesicle bulbous, aculeus shorter than vesicle, stout, strongly curved with tip angled > 90° relative to rostrocaudal axis; telsin L/D < 2.4, telsin W/ metasoma V W > 0.5; subocular tubercle absent. Chelicerae with typical buthid dentition (Vachon, 1963), fixed finger with two denticles on ventral surface. Pedipalps relatively short, chelae narrower than patella; patella with strongly costate dorsomedian and dorsoexternal carinæ; chela manus with carinae V1, VA, D3 and D4 strongly costate, VA complete. Trichobothrial pattern neobothiotoxic type C (Vachon, 1974); femur with petite ‘trichobothrium’ d₅ absent on dorsal surface, d₁-d₃, d₄ in β-configuration (Vachon, 1975), e₁ proximal to d₅ positioned at base of femur; patella with petite ‘trichobothrium’ d₅ reduced or absent, d₄ located between dorsomedian and dorsointernal carinae, esb₁ near esb₂ (< 0.18 distance to em), em about midway between esb₂ and et; chela manus with Eb₁ proximal to Eb₂, V₁-V₄ axis slightly inclined internally, eb positioned on distal manus, not fixed finger; fixed finger with db positioned in middle 30-60% of finger, proximal to est; dentate margins of pedipalp fingers straight, without scalloping or lobe/ notch combination; movable finger equipped with 7–9 rows of median denticles.
Figures 1–4: Orthochiroides insularis, Socotra Island. Figures 1–2. Male, dorsal (1) and ventral (2) views. Figures 3–4. Female, dorsal (3) and ventral (4) views. Scale bar: 10 mm.
arranged almost linearly, non-imbricated, each flanked by a single external (ED) and internal (ID) accessory denticle; 5 subterminal denticles present. Mid-ventral aspect of tarsomere II of legs sparsely setose with 1 or 2 rows of short spiniform setae; tibial spurs present on legs III–IV.

COMMENTS. Hemispermatophore characters are omitted from the generic diagnosis. We examined hemispermatophores from only one species, Orthochiroides insularis. However, general features such as a sperm hemiduct divided into 3 lobes, and a folded flagellum with a broad, laminate pars recta and a narrow cylindrical pars seminalis have been studied in genera of the Orthochiridae. We examined hemispermatophores from only one species, Orthochiroides insularis (Pocock, 1899) (Figs. 1–4, 82, 87–94, 111, 139–133, 140–142, 166, 205–211; Tables 1, 3).


Type locality and type repository. Socotra, Mt. Raggit; BMNH.

Type material examined. Yemen, Socotra Island, Hadibu Plain (Mt. Raggit, 1000 feet), 1♀ (holotype), leg. Grant and Forbes, BMNH No. 1899.7.4.180.

Other material examined (FKCP). Yemen, Socotra Island, Qalansiyah env., Ditwah (lagoon), 23 m a. s. l., 12°41′42″N 53°30′08″E, 9.XII.2003, 1♀ (Figs. 1–2, 82, 87–90, 111), leg. D. Král; Gubbah village env., 7 m a. s. l., 12°36′35″N 53°46′56″E, 23.XI.2003, 1♀ (Figs. 3–4, 91–94, 112) 1juv., leg. D. Král; Qaariah village env., 11 m a. s. l., 12°38′05″N 54°12′39″E, 28.XI.2003, 1♂, leg. J. Farkač.

Diagnosis (adult ♀). Total length of adults 22–32 mm. Petite ‘trichobothria’ d₁ on dorsal surfaces of pedipalp femur and patella reduced or absent. Chela smooth, carinae E and D₁ weak or obsolete on manus. Pectinal tooth count: 18–19 (♂), 16–18 (♀). Movable finger of pedipalp chela with 7 rows of median denticles, flanked by 7 ID and 7 OD. Metasoma V–V laterally and ventrally punctate, without carinae except for smooth, rugose dorsosubmedian carinae, and ventrolateral carinae on metasoma V which are anteriorly smooth and posteriorly denticulate. Metasoma III either laterally and ventromedially punctate (♂), or laterally punctate and ventromedially granulate-reticulate (♀). Punctae small, with diameters less than the mean distance between their centroid, together occupying much less area than smooth surfaces between them. Dorsal surfaces of all metasomal segments smooth, without punctae or granules. Aculeus much shorter than vesicle, aculeus L/telson L ~0.20. Color uniformly brown to black. Femur, patella and chela manus of pedipalp brown to black; chela fingers, and tibia and tarsomeres of legs yellowish brown to green. Tergites roughly granulated. Stermites IV–V with lateral surfaces matte, almost smooth. Sternite VII matte, almost smooth, with four distinct granulated carinae. Pedipalp, metasoma and telson glabrous, only metasoma V may have dorsolateral rows of setae, which can also occur on dorsal surface of telson. Moderately developed tibial spurs present on legs III and IV. Tarsomere I of legs I–III with 4–6 long setae in both sexes.

Distribution. Yemen, north part of Socotra Island (Fig. 119).

Comments. Fet & Lowe (2000: 194) listed “Orthochirus socotrensis” cited by Francke (1977) as a possible lapsus and synonym of Orthochirus bicolor insularis. However, Francke (1977: 112) cited the presence of three endemic species of scorpion on Socotra Island: Hemiscorpius socotranus Pocock, 1899, Butheolus insularis Pocock, 1899 and Orthochirus socotrensis (Pocock). He omitted Butheolus socotrensis Pocock, 1899 (= Buthotus socotrensis = Hottentotta socotrensis), also known from Socotra Island. It is possible that Francke cited Orthochirus socotrensis in error, instead of Butheolus sociotrensis, as Pocock did not describe Orthochirus socotrensis (see also Vachon, 1979: 237).
Table 1. Comparative measurements of Orthochiroides somalilandus sp. n. and O. vachoni specimens. Abbreviations: length (L), width (W, in carapace it corresponds to posterior width), depth (D).

<table>
<thead>
<tr>
<th>Dimensions (mm)</th>
<th>O. somalilandus sp. n.</th>
<th>O. somalilandus sp. n.</th>
<th>O. vachoni</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>♂ holotype</td>
<td>♀ paratype</td>
<td>♂ Socotra</td>
</tr>
<tr>
<td>Carapace</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L / W</td>
<td>3.29 / 3.90</td>
<td>3.86 / 4.97</td>
<td>3.03 / 3.74</td>
</tr>
<tr>
<td>Mesosoma</td>
<td>7.15</td>
<td>9.50</td>
<td>6.64</td>
</tr>
<tr>
<td>Tergite VII</td>
<td>1.87 / 3.93</td>
<td>2.29 / 4.82</td>
<td>1.87 / 3.76</td>
</tr>
<tr>
<td>Metasoma + telson</td>
<td>16.03</td>
<td>17.89</td>
<td>15.55</td>
</tr>
<tr>
<td>Segment I</td>
<td>1.91 / 2.65 / 2.19</td>
<td>2.11 / 3.00 / 2.54</td>
<td>1.84 / 2.49 / 2.20</td>
</tr>
<tr>
<td>Segment II</td>
<td>2.24 / 2.68 / 2.19</td>
<td>2.51 / 2.99 / 2.38</td>
<td>2.16 / 2.60 / 2.16</td>
</tr>
<tr>
<td>Segment III</td>
<td>2.44 / 2.75 / 2.28</td>
<td>2.68 / 3.15 / 2.43</td>
<td>2.27 / 2.74 / 2.32</td>
</tr>
<tr>
<td>Segment IV</td>
<td>2.94 / 2.86 / 2.31</td>
<td>3.23 / 3.13 / 2.46</td>
<td>2.75 / 2.83 / 2.36</td>
</tr>
<tr>
<td>Segment V</td>
<td>3.43 / 2.71 / 2.18</td>
<td>3.83 / 3.13 / 2.40</td>
<td>3.44 / 2.78 / 2.18</td>
</tr>
<tr>
<td>Telson</td>
<td>3.07 / 1.46 / 1.36</td>
<td>3.53 / 1.77 / 1.61</td>
<td>3.09 / 1.44 / 1.33</td>
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<tr>
<td>Pedipalp</td>
<td>8.46</td>
<td>9.69</td>
<td>8.29</td>
</tr>
<tr>
<td>Femur</td>
<td>1.90 / 0.99</td>
<td>2.18 / 1.02</td>
<td>1.86 / 0.81</td>
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<tr>
<td>Patella</td>
<td>2.75 / 1.30</td>
<td>3.21 / 1.49</td>
<td>2.75 / 1.20</td>
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<tr>
<td>Chela</td>
<td>3.81</td>
<td>4.30</td>
<td>3.68</td>
</tr>
<tr>
<td>Manus</td>
<td>0.98 / 0.95</td>
<td>1.16 / 1.14</td>
<td>0.92 / 0.91</td>
</tr>
<tr>
<td>Movable finger</td>
<td>2.51</td>
<td>2.73</td>
<td>2.43</td>
</tr>
<tr>
<td>Total</td>
<td>26.47</td>
<td>31.25</td>
<td>25.22</td>
</tr>
</tbody>
</table>

for smooth, rugose dorsosubmedian carinae, and ventrolateral carinae on metasoma V which are anteriorly obsolete and posteriorly weakly denticulate. Punctae small with diameters less than the mean distance between their centroids, together occupying much less area than smooth surfaces between them. Dorsal surfaces of all metasomal segments smooth without punctae or granules. Aculcus much shorter than vesicle, aculeus L/telson L ~0.20. Color of mesosoma dark green, metasoma and telson reddish brown; pedipalps and legs yellow to yellowish green, lighter on more distal segments. Tergites roughly granulated. Sternites IV–V with lateral surfaces coarsely granulated. Sternite VII with moderately dense, coarse granulation throughout, and four granulated carinae. Pedipalp, metasoma and telson glabrous, only metasoma V may have dorsolateral rows of setae, which can also occur on dorsal surface of telson. Moderately developed tibial spurs present on legs III and IV. Tarsomere I of legs I–III with 4–6 long setae in both sexes.

**Distribution.** Yemen, south part of Socotra Island (Fig. 119).


**Type locality and type repository.** Somaliland, Mader Mage village, between Eregavo and Maid, 10°48’03”N 47°17’46”E, 1389 m a. s. l. (Fig. 76); FKCP.

**Type material examined (FKCP).** Somaliland, Mader Mage village, between Eregavo and Maid, 10°48’03”N 47°17’46”E, 1389 m a. s. l. (locality No. 18SD), 23.VIII.2018, 3♂ (holotype and paratypes, DNA Nos. 1543, 1544); leg. F. Kovářík; 5 km S of Maid, 10°59’46”N 47°08’14”E, 182 m a. s. l. (Locality No. 18SF; Fig. 77), 25.VIII.2018, 3♂ (paratypes, DNA No. 1539, Figs. 70–73), leg. F. Kovářík.

**Etymology.** Named for its geographic distribution.

**Diagnosis (adult ♂♀).** Total length of adults 24–32 mm. Petite ‘trichobothria’ d on dorsal surfaces of pedipalp femur and patella reduced or absent. Chela with carinae E and D1 strongly costate, intercarinal surfaces sparsely granulate. Pectinal tooth count: 16–18 (♂), 15 (♀). Movable finger of pedipalp chela with 9 rows of median denticles, 9 ID and 9 OD. Metasoma V laterally and ventromedially rugose-reticulate and finely granulate. Metasoma III–IV laterally rugose-reticulate and finely granulate, ventromedially granulate. Inter-reticular depressions large, irregularly shaped, with diameters comparable to the mean distance between their centroids, together occupying much more area than the smooth or granular reticulations between them. Weak, finely granulated median lateral carinae present and complete on metasoma I–II, indistinct and masked by reticulate morphosculpture on metasoma III–IV. Dorsal surfaces of all metasomal segments smooth medially, granulate-reticulate or rugose-reticulate laterally; border between dorsal and dorsolateral surfaces of metasoma IV–V indistinct, not clearly demarcated by dorsosubmedian carina or abrupt transition.
Figures 9–12: Orthochiroides somalilandus sp. n. Figures 9–10. Male holotype, dorsal (9) and ventral (10) views. Figures 11–12. Female paratopotype, dorsal (11) and ventral (12) views. Scale bar: 10 mm.
Figures 13–18: Orthochiroides somalilandus sp. n. Figures 13–15. Female paratopotype, metasoma and telson, lateral (13), ventral (14), and dorsal (15) views. Figures 16–18. Male holotype, metasoma and telson, lateral (16), ventral (17), and dorsal (18) views. Scale bars: 10 mm (13–15, 16–18).
Figures 19–37: Orthochiroides somalilandus sp. n., pedipalp. Figures 19–27. Male holotype, chela, dorsal (19), external (20), and ventral (21) views. Patella, dorsal (22), external (23) and ventral (24) views. Femur and trochanter, internal (25), dorsal (26), and ventral (27) views. Figures 28–37. Female paratopotype, chela, dorsal (28), external (29), and ventral (30) views. Patella, dorsal (31), external (32) and ventral (33) views. Femur and trochanter, internal (34), dorsal (35), and ventral (36) views. Movable finger dentation (37). Trichobothrial pattern indicated in Figures 19–23, 25–26 by white circles.
Figures 38–45: Orthochoiroides somalilandus sp. n. Figures 38, 40, 42–45. Male holotype, carapace and tergites I–V (38), sternopercinal region and sternites (40), and left legs I–IV, retrolateral aspect (42–45). Figures 39, 41. Female paratopotype, carapace and tergites I–V (39), sternopercinal region and sternites (41).
in morphosculpture. Aculeus slightly shorter than vesicle, aculeus L/telson L > 0.30. Color uniformly black except for pedipalp chela fingers and tarsomeres of legs, which are yellow to yellowish brown. Tergites roughly granulated. Sternite VII densely granulated, with four granulated carinae. Pedipalp, metasoma and telson glabrous. Moderate to small tibial spurs present on legs III and IV. Tarsomere I of legs I–III with 3–4 long setae in both sexes.

**Description.** Total length of adults 24–32 mm in both sexes. For habitus, see Figs. 9–12, 50–51.

**Coloration** (Figs. 9–12, 50–51). Color uniformly black, except for pedipalp chela fingers and tarsomeres of legs which are yellow to yellowish brown. Telson reddish brown.

**Carapace and mesosoma** (Figs. 38–41). Carapace surface roughly granulated except for several smooth furrows. Superciliary carinae granulated, extending partially into preocular triangle. Tergites I–VI roughly granulated, with weak median and lateral carinae. Sternite VII densely granulated with four granulated carinae; other sternites finely granulated or shagreened in lateral areas. Sternite VI with median carinae granulated (♂), or smooth (♀), lateral carinae granulated in both sexes. Glossy smooth patch present on median-posterior zone of sternites IV–VI (♂), or sternites III–VI (♀). Posterior margins of sternites III–VI (♂), or III–V (♀) equipped with rows of non-contiguous clavate or digitate denticles. Pectinal tooth count: 16–18 (3 x 16, 3 x 17, 4 x 18) (♂), 15 (♀). Female pectine teeth notably shorter than male teeth, mid-pectine sensillar margin L/tooth W 1.63 (♀), 2.45 (♂).

**Hemispermatophore** (Figs. 46–49). Flagelliform. Trunk narrow, elongate, broadened proximally. Flagellum folded, with thicker, laminate pars recta, 0.45 times length of trunk, and thinner, cylindrical, hyaline pars reflecta, 0.61 times length of trunk. Capsule short, 0.15 times length of trunk. Sperm hemiduct divided into 3 lobes; posterior lobe longest, lanceolate; median lobe shortest, laminate; median and anterior lobes apically acuminate. Basal lobe an apically curved hook with a broad base.

**Metasoma and telson** (Figs. 13–18). Metasoma I–II with 10 granulated carinae. Metasoma III–IV without distinct median lateral carinae. Metasoma V with dorsosubmedian carinae smooth, rugose, irregular; ventrolateral carinae weak, granulated, restricted to posterior third of segment, other carinae absent. Metasoma I with intercarinal surfaces finely granulated.
Figures 50–51. Orthochiroides somaliandus sp. n., paratypes male (50) and female (51) in vivo (laboratory photographs on standard background).
Figures 52–53: Orthochiroides somalilandus sp. n., localities. Figure 52. Type locality, Somaliland, Mader Mage vill., between Eregavo and Maid. Figure 53. Somaliland, 5 km S of Maid.

Pedipalps (Figs. 19–37). Petite ‘trichobothrium’ d₁ on dorsal surface of pedipalp femur absent. Femur with five granulated carinae, intercarinal surfaces densely granulated. Patella with seven moderately to strongly costate, smooth or granulated carinae; intercarinal surfaces with sparse fine granulation, or almost smooth. Chela with six complete, strongly costate, smooth or weakly crenulate carinae (V₁, V₂, E, D₁, D₃ and D₄), with D₁, D₃ and D₄ extending along full length of fixed finger; intercarinal surfaces with sparse fine granulation, or almost smooth. All pedipalp segments glabrous, bearing only several setae. Movable fingers with 9 rows of denticles, 9 ID and 9 OD.

Legs (Figs. 42–45). Legs III–IV with moderate to small tubial spurs. Femur with four granulated carinae; patella with five carinae. Patella with only a few macrosetae. Tibia with macrosetae on the retroinferior aspect of legs I–II. Tarsomere I of legs I–III with 3–4 long macrosetae in both sexes, of legs IV with 2 setae. Tarsomeres I–II of all legs with two irregular rows of macrosetae on proinferior and retroinferior aspects.

Measurements. See Table 1.

Comments on locality and life strategy. The type locality is a montane slope with trees and bushes (Fig. 52, fig. 18SF). The minimum recorded temperature was 24°C. The minimum recorded humidity was 37%. The locality was first recorded a minimum nighttime temperature of 24°C. At this locality, the type specimens were collected among rocks near Mader Mage village at an approximate elevation of 1,389 m a. s. l. At this locality, the other two species of Orthochiroides found on granulate. Metasoma III–IV laterally rugose-reticulate and finely granulate, ventromedially finely granulate. Metasoma V laterally and ventromedially rugose-reticulate, finely granulate in posterior ventral area. Reticulations on metasoma III–V thinner and partially granulated in male, thicker and polished in female. Dorsal surfaces of metasoma I–IV smooth medially, of metasoma V smooth posterio-medially; lateral areas of dorsal surfaces granulate on segment I, granulate reticulate on II, rugose-reticulate on III–V. Entire metasoma and telson glabrous. Telson smooth and punctate.

Pedipalps (Figs. 19–37). Petite ‘trichobothrium’ d₁ on dorsal surface of pedipalp femur absent. Femur with five granulated carinae, intercarinal surfaces densely granulated. Patella with seven moderately to strongly costate, smooth or granulated carinae; intercarinal surfaces with sparse fine granulation, or almost smooth. Chela with six complete, strongly costate, smooth or weakly crenulate carinae (V₁, V₂, E, D₁, D₃ and D₄), with D₁, D₃ and D₄ extending along full length of fixed finger; intercarinal surfaces with sparse fine granulation, or almost smooth. All pedipalp segments glabrous, bearing only several setae. Movable fingers with 9 rows of denticles, 9 ID and 9 OD.

Legs (Figs. 42–45). Legs III–IV with moderate to small tubial spurs. Femur with four granulated carinae; patella with five carinae. Patella with only a few macrosetae. Tibia with macrosetae on the retroinferior aspect of legs I–II. Tarsomere I of legs I–III with 3–4 long macrosetae in both sexes, of legs IV with 2 setae. Tarsomeres I–II of all legs with two irregular rows of macrosetae on proinferior and retroinferior aspects.

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Pedipalps (Figs. 19–37). Petite ‘trichobothrium’ d₁ on dorsal surface of pedipalp femur absent. Femur with five granulated carinae, intercarinal surfaces densely granulated. Patella with seven moderately to strongly costate, smooth or granulated carinae; intercarinal surfaces with sparse fine granulation, or almost smooth. Chela with six complete, strongly costate, smooth or weakly crenulate carinae (V₁, V₂, E, D₁, D₃ and D₄), with D₁, D₃ and D₄ extending along full length of fixed finger; intercarinal surfaces with sparse fine granulation, or almost smooth. All pedipalp segments glabrous, bearing only several setae. Movable fingers with 9 rows of denticles, 9 ID and 9 OD.

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Pedipalps (Figs. 19–37). Petite ‘trichobothrium’ d₁ on dorsal surface of pedipalp femur absent. Femur with five granulated carinae, intercarinal surfaces densely granulated. Patella with seven moderately to strongly costate, smooth or granulated carinae; intercarinal surfaces with sparse fine granulation, or almost smooth. Chela with six complete, strongly costate, smooth or weakly crenulate carinae (V₁, V₂, E, D₁, D₃ and D₄), with D₁, D₃ and D₄ extending along full length of fixed finger; intercarinal surfaces with sparse fine granulation, or almost smooth. All pedipalp segments glabrous, bearing only several setae. Movable fingers with 9 rows of denticles, 9 ID and 9 OD.

Legs (Figs. 42–45). Legs III–IV with moderate to small tubial spurs. Femur with four granulated carinae; patella with five carinae. Patella with only a few macrosetae. Tibia with macrosetae on the retroinferior aspect of legs I–II. Tarsomere I of legs I–III with 3–4 long macrosetae in both sexes, of legs IV with 2 setae. Tarsomeres I–II of all legs with two irregular rows of macrosetae on proinferior and retroinferior aspects.

Measurements. See Table 1.

Comments on locality and life strategy. The type locality is a montane slope with trees and bushes (Fig. 52, fig. 18SF). The minimum recorded temperature was 24°C. The minimum recorded humidity was 37%. The locality was first recorded a minimum nighttime temperature of 24°C. At this locality, the type specimens were collected among rocks near Mader Mage village at an approximate elevation of 1,389 m a. s. l. At this locality, the other two species of Orthochiroides found on granulate. Metasoma III–IV laterally rugose-reticulate and finely granulate, ventromedially finely granulate. Metasoma V laterally and ventromedially rugose-reticulate, finely granulate in posterior ventral area. Reticulations on metasoma III–V thinner and partially granulated in male, thicker and polished in female. Dorsal surfaces of metasoma I–IV smooth medially, of metasoma V smooth posterio-medially; lateral areas of dorsal surfaces granulate on segment I, granulate reticulate on II, rugose-reticulate on III–V. Entire metasoma and telson glabrous. Telson smooth and punctate.

Pedipalps (Figs. 19–37). Petite ‘trichobothrium’ d₁ on dorsal surface of pedipalp femur absent. Femur with five granulated carinae, intercarinal surfaces densely granulated. Patella with seven moderately to strongly costate, smooth or granulated carinae; intercarinal surfaces with sparse fine granulation, or almost smooth. Chela with six complete, strongly costate, smooth or weakly crenulate carinae (V₁, V₂, E, D₁, D₃ and D₄), with D₁, D₃ and D₄ extending along full length of fixed finger; intercarinal surfaces with sparse fine granulation, or almost smooth. All pedipalp segments glabrous, bearing only several setae. Movable fingers with 9 rows of denticles, 9 ID and 9 OD.

Legs (Figs. 42–45). Legs III–IV with moderate to small tubial spurs. Femur with four granulated carinae; patella with five carinae. Patella with only a few macrosetae. Tibia with macrosetae on the retroinferior aspect of legs I–II. Tarsomere I of legs I–III with 3–4 long macrosetae in both sexes, of legs IV with 2 setae. Tarsomeres I–II of all legs with two irregular rows of macrosetae on proinferior and retroinferior aspects.

Measurements. See Table 1.
Figures 54–57: Orthochiroides vachoni, paratypes, FKCP. Figures 54–55. Male, dorsal (78) and ventral (79) views. Figures 56–57. Female, dorsal (80) and ventral (81) views. Scale bar: 10 mm.
Socotra (O. insularis and O. socotrensis) are endemic, as are two other Socotra scorpions, Hemiscorpius socotranus Pocock, 1889 and Hottentotta socotrensis (Pocock, 1889). This suggests that ‘O. vachoni’ from Socotra might also be an endemic species different from O. vachoni on the mainland. However, the Socotra specimen is virtually identical to all paratype males of O. vachoni that we have examined (Figs. 58–81, Tab. 1). There are only a few minor differences that lie well within the scope of intraspecific variation typical for other buthids (i.e., sternite VII fine granulation slightly more intense; metasoma III-IV median lateral carinae slightly less visible and partially obscured in both by strong reticulate morphosculpture; small morphometric differences in metasoma V). In our opinion, these differences are insufficient for the diagnosis of a new species.

The other mainland African species O. somalilandus sp. n. differs more from O. vachoni, than does the specimen from Socotra. The collection of an extensive type series of O. vachoni from southern Somalia (Kovařík, 1998), and the presence of a closely similar species in Somaliland, indicate that this species occurs naturally on the mainland. We cannot exclude the possibility that the Socotra record is spurious and possibly an introduction by human transport. Other non-endemic flora and fauna occur on or have been introduced to the island (e.g., Hůla & Niedobová, 2020; Senan et al., 2010; Witt et al., 2020). The range of distribution of O. vachoni in Somalia is unknown. It may not be confined to its type locality, and could extend further.

Figures 58–59. Orthochiroides vachoni, male from Socotra, dorsal (58) and ventral (59) views.
up the coast closer to Socotra. Without further information, we take a conservative approach and provisionally classify the Socotra record as *O. vachoni*.

In their paper synonymizing *Orthochiroides* with *Orthochirus*, Lourenço & Ythier (2021) examined a single male paratype of *Orthochiroides vachoni* deposited in MNHN, and illustrated its metasomal segment V and telson in lateral view (Lourenço & Ythier, 2021: 344, fig. 20). They depicted the morphosculpture on the lateral surface of metasoma V as consisting of numerous small, rounded depressions or punctae of uniform size and shape, with diameters much less than the mean distance between their centroids, that together occupy much less area than the smooth surfaces between them. This is similar to the punctate morphosculpture on metasoma V of most species of *Orthochirus*. Their illustration differs from the illustration of a rugose-reticulate morphosculpture on the lateral surface of the male holotype published in the original description of *O. vachoni* (Kovařík, 1998: 117, fig. 1). In that figure, the depressions are depicted as more irregular in size and shape, including larger diameters comparable to the mean distance between their centroids, together occupying as much or more area than the surface between them. Kovařík (1998: 117–119, figs. 4–6) described the larger depressions on the metasoma of *O. vachoni* and emphasized their difference from the punctae of *Orthochirus*. This morphosculpture is documented again here photographically for comparison to fig. 20 of Lourenço & Ythier (2021) (cf. Figs. 60–63, 84–86). Kovařík (1998) analyzed the entire type series of *O. vachoni* consisting of 38 specimens, including the paratype deposited in MNHN, and did not report any examples of fine punctate morphosculpture on the metasoma. We regard fig. 20 of Lourenço & Ythier (2021) to be a gross misrepresentation of the true anatomy of *O. vachoni*.
Figures 64–81: Orthochiroides vachoni, male from Socotra. Figures 64–75. Pedipalp. Chela, dorsal (64), external (65), and ventral (66) views. Patella, dorsal (67), external (68) and ventral (69) views. Femur and trochanter, internal (70), dorsal (71), ventral (72), and external (73) views. Movable (74) and fixed (75) finger dentation. Trichobothrial pattern indicated in Figures 64–68, 70–71 by white circles. Figures 76–77. Carapace and tergites I–VI (76), sternopectinal region and sternites (77). Figures 78–81. Right legs I–IV, retrolateral aspect.
Key to species of Orthochiroides

1. Movable finger of pedipalps with 7 rows of median denticles, 7 ID and 7 OD (Figs. 111–114); metasoma IV–V of adults ventrally smooth and punctate; punctae on ventral surface of metasoma V small, with diameters less than mean distance between their centroids, occupying less area than smooth surfaces between them (Figs. 82–83); pedipalps more slender, patella L/W 2.5–3.0 ........... 2
   – Movable finger of pedipalps with 8–9 rows of median denticles, 9 ID and 9 OD (Figs. 115–118); metasoma IV–V of adults ventrally rugose-reticulate; inter-reticular depressions or lacunae on ventral surface of metasoma V large, irregularly shaped, with diameters comparable to mean distance between their centroids, occupying more area than raised reticular surfaces (Figs. 13–18, 60–63, 84–86); pedipalps more stout, patella L/W 2.1–2.3 ........ 3

2. Pedipalps with patella and chela manus brown or black, darker than fingers (Figs. 87–94); sternite VII matte, almost smooth ........................................... O. insularis (Pocock, 1899)
   – Pedipalps with patella and entire chela yellow (Figs. 95–102); sternite VII coarsely granulated ......................................................... O. socotrensis Kovařík, 2004

3. Border between dorsomedian and dorsolateral surfaces of metasoma IV–V distinct, demarcated by dorsosubmedian carina; dorsal surfaces of metasoma III–V weakly rugose laterally (Fig. 86); metasoma IV ventromedially rugose-granulate (Fig. 85) ...................... O. vachoni Kovařík, 1998
   – Border between dorsomedian and dorsolateral surfaces of metasoma IV–V indistinct, not demarcated by dorsosubmedian carina; dorsal surfaces of metasoma III–V strongly rugose or rugose-reticulate laterally (Fig. 18); metasoma IV ventromedially granulate (Fig. 17). ............................................... O. somalilandus sp. n.
Morphological characters separating *Orthochiroides*
Kovařík, 1998 from *Orthochirus* Karsch, 1891

Kovařík (1998: 116) separated *Orthochiroides* (then monotypic, with type species *Orthochiroides vachoni*) from *Orthochirus* on the basis of telson shape (i.e., bulbous or “inflated” vs. slender respectively) and morphosculture of metasoma V (larger “punctae” occupying a major part of the surface, vs. smaller punctae occupying a minor part of a smooth surface, respectively). The posterior metasoma of *Orthochiroides vachoni* exhibits a distinctive and unique rugose-reticulate morphosculture. The “punctae” of Kovařík (1998) are herein described as inter-reticulate depressions or lacunae. Kovařík (1998: 119–120) also reported an ontogenetic character, in which the metasomal segments of juveniles of *Orthochiroides vachoni* are granulated, and lack the reticulate morphosculture and depressions characteristic of adults. In contrast, the smooth, punctate morphosculture of the metasomal segments typical of adult *Orthochirus* is already well expressed in juveniles of that genus. Other characters separating *Orthochiroides* from *Orthochirus* were implicit in the generic diagnosis of Kovařík (1998: 116): e.g., “four pairs of lateral eyes” and “six pronounced keels on the tibia of pedipalps” (= 6 strong costate carinae on the pedipalp chela).

Kovařík (2004: 23–24) extended the genus *Orthochiroides* to encompass *Butheolus insularis* Pocock, 1899, and described another species, *Orthochiroides socotrensis* Kovařík, 2004. The metasomal segments of these two species exhibit a smooth, punctate morphosculture, similar to that of *Orthochirus*. In a revised diagnosis of *Orthochiroides*, Kovařík (2004: 23) characterized metasoma IV–V as “ventrally punctate” in a broad sense to also cover the different rugose-reticulate condition in *Orthochiroides vachoni*. The only character mentioned that separated *Orthochiroides* from *Orthochirus* was the shape of the telson.

Lourenço & Ythier (2021: 339) dismissed almost all of the diagnostic characters proposed for the genus *Orthochiroides* by Kovařík (1998, 2004) as being “without any generic value”, discussing only two possibly valid characters: (1) four pairs of lateral eyes, which they rejected because they observed only three pairs in the male paratype that they examined; and (2) a bulbous telson, which they confirmed in their male paratype, but dismissed as “not sufficient for the definition of a genus”. In the first case, surveys of the comparative anatomy of lateral eyes in buthids (Loria & Prendini, 2014; Yang et al., 2013) found that five pairs were present in the majority of genera and species, while a minority had reduced counts of 2–4 pairs. Intraspecific variation in lateral eye counts could also occur. Earlier studies may have underestimated the counts, as the typical 5-eye pattern includes 3 larger ocelli (PLMa, MLMa, ALMa) which are readily visible, plus two smaller ocelli (PDMI, ADMi) which are more difficult to detect (“Type 5” configuration of Loria & Prendini, 2014). We re-examined the lateral eyes of *Orthochiroides* and detected up to five pairs in a Type 5 configuration, although in some cases only 3 or 4 were discernible (either PDMI or ADMi may be indistinct). We therefore consider Type 5 to be the basic pattern in *Orthochiroides*, a pattern that is also found in *Orthochirus*. We agree that this character does not separate *Orthochiroides* from *Orthochirus*. However, it does contribute some diagnostic value by excluding those buthid genera with 2–4 pairs of lateral eyes. In the second case, although the difference between telson shapes of *Orthochiroides* and *Orthochirus* was confirmed by Lourenço & Ythier (2021: 339), they nevertheless rejected it as a diagnostic character for *Orthochiroides*. Below, we examine this character and six other characters that we propose for the differential diagnosis of *Orthochiroides* vs. *Orthochirus*. Character numbers and states refer to the list in Table 5.

(i) Carapace shape: *Orthochiroides*: strongly trapezoidal, posterior width/ anterior width > 2.2 (♂), > 2.3 (♀); *Orthochirus*: moderately trapezoidal, posterior width/ anterior width < 2.2 (♂), < 2.3 (♀) (character 6, states 2 and 1). The dorsal profile of a scorpion carapace varies from being almost parallel-sided, to having lateral margins anteriorly convergent. The former shape has been loosely termed ‘sub-rectangular’ and the latter ‘trapezoidal’, although the distinction is subjective without quantitative rules for determining which is applicable. In order to objectively compare carapace shapes of *Orthochiroides* and *Orthochirus*, we measured two morphometric ratios, carapace W/L and carapace posterior W/ anterior W. These ratios parametrize the possible ranges of trapezoidal geometries, mapping them into a 2D morphospace. More ‘strongly’ trapezoidal carapaces are characterized by higher values of these ratios. We analyzed and compared the ratios for *Orthochiroides* (4 spp., 100% of named species) and *Orthochirus* (45 spp., 85% of named species). In Figs. 130–131, ratios for different species of male and female *Orthochiroides* and *Orthochirus* are shown in bivariate scatter plots. Across *Orthochirus* species, posterior W/ anterior W was positively correlated with W/ L (P = 0.00014) in females, but not in males (P = 0.111). In both sexes of *Orthochiroides*, the ratio of posterior W/ anterior W tended to be higher for higher values of W/ L, although the trend was not statistically significant given the small number of species. However, the important result for differential diagnosis is that the morphospace domains occupied by carapaces of *Orthochiroides* and *Orthochirus* are disjunct in both sexes. The two genera can be reliably separated by threshold values of the ratio of posterior W/ anterior W.

(ii) Telson shape: *Orthochiroides*: vesicle bulbous, telson L/D < 2.5 (♂), telson W/ metasoma V W < 0.5 (♀); *Orthochirus*: vesicle slender, pyriform, telson L/D > 2.5 (♂), telson W/ metasoma V W > 0.5 (♀) (character 40, states 1 and 2; character 43, states 0 and 1) (cf. Kovařík, 1998: 116: *Orthochiroides*, “differs in shape of the telson, which is highly inflated”).

The morphometric ratio of telson L/ telson D is a measure of telson elongation. Bulbous telsons have lower values, and slender telsons higher values. We analyzed and compared the ratio for *Orthochiroides* (4 spp., 100% of named species) and
Figures 120–129: Comparison of metasoma V and telson of five ingroup genera in lateral view. **Figures 120–121.** *Orthochiroides somalilandus* sp. n., male holotype (120) and female paratopotype (121). **Figures 122–123.** *Orthochirus scrobiculosus* (Grube, 1873), male topotype (NMB, 122) and female holotype (123). **Figures 124–125.** *Neobuthus eritreaensis* Lowe et Kovařík, 2016, male (124) and female (125) paratypes. **Figures 126–127.** *Butheolus gallagheri* Vachon, 1980, male (126) and female (127) from Oman, Mirbat, 17°02.19’N 54°38.75’E, 54 m a. s. l. (FKCP). **Figures 128–129.** *Gint amoudensis* Kovařík et al., 2018, male holotype (128) and female paratype (129).
Orthochirus (39 spp., 73% of named species). The distribution of the ratio was widely disjunct between Orthochiroides and Orthochirus, with consistently higher values in the latter for both sexes. Two additional morphometric trends may distinguish the two genera: the bulbous telson of Orthochiroides is typically wider than that of Orthochirus, and the posterior metasomal segments of Orthochirus exhibit greater lateral expansion compared to those of Orthochiroides. We analyzed the ratio of telson W/ metasoma V W, and found it to be invariably larger for female Orthochiroides, and mostly larger for males (overlapping for only one species of Orthochirus). Bivariate scatter plots of the two ratios revealed the wide disjunction between the two genera in 2D morphospace (Figs. 132–133). See also Figs. 120–121 vs. figs. 122–123.
Figures 134–135: Morphometric analysis of scorpion telson shape. Bivariate scatter plots of principal component scores, PC1 vs. PC2 (134) and PC3 vs. PC4 (135) explaining 94.36% of the variance of telson lateral profiles parametrized by 7 variables (136): LV (vesicle length), LVA (anterior vesicle length), HV (vesicle depth), HA (aculeus height), SA (aculeus ventral curve length), BT (aculeus basal thickness), and AT (area of subaculear tubercle). First 6 variables normalized to telson length (TL), AT normalized to area of vesicle bounding box (LV.HV). Axis labels list associated variables with factor loadings > 50% (positive or negative). Sample: 2562 profiles; families: 20; genera, 224; species, 1866 (1858 extant, 8 extinct); sex, 1340 ♂ + 1213 ♀ + 9 ?. Data compilation as in Lowe et al, 2019.
Figures 136–141: Variation in telson shape of scorpion genera. Bivariate scatter plots of principal component scores, PC1 vs. PC2 (136, 138, 140) and PC3 vs. PC4 (137, 139, 141) for selected genera: Ananteris (83 spp.), Bothriurus (35 spp.), Buthus (43 spp.) and Tityus (199 spp.) (136–137); Diplocentrus (32 sp.), Euscorpius (43 spp.), Microtityus (32 sp.), Parabuthus (22 sp.) and Reddyanus (27 spp.) (138–139); Baloorthochirus (1 sp.), Fetilinia (1 sp.), Orthochiroides (4 spp.) and Orthochirus (43 spp.) (140–141).
Figures 142–165: Ventral profiles of metasoma and telson of male Orthochiroides and Orthochirus. **Figures 142–146.** Orthochiroides insularis (142), O. socotrensis (143), O. vachoni (143–145), O. somalilandus (146), O. socotrensis (143). **Figures 147–165.** Orthochirus afar (147), O. fomichevi (148), O. formozovi (149), O. gantenbeini (150), O. hormozganensis (151), O. iranus (152), O. iragus (153), O. kermanensis (154), O. kryzhanovskiy (155), O. kucerai (156), O. melanurus (157), O. mesopotamicus (158), O. nordmanni (159), O. olivaceus (160), O. persa (161), O. semnanensis (162), O. stockwelli (163), O. vignolii (164), O. zagrosensis (165). Colored arrows: character 29, antero-lateral corners of metasoma IV (green = flush, 0; blue = wedge-shaped, 1); character 32, widening of metasoma III near posterior margin (orange = graded, 0; red = abrupt, 1). Figs. 147–165 after: Kovařík et al., 2016, 2019b, 2020a, 2020b; Kovařík & Navidpour, 2020; Navidpour et al., 2019.
Figures 205–210: Phylogenetic analysis of Orthochiroides, and Orthochirus and related buthid genera. Figures 205–209. Examples of most parsimonious trees ($T_0$, $T_1$, $T_2$, $T_3$, and $T_4$) retrieved by cladistic analyses of 43 morphological characters under equal and implied weights. Numbers above indicate jackknife supports (36% resampling probability), those below relative Bremer supports, computed for implied weights (IW), concavity constant $k = 1$ (207), $k = 10$ (205), $k = 6$ (206), $k = 1$ (208) and $k = 3$ (209). Gray panels 1, 2 and 3: clades of interest. Figure 210. Phylogenetic trees of a subset of the exemplar species in Figs. 205–209, reconstructed by Bayesian inference (BEAST) and maximum likelihood (ML) analyses of multilocus sequence data (Štundlová et al., 2022) (node supports indicated).
Principal components analysis reduced the dimensions to 4 variables (PC1–PC4) explaining a cumulative 94.36% of the variance (eigenvalues: 3.7509, 1.4463, 0.9487, 0.4590; respective variances: 53.58%, 20.66%, 13.55%, 6.56%). On the basis of >50% loading of variables on principal components, we interpreted PC1 as related positively to aculeus length, height and thickness, and negatively to vesicle length, PC2 negatively to vesicle anterior length, and PC3 positively to vesicle depth, and negatively to subaculear tubercle size (Table 2). The distribution of scores for the entire order is shown in bivariate scatter plots of PC1 vs. PC2 (Fig. 134) and PC3 vs. PC4 (Fig. 135). In Figs. 136–139, scores for 9 selected genera are mapped in these 2D subspaces of the 4D reduced morphospace. It is evident that each genus is distributed within a specific domain of morphospace, corresponding to its range of expressed telson shapes. Domains of different genera may be broadly overlapped, reflecting similarities of shapes among the various genera. Some genera are spread over larger domains, while others are restricted to smaller domains. The former may indeed be characterized as having “strong variations in the shape of the telson”. For example, the speciose New World buthid genus Tityus is widely distributed in telson morphospace, although it has been proposed to be divided into subgenera (Lourenço, 2006) and it remains to be seen if future studies will justify splitting it into more than one genus.

The diagnosis of Orthochiroiodes as a genus distinct from Orthochirus is, however, based not merely on a difference in telson shapes, but specifically on the distribution of their shapes in morphospace. Figs. 140–141 show the distributions of principal components scores of these two genera. If Orthochiroiodes were a synonym of Orthochirus, then this single genus would exhibit a split distribution of two widely separated clusters in PC1-PC2 subspace (Fig. 140, red vs. green
Table 2. Factor loadings of principal components extracted from seven biometric variables quantifying telson shape. See Figs. 134–135 for variable abbreviations.

<table>
<thead>
<tr>
<th>Taxon</th>
<th>PC1</th>
<th>PC2</th>
<th>PC3</th>
<th>PC4</th>
<th>PC5</th>
<th>PC6</th>
<th>PC7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compobuthus eritreensis</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Orthochiroides vachoni</td>
<td>0.110</td>
<td>0.020</td>
<td>0.001</td>
<td>0.023</td>
<td>0.027</td>
<td>0.010</td>
<td>0.010</td>
</tr>
<tr>
<td>Orthochiroides somalilandus</td>
<td>0.110</td>
<td>0.020</td>
<td>0.001</td>
<td>0.011</td>
<td>0.000</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Orthochiroides socotrensis</td>
<td>0.110</td>
<td>0.020</td>
<td>0.011</td>
<td>0.121</td>
<td>0.100</td>
<td>0.000</td>
<td>0.220</td>
</tr>
<tr>
<td>Orthochiroides insularius</td>
<td>0.110</td>
<td>0.020</td>
<td>0.011</td>
<td>0.120</td>
<td>0.100</td>
<td>0.100</td>
<td>0.220</td>
</tr>
<tr>
<td>Neobuthus eritreensis</td>
<td>0.010</td>
<td>0.121</td>
<td>0.100</td>
<td>0.220</td>
<td>0.110</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Neobuthus kutscheri</td>
<td>0.010</td>
<td>0.121</td>
<td>0.100</td>
<td>0.220</td>
<td>0.110</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Gint banfasae</td>
<td>0.000</td>
<td>0.100</td>
<td>0.001</td>
<td>0.203</td>
<td>0.111</td>
<td>0.011</td>
<td>0.330</td>
</tr>
<tr>
<td>Gint maidensis</td>
<td>0.000</td>
<td>0.100</td>
<td>0.101</td>
<td>0.200</td>
<td>0.111</td>
<td>0.011</td>
<td>0.330</td>
</tr>
<tr>
<td>Somalibuthus sabae</td>
<td>0.000</td>
<td>0.100</td>
<td>0.112</td>
<td>0.230</td>
<td>0.111</td>
<td>0.011</td>
<td>0.330</td>
</tr>
<tr>
<td>Xenobuthus anthracinus</td>
<td>0.110</td>
<td>1.110</td>
<td>0.111</td>
<td>2.130</td>
<td>1.110</td>
<td>0.110</td>
<td>0.330</td>
</tr>
<tr>
<td>Xenobuthus xanthus</td>
<td>0.110</td>
<td>1.110</td>
<td>0.111</td>
<td>2.130</td>
<td>1.110</td>
<td>0.110</td>
<td>0.330</td>
</tr>
<tr>
<td>Butheolus gallagheri</td>
<td>0.111</td>
<td>0.100</td>
<td>1.111</td>
<td>2.110</td>
<td>1.100</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Butheolus harrisoni</td>
<td>0.111</td>
<td>0.100</td>
<td>0.001</td>
<td>2.102</td>
<td>1.100</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Butheolus thalassinus</td>
<td>0.110</td>
<td>1.100</td>
<td>0.001</td>
<td>1.170</td>
<td>1.100</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Baloorthochirus becvari</td>
<td>0.110</td>
<td>1.100</td>
<td>0.001</td>
<td>2.070</td>
<td>1.110</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Feltinia dentator</td>
<td>0.110</td>
<td>0.010</td>
<td>0.111</td>
<td>2.170</td>
<td>1.110</td>
<td>1.001</td>
<td>0.000</td>
</tr>
<tr>
<td>Orthochirus glabrifrons</td>
<td>0.110</td>
<td>0.100</td>
<td>0.011</td>
<td>2.020</td>
<td>1.110</td>
<td>0.000</td>
<td>0.221</td>
</tr>
<tr>
<td>Orthochirus innesi</td>
<td>0.110</td>
<td>0.100</td>
<td>0.111</td>
<td>2.120</td>
<td>1.110</td>
<td>0.000</td>
<td>0.221</td>
</tr>
<tr>
<td>Orthochirus afar</td>
<td>0.110</td>
<td>0.100</td>
<td>0.111</td>
<td>2.120</td>
<td>1.110</td>
<td>1.000</td>
<td>2.221</td>
</tr>
</tbody>
</table>

Table 3. Character matrix for phylogenetic analysis of Orthochiroides and exemplar species of Orthochirus and similar buthid genera. Numbered characters as defined in the Table 3.

Symbols). The disjunct distribution mirrors that of Figs. 132–133. On the other hand, the wide variations in telson shapes of other genera (Figs. 136–139) were not clearly segregated into discrete clusters, but were more or less continuously dispersed with intermediate forms linking the varied shapes. This was not the case for Orthochiroides and Orthochirus, which lack any intermediate forms between the bulbous shape of the former, and the slender shape of the latter. We mapped morphospace distributions of 97 other scorpion genera, each represented by samples of more than 4 data points, and found none with a similarly conspicuous separation into two or more clusters. A potential clustering into two groups was only seen in Uroplectes Peters, 1861, a genus that has not been revised recently and whose monophyly is untested. A small minority of species with telson shapes that are highly divergent from those of their congeners seem to be isolated cases that have evolved specialized autapomorphies, e.g., Urodacus mckenziei Volschenk et al, 2000, U. megamastigus L. E. Koch, 1977, and Chaerilus pictus (Pocock, 1890). Synonymy of Orthochiroides with Orthochirus would manufacture a genus with two highly divergent telson shapes, a property that is absent in almost all other currently recognized scorpion genera. Maintaining them as distinct genera complies with the prevalent taxonomic convention.

(iii) Metasoma III–IV. anterior corners: Orthochiroides: corners flush with anterior margin, or almost so, not projecting forward and outward; Orthochirus: corners wedge-shaped, projecting forward and outward ($\hat{3}$) (characters 29, states 0 and 1).

In ventral profile, the anterior corners of metasoma III–IV in Orthochiroides are level with, or extend only slightly
forward from the antero-ventral margin, and not outward (Figs. 142–146, 166–167; green arrows, character state 0). In contrast, the anterior corners of metasoma III–IV in Orthochirus extend distinctly forward and outward from the antero-ventral margin as wedge-shaped processes (Figs. 147–150, 166–169; red arrows, character state 1).

(iv) Metasoma II–III, posterior width: Orthochiroides: widening gradually from posterior margin; Orthochirus: widening abruptly from posterior margin (character 33, states 0 and 1).

In ventral profiles, the posterior ends of metasoma II–III in Orthochiroides widen gradually with increasing forward distance from the posterior margin (Figs. 142–146, 166–167; orange arrows, character state 0). In contrast, the posterior ends of metasoma II–III in Orthochirus are abruptly widened. A narrower annulus at the posterior margin connects to a widely flared section in front of it (Figs. 147–150, 168–170; red arrows, character state 1). This leaves a gap or recess at each posterior corner to accommodate the anterior corner wedge of the next segment. The different, more specialized construction in Orthochirus allows consecutive segments to be more tightly interlocked when the metasoma is folded or coiled.


(vi) Sternites III–VII, posterior margins: Orthochiroides: coarsely denticulate, armed with fringes of non-contiguous, enlarged blunt denticles; Orthochirus: finely microdenticulate or smooth (character 19, states 1, 2 and 0).

The sternite posterior margins in Orthochirus are similar to those of most other buthids, in being either smooth or bearing very fine, triangular microdenticles that are densely spaced, separated from each other by less than their widths. The sternite marginal denticles in Orthochiroides are much

**Table 4.** Statistics of most parsimonious trees (MPTs) retrieved by cladistic analysis of the character matrix in Table 3, rooted by outgroup taxon *Compsobuthus eritreaensis*. **Weights:** different weighting schemes tested; **EW/ IW:** equal weights/ implied weights, all characters reversible; **EW*/ IW*: equal weights/ implied weights, characters 1 & 2 irreversible; **k:** concavity constant for implied weighting; **N_MPT:** number of MPTs; **MPTs:** list of MPTs retrieved under each weighting scheme: **T**, i = 0–9; **Steps:** tree length in total number of character transformation steps; **AH:** adjusted homoplasy (for implied weights); **CI:** tree consistency index; **RI:** tree retention index; **J:** average jackknife support from 4,000 pseudoreplicates, 36% resampling probability; **Ji, i = 1–3:** node support for clades 1–3 indicated by the shaded panels in Figs. 205–209, 211; **J:** ((Orthochiroides insularis, O. sociotatus), (O. somalilandus, O. vachonii)); **J2:** (Baloorthochirus becvari, (Fetilinia dentator, (Orthochirus glabripennis, (O. afra, O. innesi)))); **J3:** (Fetilinia dentator, (Orthochirus glabripennis, (O. afra, O. innesi)))). Gray cells indicate **J1, J2 or J3 support > 50%**, blank cells indicate clade for **J1** was not recovered.
larger, blunt, clavate or digitate in shape, and are well separated from each other by more than their widths. Similar enlarged marginal denticles are also found on the sternites of Butheolus and Xenobuthus, and serve as a diagnostic character.

In Butheolus and Xenobuthus, the enlarged denticles are sexually dimorphic, with males bearing larger denticles than females. In Orthochiroideos, Orthochirus and Neobuthus, these carinae on the manus are well developed (character 13, states 0 and 1).

(vii) **Pedipalp chela manus, carinae:** Orthochiroideos: carinae V1, VA, D3 strong and costate, E either strong and costate, or weak; Orthochirus: carinae V1, VA, D3 weak, E obsolete (character 14, states 0 and 1; character 15, states 0/1 and 2) (cf. Kovářík, 1998: 116: Orthochiroideos, “six pronounced keels on the tibia of pedipalps”).

The ventroexternal (V1), ventral accessory (VA) and dorsal secondary (D3) carinae are strongly developed in all four species of Orthochiroideos (Figs. 19–21, 28–30, 64–66, 87–88, 91–92, 95–96, 99–100, 103–104, 107–108). The external secondary carina (E) is strongly developed in Orthochiroideos vachoni and O. somalilandus sp. n. (Figs. 19–21, 28–30, 64–66, 103–104, 107–108), and weakly developed in O. insularis and O. socotrensis (Figs. 87–89, 91–92, 95–96, 99–100). In Orthochirus, these carinae on the manus are attenuated, either reduced to weak, smooth carinae (V1, VA, D3) or obsolete (E).

<table>
<thead>
<tr>
<th>Character</th>
<th>Steps</th>
<th>CI</th>
<th>RI</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 Base color: lighter, yellow to light brown (0); darker, dark brown to black (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.714</td>
</tr>
<tr>
<td>1 Carapace, antero-submedian carinae: strong, granulate (0); reduced or obsolete (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.867</td>
</tr>
<tr>
<td>2 Carapace, central median carinae: strong (0); reduced or obsolete (1).</td>
<td>4</td>
<td>0.250</td>
<td>0.769</td>
</tr>
<tr>
<td>3 Carapace, supraocular carinae, : granulate (0); smooth (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.714</td>
</tr>
<tr>
<td>4 Carapace granulation density: dense (0); sparse (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>5 Carapace, granulation size: fine (0); coarse (1).</td>
<td>4</td>
<td>0.250</td>
<td>0.571</td>
</tr>
<tr>
<td>6 Carapace shape: subrectangular (0); moderately trapezoidal (1); strongly trapezoidal (2).</td>
<td>3</td>
<td>0.667</td>
<td>0.875</td>
</tr>
<tr>
<td>7 Cheliceral fixed finger, ventral accessory denticles: 2 (0); 1 (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.500</td>
</tr>
<tr>
<td>8 Pedipalp femur, petite 'trichobothrium' : present (0); absent (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.600</td>
</tr>
<tr>
<td>9 Pedipalp femur, trichobothrium e position: proximal (0), submedian (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>10 Pedipalp patella, dorsomedian carina, : complete (0); incomplete (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.500</td>
</tr>
<tr>
<td>11 Pedipalp patella, dorsomedian carina, : granulate (0); smooth (1).</td>
<td>4</td>
<td>0.250</td>
<td>0.625</td>
</tr>
<tr>
<td>12 Pedipalp chela manus, dorsoexternal carina, : granulate (0); smooth (1); absent (2).</td>
<td>5</td>
<td>0.400</td>
<td>0.625</td>
</tr>
<tr>
<td>13 Pedipalp chela manus, trichobothrium F2 position: external (0); medial (1); internal (2).</td>
<td>2</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>14 Pedipalp chela manus V1, VA, D3 carinae: strong (0); weak or obsolete (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.800</td>
</tr>
<tr>
<td>15 Pedipalp chela manus E carina: strong (0); weak (1); obsolete (2).</td>
<td>3</td>
<td>0.667</td>
<td>0.750</td>
</tr>
<tr>
<td>16 Pedipalp chela fixed finger 'trichobothrium' dt vs. et: distal (0); level (1); proximal (2).</td>
<td>7</td>
<td>0.286</td>
<td>0.500</td>
</tr>
<tr>
<td>17 Hemispermatophore basal lobe: broad hook (0); narrow hook (1); laminate hook (2); scoop (3).</td>
<td>4</td>
<td>0.750</td>
<td>0.833</td>
</tr>
<tr>
<td>18 Tergite V-VI carination: tricarinate (0); obsolete (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.333</td>
</tr>
<tr>
<td>19 Sternite V antero-medial fine granulation, : present (0); absent (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.333</td>
</tr>
<tr>
<td>20 Sternite VI submedian carinae: distinct (0); obsolete (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>21 Sternite VI lateral carinae, : distinct (0); obsolete (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.857</td>
</tr>
<tr>
<td>22 Sternite VI antero-medial surface texture: matte (0); glossy (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.800</td>
</tr>
<tr>
<td>23 Sternite VII submedian carinae: present (0); absent (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>24 Sternite VII lateral carinae: present (0); absent (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.000</td>
</tr>
<tr>
<td>25 Metasoma I dorsal granulation, : present (0); absent (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.500</td>
</tr>
<tr>
<td>26 Metasoma I ventromedial surface: granulate (0); smooth (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.750</td>
</tr>
<tr>
<td>27 Metasoma I ventrous submedian carinae: strong (0); weak/obsolete (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>28 Metasoma III-IV, anterior corners: flush (0); wedge-shaped (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>29 Metasoma III-IV, anterior corners: flush (0); reduced (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>30 Metasoma II, lateral surface: granulate (0); rugose-reticulate (1); punctate (2); smooth (3).</td>
<td>4</td>
<td>0.500</td>
<td>0.667</td>
</tr>
<tr>
<td>31 Metasoma II, lateral surface: granulate (0); rugose-reticulate (1); punctate (2); smooth (3).</td>
<td>5</td>
<td>0.600</td>
<td>0.714</td>
</tr>
<tr>
<td>32 Metasoma III, widening from posterior margin: continuous (0); abruptly stepped (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>33 Metasoma III, widening from posterior margin: continuous (0); abruptly stepped (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>34 Metasoma V ventrolateral carinae: granulate (0); lobate (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.500</td>
</tr>
<tr>
<td>35 Metasoma V dorsosubmedian carinae: granulate (0); smooth (1); obsolete (2).</td>
<td>6</td>
<td>0.333</td>
<td>0.500</td>
</tr>
<tr>
<td>36 Metasoma V dorsal carina: width increase: &lt; 10% (0); &gt; 10% (1).</td>
<td>1</td>
<td>1.000</td>
<td>1.000</td>
</tr>
<tr>
<td>37 Metasomal macrosetae, : length: : &gt; (0); &lt; (1).</td>
<td>3</td>
<td>0.333</td>
<td>0.500</td>
</tr>
<tr>
<td>38 Telson vesicle ventral surface, punctation: absent (0); weak (1); strong (2).</td>
<td>4</td>
<td>0.500</td>
<td>0.714</td>
</tr>
<tr>
<td>39 Telson vesicle ventral surface, granulation: moderate to strong (0); weak or absent (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.667</td>
</tr>
<tr>
<td>40 Telson vesicle lateral profile, : hemi-elliptic (0); bulbous (1); slender, pyriform (2).</td>
<td>4</td>
<td>0.500</td>
<td>0.800</td>
</tr>
<tr>
<td>41 Telson vesicle posterior slope: moderate (0); steep (1); shallow (2).</td>
<td>3</td>
<td>0.667</td>
<td>0.833</td>
</tr>
<tr>
<td>42 Telson vesicle W/ metasoma V W: &gt; 0.5 (0); &lt; 0.5 (1).</td>
<td>2</td>
<td>0.500</td>
<td>0.667</td>
</tr>
</tbody>
</table>

Table 5. Lengths (Steps), consistency indices (CI) and retention indices (RI) of individual characters mapped to the most parsimonious tree, Tg (EW).

The proposed synonymy of Orthochiroides with Orthochirus by Lourenço & Ythier (2021: 339) equates to the hypothesis that these two genera together comprise a monophyletic group. To test this hypothesis, we conducted a cladistic analysis of an ingroup that included all species of Orthochiroides, plus selected exemplar species representing Orthochirus and other morphologically similar genera of small buthids (Neobuthus Hirst, 1911, Gint Kovařík et al, 2013, Butheolus Simon, 1882, Xenobuthus Lowe, 2018, Balooorthochirus Kovařík, 1996 and Fettlinia Lowe & Kovařík, 2021; Table 3). As outgroup taxon we selected Compsobuthus eritreensisus Kovařík, Lowe, Plíšková & Šťáhlavský, 2016. This selection was guided by results of a molecular phylogenetic study in which this species, along with other Compsobuthus species, was placed in the immediate sister clade of a larger clade including all of our DNA-sampled ingroup taxa (Štundlová et al., 2022). We analyzed a matrix of 43 discrete morphological characters (Tables 3, 5) These characters were selected on the basis of: (i) their relative intrageneric stability for ingroup taxa, with most having minor or no interspecific variation within each genus; and (ii) their systematic variation across genera of the ingroup, which was potentially informative about higher level relationships. They included 25 binary characters and 18 unordered multistate characters. Phylogenetically uninformative characters were excluded from the analyses. Two binary characters (characters 1 and 2: loss of antero-submedian and central median carinae on the carapace) were tested under assumptions of both reversibility and irreversibility. Intrageneric stability ensured that results about intergeneric relationships would not be sensitive to our representation of the larger genera (Neobuthus and Orthochirus) by a minority of their named species. Additional exemplars of these genera would have identical, or almost identical, character scores as those already included, and are therefore expected to only introduce polytomies at their respective genus nodes.

We conducted 22 analyses, assuming either equal weights, or implied weights with varying strengths of concavity ($k = 1–6, 8, 10, 15, 30$). In total, nine distinct most parsimonious trees ($T_i, i = 0–8$) were retrieved (Table 4). A monophyletic including all species of Orthochiroides, clade 1 = ((Orthochiroides insularis, O. socotrensis), (O. somalilandus, O. vachoni)), was resolved in 14/22 analyses. Clade 1 was separated from, and basal to, other ingroup taxa. It received significant support in 9/11 analyses that assumed irreversibility of characters 1 and 2 (e.g., upper gray panels in Figs. 205–206, 209). If reversibility was permitted, the carapace carinae were initially lost for the entire ingroup and later regained in Gint, Somalibuthus and Xenobuthus. Irreversibility assumes that once lost, a compound structure like a carina is less likely to be reconstructed de novo in these genera, which also possess other plesiomorphic characters, than it is to be lost independently in the more specialized lineages (clades 1, 2 and 3), which also possess other apomorphic characters. In most cases, imposing irreversibility had little effect on tree lengths, but increased support for clade 1 (Orthochiroides) which gained two synapomorphies (i.e., carinal losses).

In all of the analyses and tree topologies, either clade 2 = (Balooorthochirus becvari, (Fettlinia dentator, (Orthochiroides glabrifrons, (O. afar, O. ‘innesi’))), or clade 3 = (Fettlinia dentator, (Orthochiroides glabrifrons, (O. afar, O. ‘innesi’))), or both, were resolved with significant support (jackknife statistic 50% – 71%) (Table 4, gray cells). This result was independent of reversibility of characters 1 and 2. Clades 2 and 3 excluded all species of Orthochiroides, and included Orthochirus and either one or both of two other genera, Balooorthochirus and Fettlinia. This implies that a group including only Orthochiroides and Orthochirus is polyphyletic, thereby refuting the proposed synonymy of these two genera. A genus including both Orthochiroides and Orthochirus would also need to include Balooorthochirus and Fettlinia, which have very different metasomal morphologies. Even then, it would require fusion of clades 1 and 2, which were not resolved as sister groups in any of our analyses. A monophyletic genus that includes both Orthochirus and Orthochirodes necessitates the inclusion the entire set of ingroup taxa (Figs. 205–209, 211).

A subset of our ingroup taxa was included in an extensive DNA analysis of buthid phylogeny (Štundlová et al., 2022). The relationships among these taxa reconstructed by Bayesian inference and maximum likelihood analysis applied to the entire buthid dataset are shown in Fig. 210. The species representing Orthochiroides occupied a basal position, separated from the lineages containing Orthochirus (shaded panels in Fig. 210). This separation, as well as the overall topology, were consistent our morphological analyses. Taken together, these findings further confirm the status of Orthochiroides as a genus separate from Orthochirus, as originally proposed by Kovařík (1998, 2004). The synonymy of Orthochiroides with Orthochirus by Lourenço & Ythier (2021) is hereby invalidated, and we reinstate Orthochirodes Kovařík, 1998 stat. rev., with a differential diagnosis provided by the seven characters (i) – (vii).

Off the nine different trees obtained in our analyses, the most frequent was $T_6$ (e.g., Fig. 205). It was retrieved repeatedly under equal weights and under implied weights with moderate to weak concavity constants. It also enjoyed the highest support values for clades 1, 2 and 3, and was our preferred phylogenetic hypothesis. Unambiguous synapomorphies for $T_6$ are mapped in Fig. 211.

Discussion

At first glance, the basal position of Orthochiroides relative to other ingroup taxa and its separation from Orthochirus revealed by cladistic analysis, may not be obvious. The general habitus, small size, dark coloration, trapezoidal carapace, abbreviated pedipalps, and stout metasomal segments with heavy sclerotization and specialized morphosculpture (pins or
depressions) on more posterior segments, all seem to suggest a closer relationship with *Orthochirus*. However, the weight of evidence from a larger set of characters analyzed collectively, as well as from wholly independent DNA analyses, imply that these similarities are superficial and are likely to have arisen by convergence. Small size and dark coloration are characters found in many other genera and families of scorpions. A trapezoidal carapace and abbreviated pedipalps are characters shared with most of the other ingroup genera of small buthids (*Neobuthus, Xenobuthus, Butheolus, Balooorthochirus* and *Fetilinia*). Moreover, the carapaces of *Orthochiroidea* and *Orthochirus* occupy disjunct geometric domains in trapezoidal morphospace. The heavy sclerotization and specialized morphosculpture of metasomal segments in *Orthochiroidea* and *Orthochirus* are not shared with other ingroup members, and could be synapomorphies relating the two genera. However, heavy sclerotization and punctate sculpture of the posterior metasoma, similar to that of *Orthochirus* spp., *Orthochiroidea insularis* and *Orthochiroidea socotrensis*, also occurs in the ‘*Buthus*’ group genus *Microbuthus* Kraepelin, 1898. Other characters, including uniquely modified pedipalps, imply that *Microbuthus* is more closely related to *Picobuthus* Lowe, 2010, a genus without punctate metasomal segments (Lowe, 2010). Indeed, DNA analysis placed *Microbuthus* and *Picobuthus* in a lineage separate from *Orthochiroidea*, *Orthochirus* and the other ingroup taxa analyzed here (Štundlová et al., 2022). Other buthid genera with punctate posterior metasomal segments include *Butheoloides* Hirst, 1925, *Isometroides* Keyserling, 1885, *Karasbergia* Hewitt, 1913, and some species of *Uroleutes* (Hirst, 1925; Lourenço, 2001; Prendini, 2004, 2015). These all reside in clades external to the ‘*Buthus*’ group (Fet et al., 2005; Štundlová et al., 2022). The punctate metasoma admits a functional interpretation that can explain why it may be favored by parallel evolution and susceptible to homoplasy (Lowe, 2010). In buthids with weak pedipalps, the posterior metasoma combined with the telson function as the main offensive and defensive weapon, and is expected to be under selection pressure to develop thicker sclerotization. It may also have sensory functions (Fet et al., 2003), requiring deployment of sensillae over its cuticular surface. The punctae or pits simply represent foci where the exoskeleton is thinned to facilitate the connections of internal sensory neurons to external sensillae.

A character that we did not include in our diagnosis of *Orthochiroidea* is the shape of the hemispermatophore basal lobe. Basal lobe shapes are potential diagnostic characters because they appear to be more or less conserved within many ‘*Buthus*’ group genera that have been studied (e.g., Kovařík et al., 2018). In *Orthochiroidea somalilandus* sp. n., the basal lobe is a broad hook with a triangular profile, projecting strongly out from the convex surface of the capsule (Figs. 46–48, 193–196). In several species of *Orthochirus* that we examined, the basal lobe is a ‘laminate’ hook that is more elongated with a lower profile (Figs. 197–204). Similar shapes of the basal lobe in *Orthochirus* have been recorded for a few species in the literature (Levy & Amitai, 1980: 99, fig. 90; Vachon, 1952: 227, fig. 309). The hemispermatophore has not been described for other species of *Orthochiroidea* and *Orthochirus*, and more studies are needed to determine if these differences can be generalized as diagnostic characters. In our data matrix, hook shapes of the other three species of *Orthochiroidea*, and of *Balooorthochirus* and *Fetilinia* were scored as unknowns (?) and the character did not contribute much to the analysis. However, we note that a broad hook is also present in the outgroup genus *Compsobuthus* (Fig. 189). In *Neobuthus* and *Gint*, the basal lobe is also broad but the apical hook is less sharp (Figs. 190–192). *Orthochiroidea* is more similar to these other genera in the form of its basal lobe, than to *Orthochirus*.

Under our preferred hypothesis, *T*<sub>0</sub>, *Orthochiroidea* is monophyletic, and its sister node is supported by six unique synapomorphies (Fig. 211). These involve coarsening of carapace granulation, attenuation and loss of carinae on the pedipalp and sternite VI, and development of glossy cuticle on sternite VI. At the next level of the tree, the branch leading to *Orthochirus* is supported by two synapomorphies (one unique). These both involve the carapace, i.e., a moderately trapezoidal shape, and loss of antero-submedian carinae. At the next bifurcation, clade 2 = (*Balooorthochirus*, *Fetilinia*, *Orthochirus*) diverges from its sister group containing *Butheolus* and *Xenobuthus*. Clade 2 is supported by six additional synapomorphies (3 unique). These involve loss of centromedian carinae on the carapace, loss of petite ‘trichobothrium’ d, on the dorsal femur, stepped widening of metasoma II–III posterior margins, and a telson with slender pyriform vesicle and shallow posterior slope. The metasoma and telson characters are unique to clade 2. Within clade 2, clade 3 contains *Fetilinia* and *Orthochirus* and is supported by four additional synapomorphies (one unique). These involve reduction of granule size on the carapace (a reversion from coarse to fine granulation), smoothening of pedipalp patella carinae, and the forward and outward extension of the anterior corners of metasoma III–IV. The latter character is unique, more weakly developed in *Fetilinia* (a partial or intermediate condition), and more strongly so in *Orthochirus*. We postulate an evolutionary sequence in which stepped widening of metasoma II–III posterior margins developed first in the ancestor of *Balooorthochirus*, followed by addition of a weak metasoma III–IV anterior corner flaring in the ancestor of *Fetilinia*. Finally, stronger anterior corner flaring in the ancestor of *Orthochirus* enabled a closer interlocking of successive metasomal segments, which is a unique structural adaptation of the genus which never evolved in *Orthochiroidea*. The separation of clade 2 from *Orthochiroidea* is supported by a total of 13 synapomorphies (10 unique), and the separation of clade 3 by a total of 17 synapomorphies (11 unique).

Under our preferred hypothesis, *T*<sub>0</sub>, the pair of species of *Orthochiroidea* endemic to Socotra Island (*O. insularis* and *O. socotrensis*) form a sister group with respect to the group composed of the pair of species found on the Horn of Africa mainland (*O. somalilandus* sp. n. and *O. vachoni*) (Fig. 211). The two sister groups are united under clade 1 by five
non-unique synapomorphies. These involve dark coloration, reduction of carinae on the carapace, development of a fringe of enlarged, sexually non-dimorphic denticles on the posterior margins of sternites III–V, and a bulbous telson. However, compared to the mainland species, the two Socotra species are more derived with five synapomorphies. These involve smoothening of pedipalp patella carinae, obsolence of pedipalp fixed finger trichobothrium dt with respect to et, and a punctate metasoma II lateral surface. The punctate surface morphosculture of the Socotra species extends over lateral and ventral metasoma III–V (Figs. 82–83), surfaces that are rugose-reticulate in the two mainland species (Figs. 13–14, 16–17, 84–85). The rugose-reticulate condition can be viewed as an intermediate stage of enhanced sclerotization of metasomal segments, in which the network of thickened wrinkles or ridges occupies less total surface area than the lacunae between them (Kovařík, 1998). In the Socotra species, the thickened exoskeleton has expanded to occupy a greater total surface area than the lacunae, which have contracted into smaller punctae or pits. Some of the most parsimonious trees retrieved under implied weights with strong concavity placed smaller punctae or pits. Some of the most parsimonious trees retrieved under implied weights with strong concavity placed

Comparative material examined

_Baloothorhocirus becvari_ Kovařík, 1996

**Pakistan**: SE Balochistan, Khurkha, 38 km S of Uthal, 24.1V. 1993, 1♂ (holotype) (FKCP), leg. S. Bečvář.

_Buthothes gallagheri_ Vachon, 1980

**Oman**: main road above Khor Rori Beach, UV detection on ground, densely vegetated wadi, E of Taqah; warm and humid with many insects, 17°03.22′N 54°25.33′E, 50 m a. s. l., 18.X.1993, 21:24 h, 4♂11♀ (GLPC, NHMB, ONHM), leg. G. Lowe; Mirbat, 17°02.19′N 54°38.75′E, 54 m a. s. l., 4♂11♀ (FKCP).

_Buthothes harrisoni_ Lowe, 2018

**Oman**: Jabal Qara; north slopes, Nejd, UV detection, rocky wadi & rocky slopes, 17°17.83′N 54°05.11′E, 800 m a. s. l., 16.X.1993, 22:38 h, 4♂7♀ (NHMB, USNM), leg. G. Lowe.

_Buthothes thalassinus_ Simon, 1882


_Compsobuthus eritreaensis_ Kovařík, Lowe, Plíšková & Šťáhlavský, 2016

**Eritrea**: near Massawa, 15°36′58.7″N 39°22′32.8″E, 74 m a. s. l., 4.-5.XI.2015 (Locality 15EI), 7♂12♀ (holotype and paratypes) (FKCP), 6♂ hemispermatophore No. 877 (GLPC), leg. F. Kovařík.

_Fetilina dentator_ Lowe & Kovařík, 2021

**Pakistan**: Khyber Pakhtunkhwa (formerly North-Western Frontier) Province, Karak, 33.102°N 71.049°E, 23.VI.2010, 1♂ subadult (holotype) 1♀ juvenile (paratype) (FKCP), leg. Z. Ahmed.

_Gint amoudensis_ Kovařík, Lowe, Just, Awale, Elmi & Šťáhlavský, 2018

_Somalliland_: Borama, Amoud University campus, 09°56′49″N 43°13′23″E, 1394 m a. s. l. (Locality No. 17SR =17SA), 9-13. IX.2017, 1♂ (holotype), 6♂7♀ juvs.2♂ juvs. (FKCP), leg. F. Kovařík.

_Gint banfasae_ Kovařík & Lowe, 2019

_Somalliland_: Shanshade vill., 08°39′35″N 45°55′49″E, 790 m a. s. l. (Locality No. 18SJ), 29–31.VIII.2018, 66♂8♀4juvs. (holotype and paratypes), Nos. 1530, 1531, 1532, 1533, 1534) (FKCP), 2♂ (GLPC), leg. F. Kovařík et al.

_Gint maidensis_ Kovařík, Lowe, Just, Awale, Elmi & Šťáhlavský, 2018

_Somalliland_: Maid, 11°00′03″N 47°06′30″E, 52 m a. s. l. (Locality No. 17SN), 3-4.IX.2017, 11♂9♀ im.♂1im.♀ (holotype, paratypes, Nos. 1321, 1324, 1336) (FKCP), hemispermatophores Nos. 1321, 1324, 1336 (GLPC), leg. F. Kovařík.

_Neobuthus eritreaensis_ Kovařík & Lowe, 2016

**Eritrea**: near Massawa, 15°36′58.7″N 39°22′32.8″E, 74 m a. s. l., 4.XI.2015 (Locality No. 15EI), 1♂ (holotype) 4♂1♀ im.♂1im.♀ (paratypes) (FKCP), 1♂1♀ (paratypes), 6♂5♀ juvs. (paratypes) (FKCP), 2♂3♀3♂✿ juvenile (paratypes) (GLPC), leg. F. Kovařík et al.
Orthochirus afar Kovářík & Lowe, 2016

**Ethiopia**: Afar Region, Gewane, 10°09’38”N 40°39’45”E, 631 m a. s. l. (Locality No. 12EO), 23.-24.XI.2012, 1♂ holotype, 1♀ paratypes, 1♂juv. paratype (FKCP), leg. F. Kovářík; 11°43’22”N 40°56’52”E, 457 m a. s. l. (Locality No. 12EM), 20.XI.2012, 1♂juv. (paratype) (FKCP), leg. F. Kovářík.

**Somaliland**: Gerissa, N of Borama, 10°36’01”N 43°26’07”E, 245 m a. s. l. (Locality No. 17ST), 11.-12.IX.2017 (FKCP), ♂ hemispermatophores No. 1301, 1328 (GLPC), leg. F. Kovářík.

Orthochirus glabrifrons (Kraepelin, 1903)


Orthochirus cf. ‘grlabifrons’ (Kraepelin, 1903)

**Oman**: W of Wadi Andhur, 17.6877488°N 54.5917380°E, X.2019, 1♂1♀, ♂ hemispermatophore No. 1389 (GLPC), leg. M. Stockmann; Kawdaki, ♂ hemispermatophore No. 1219 (GLPC).

Orthochirus cf. ‘innesi’ Simon, 1910


Orthochirus scrobiculosus (Grube, 1873)

**Turkmken**: Balkan Province, Krasnovodsk [now Türkmenbashı, 40.02°N 52.97°E], ♂ lectotype, leg. Dr. [Gustav] Radde, [June 1870], MNHW No. 531.

Orthochirus sp.


Somalithus sabae Kovářík & Njoroge, 2021


**Somalithus xanthus** Lowe, 2018


**Acknowledgements**

Thanks are due to V. Bejiček, Jan Farkač, David Král, and Karel Šťastný who collected scorpions on Socotra Island. Thanks are due to Petra Frydlová, Daniel Frynta, David Král, Petr Kabátek and David Sommer (Czech Republic), Abdiqadir Abdilahi and Abdisalaan Shabele (Republic of Somaliland) who participated and helped in the expeditions to Somaliland. Thanks to Mohamud Yousuf Muse (President of University of Hargeisa), Mohamed A. Sulub (Director, Corporate Communication Directorate, University of Hargeisa), Sulieman Ahmed Gulair (President of Amound University), Ahmed A. Boqore (Vice President, Academic Affairs of Amound University), Shukuri Haji Ismail and Abdinasir Hussein (Ministry of Environment & Rurar Development, Hargeysa, Republic of Somaliland), and inhabitants of Mader Mage and Maid villages (Somaliland) for their help. Further, we thank two anonymous reviewers for their comments on the manuscript.

**References**


