A new species of Mouse Spider (Actinopodidae, *Missulena*) from the Goldfields region of Western Australia

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http://zoobank.org/6FB6F2EA-4F6F-44ED-A644-DB99C9411E36
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Abstract

A new species of Mouse Spider (family Actinopodidae Simon, 1892), *Missulena harewoodi*, is described from near the City of Kalgoorlie-Boulder in the Goldfields region of Western Australia. It differs from all other *Missulena* species by the unusual light grey colouration of the abdomen in combination with small body size and shiny carapace. A phylogenetic analysis of a fragment (658 bp) of the COI barcoding gene places *M. harewoodi* sp. n. in a clade with four *Missulena* species from the Pilbara region of Western Australia, more than 900 km away. *Missulena harewoodi* sp. n. is one of the many species in this genus that are currently only known from a single, or a very limited number of specimens, highlighting the paucity of fauna collections in many arid regions of Australia and the difficulties in sampling these cryptic spiders.

Introduction

The spider family Actinopodidae Simon, 1892 has a Gondwanan distribution with species found in Australia and South America (World Spider Catalog 2017). In Australia, these spiders are commonly called Mouse Spiders and widely known because they can be quite large (up to 3–4 cm in body size) and superficially resemble the venomous Funnel-web Spiders (family Hexathelidae Simon, 1892). Some species may have a potent venom although spiders appear to generally apply a dry bite in defense (e.g. Isbister 2004; Isbister and Gray 2004; Rash et al. 2000). The colour patterns in male *Missulena* belong to the most spectacular amongst all mygalomorph spiders. Most striking are the sometimes bright red fangs and/or carapace in males, iridescent blue abdomens, or an abdomen with light colour patterns (Framenau et al. 2014). Vivid colouration with typical warning colours such as red possibly relates to the diurnal activity of mate-searching *Missulena* males, whereas the majority of males in other mygalomorph families are strictly nocturnal (Harms and Framenau 2013).

The Australian fauna in this family currently comprises 16 species in the single genus *Missulena* Walckenaer, 1805 (Miglio et al. 2014; World Spider Catalog 2017). However, species-level diversity in this genus is clearly much higher, with dozens of undescribed species identified in recent barcoding analyses (e.g. Castalanelli et al. 2014). The fauna of Western Australia, where 14 of the 16 described species occur (Miglio et al. 2014), is particularly diverse. Recent barcoding projects focusing on the Pilbara region in the northwest of the state recovered 13 distinct genetic clades, only five representing described species (Castalanelli et al. 2014). The two most recent taxonomic studies (Harms and Framenau 2013, Miglio et al. 2014) on *Missulena* developed a protocol for the description of spe-
cies after several decades of taxonomic neglect and added six new species to the Australian fauna. They also provided an updated key to the males of all Australian species (Harms and Framenau 2013; Miglio et al. 2014).

Recent collections as part of environmental assessment studies in the semi-arid Goldfields region of Western Australia recovered a small male of *Missulena* with the feature of light grey abdomen both as live (VWF pers. obs.) and preserved (Fig. 1A) specimen, another facet of the striking colour variations present within this genus. Light abdomen discolourations have previously only been described from two larger species with very different distribution in Australia (*M. pruinosa* Levitt-Gregg, 1966 from tropical northern Australia and *M. brandleyi* Rainbow, 1914 from the subtropical/temperate eastern parts of the country), but the current species is smaller and the pattern is more pronounced.

In this paper, we describe this unusual species of *Missulena* based on male morphology. A phylogenetic analysis of a fragment of the COI barcoding gene is used to explore the systematic position of this species and consolidate its taxonomic concept developed based on morphology. In documenting this species, we hope to raise further awareness for the Australian Mouse Spiders by documenting another morphological rarity in its diverse fauna.

### Methods

**Morphology.** The holotype of *M. harewoodi* sp. no. was examined in 75% ethanol under Leica M205C and M80 stereomicroscopes. Digital images were taken using a Leica DFC 295 digital camera attached to the Leica M205C stereomicroscope controlled by the Leica Application Suite Version 3.8. The images were edited and formatted in Adobe Photoshop CC, release 2017. For measurement protocols and overall format of the description please see Griswold and Ledford (2001), Harms and Framenau (2013) and Miglio et al. (2014). Abbreviations for morphological characters are as follows: (EL) embolar lamella, (DET) distal embolar tooth, (BEI) basal embolar intumescence, (PME) posterior median eyes, (PLE) posterior lateral eyes, (ALE) anterior lateral eyes, (AME) anterior median eyes, (MOQ) median ocular quadrangle, (OAL) ocular area length, (OAW) ocular area width. Leg spination: (d) dorsal, (pv) proventral, (rv) retroventral, (v) ventral. WAM refers to the Western Australian Museum. The nomenclature of undescribed species follows the designation as provided by the Western Australian Museum (WAM) or Castalanelli et al. (2014).

**Molecular analyses.** Sequencing of the mitochondrial COI gene of the holotype followed the protocol described in a previous study (Harms and Framenau 2013). Phylogenetic analyses were undertaken on an alignment comprising all 84 *Missulena* sequences from previous studies and available on GenBank (Castalanelli et al. 2014; Harms and Framenau 2013), four specimens that were sampled in Western Australia since then and the holotype of *M. harewoodi* sp. n. (Table 1). COI sequences of a single specimen of *Euaegus chisoeus* Gertsch, 1939 (Dipluridae Simon, 1889) and five unidentified specimens of *Conothele* Thorell, 1878 (family Ctenizidae Thorell, 1887) were extracted from GenBank and included in the analyses as outgroups. A barcoding threshold of 9.5% pairwise sequence divergence was used for species delineation at a molecular level (Castalanelli et al. 2014). Phylogenetic analyses were executed using MrBayes Version 3.2.1 for Macintosh (Ronquist et al. 2012), with the GTR+1+G model of nucleotide substitution suggested by MrModeltest Version 3.7 (Posada and Crandall 1998) for an unpartitioned dataset. Four Markov chain Monte Carlo (MCMC) chains were run for 40,000,000 generations, sampling every 1,000 generations and discarding the first 25% of sampled trees as ‘burnin’. FigTree version 1.4.3 (Rambaut 2016) was used to visualise and edit the tree.

**Systematics**

**Family Actinopodidae Simon, 1892**

**Missulena Walckenaer, 1805**


**Missulena harewoodi** sp. n.

http://zoobank.org/3FE6A340-C871-45B8-ACFC-E9677586C417

Figs 1A–G, 2A–F, 4

**Type material.** **AUSTRALIA:** **Western Australia:** holotype male, 20 km East of Kalgoorlie, 30°44′41″S, 121°34′01″E, 14–16 April 2015, Greg Harewood, dry

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Table 1. *Missulena* specimen sequenced for this study.

<table>
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<th>WAM registration</th>
<th>Identification (WAM database)</th>
<th>Sex</th>
<th>Location</th>
<th>Latitude</th>
<th>Longitude</th>
<th>GenBank accession no.</th>
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[evolsyst.pensoft.net](http://evolsyst.pensoft.net)
Figure 1. *Missulaena harewoodi* sp. n., male holotype (WAM T142820) A, habitus, dorsal view; B, ventral view; C, eye region, dorsal view; D, carapace, lateral view; E, sternum, ventral view; F, abdomen, dorsal view; G, maxillae and labium, ventral view. Scale bars: A, B – 2.0; C – 0.1; D, E, F – 1.0; G – 0.5. Arrows point to pair of anterior setae (C) and labio-sternal sigillae (E) (see text).
pittal trap, Goldfields Blackbutt low woodland over open scrub on loam (WAM T142820).

**Etymology.** The specific epithet is a patronym in honour of Greg Harewood, the collector of the type specimen.

**Diagnosis.** The colouration of the holotype of *M. harewoodi* sp. n. is most similar to *M. pruinosa* due to the light dorsal discoulouration of the abdomen, but the species differs in the lower number of spines of the rastellum (three vs ten), smaller size (male body length 8.0 mm vs 12.5 mm) (measurements from Faulder 1995a) and narrower pedipalp tibia. Male *Missulena bradleyi* also have a light pattern on the dorsal side of the abdomen, but it is restricted to an anterior light blue patch and the species is also larger (male body length 8.0 mm vs 10.9 mm) (measurements from Faulder 1995a). Otherwise, somatic morphology most closely resembles three species with a brown carapace, *M. melissae* Miglio, Harms, Framenau & Harvey, 2014, *M. faulderi* Harms & Framenau, 2013 and *M. rutraspina* Faulder, 1995, but *M. harewoodi* sp. n. differs considerably in the color pattern of the abdomen of both live and preserved specimens (brown in *M. melissae*, grey-brown in *M. faulderi* and blue-grey in *M. rutraspina*) and the much smoother carapace (Faulder 1995b; Harms and Framenau 2013; Miglio et al. 2014).

**Description.** Adult male, based on holotype (WAM T142820). Small mygalomorph spider (total length 8.0).

**Colour:** Carapace glabrous brown to dark brown (Fig. 1A); dark brown around PME (Fig. 1C); chelicerae and fangs glabrous, brown to reddish-brown (Figs 1D, G); abdomen dorsally pale whitish laterally, medially light grey (Figs 1A, F); ventrally pale yellowish-brown (Fig. 1B); sternum yellowish-dark, darker towards margins, sigilla light brown (Fig. 1B, E); labium and maxillae light brown, with yellow-brown spots (Fig. 1G); legs glabrous brown, ventrally with olive tinge (Figs 1A, B); spinnerets olive gray (Fig. 2F).

**Carapace:** 3.52 long, 3.74 wide; clypeus 0.19; caput and eye region elevated (Fig. 1D); pars cephalica smooth, pars thoracica with bands of fine, radial furrows.

**Eyes:** OQ 3.28 times wider than long, OAW 2.13; OAL 0.65; width of posterior eye group 1.85; PME 0.178; PLE 0.18; ALE 0.23; AME 0.30; AME inter-distance 0.15; AME to ALE 0.83; AME to PME 0.21; PLE to ALE 0.40; PLE to PME 0.42; PME inter-distance 1.20; PME to ALE 0.49; two black setae anterior of AME (Fig. 1C).

**Chelicerae:** 1.62 long, 1.03 wide; with few short silvery setae medially; rastellum developed, slightly pronounced, consisting of a sclerotised process with 3 (left 4) strong conical spines and 12–14 disordered setae (Fig. 2E), 10–14 long setae extend forward from anterior margin of each chelicera and cover base of fang; inner margin of cheliceral furrow with 3 rows of teeth (Fig. 1G); pro-lateral (inner) row with ca. 9 teeth; intermediate row with 6 proximal, small spaced teeth; retrolateral (outer) row with 2 proximal teeth.

**Maxillae:** 1.56 long; 1.10 wide (Fig. 1G), with ca. 40 pointed cusuples along entire anterior margin.

**Labium:** ca. 0.82 long, 0.70 wide; conical, 11 pointed cusuples anteriorly (Figs 1G); labiosternal suture poorly developed; a pair of sigilla near labiosternal suture (Fig. 1E).

**Sternum:** 2.17 long, 2.00 wide; pear-shaped and rebordered (Fig. 1E), with dark setae of varying length, arranged irregularly but denser laterally and towards labium; 4 pairs of sigilla located more than three times their length from the border of the sternum, anterior and second pair (antior-posterior) smallest and poorly defined, third pair bigger than 2 anterior pairs and poorly defined; posterior pair biggest, roughly oval and well defined, 3 posterior sigilla slightly depressed.

**Abdomen:** 3.58 long, 3.23 wide; roughly oval (but collapsed through preservation) (Fig. 1A, F); 4 spinnerets (Fig. 2F), PLS 0.52 long, 0.43 wide, apical segment domed; PMS 0.35 long, 0.16 wide at base.

**Pedipalp:** Length of trochanter 0.76, femur 1.62, patella 1.08, tibia 2.12, tarsus 0.86; tibia with irregular black setae, densest ventrally (Fig. 2A–C); bulb pyriform (Fig. 2A–C), two strongly sclerotised sections connected by a velar median structure (“haematodocha”, Fig. 2C); embolus very slightly curved, reaches to half tibia length, with an intumesence in proximal region (BEI), a strong curvature in the duct in prolateral view; tapering and slightly twisted medially (Fig. 2C); embolus tip rounded triangular, with a lamella (EL) poorly developed and no prominent tooth (DET) (Fig. 2D).

**Legs:** With few brown setae, ventral setae of tibiae and metatarsi generally much longer and thicker than dorsal setae; dorsal; preening comb distal in tarsi, very small and plain; metatarsi and tarsi I and II ascopulate, metatarsi (along distal half) and tarsi (along whole length) of legs III and IV densely scouplate. **Leg measurements:** Leg I: femur 2.37, patella 1.54, tibia 1.84, metatarsus 1.46, tarsus 0.97, total 8.18. Leg II: 2.54, 1.44, 1.46, 1.59, 0.95, 7.98. Leg III: 2.68, 1.49, 1.29, 1.17, 1.28, 8.11. Leg IV: 3.14, 1.70, 2.10, 3.29, 1.16, 11.39. Formula 4123.

**Trichobothria:** Arranged in discontinuous rows; tibiae I–II with 2 rows of 3 in retrodorsal and prodorsal position, respectively; tibiae III with 1 rows of 2 in retrolaterodorsal; tibiae IV with 2 rows, the first row with 3 in retrolaterodorsal and the second row with 2 in proximolateral position; metatarsi with 3 in proximo-dorsal row, tarsi I with 2 in proximo-dorsal row, tarsi II with 3, III IV with 4 medio-dorsally, respectively.

**Leg spination:** Pedipalp aspinose; leg I: tibia r1v1–1–0, v3–3–7, pv1–2–0, d0–0–0; metatarsus r2v2–1–1, v2–3–5, pv0–0–0, d0–0–0; tarsus r1v4–4, v2–7–3, pv2–2–2, d0–0–0; leg II: tibia r0v0–0–0, v0–0–0, pv0–1–0, d0–0–0; metatarsus r0v0–0–0, v0–0–0, pv0–0–0, d0–0–0; tarsus r3–3–5, v1–2–2, pv1–3–2, d0–0–0; leg III: tibia r0v0–0–0, v0–3–2, pv2–2–2, d2–1–3; metatarsus r2v2–2–3, v0–0–0, pv3–3–4, d8–4–2; tarsus r3–3–5, v0–0–1, pv1–3–4, d0–2–2; leg IV: tibia r0v0–2–0, v2–4–4, pv1–2–2, d3–0–0; metatarsus r1v1–3–2, v0–0–0, pv1–3–4, d0–0–1; tarsus r4v9–13, v0–0–1, pv1–4–6, d0–0–2; patellae I with ca. 10 rasp prolaterally, II with 2 rasps prolaterally, III with ca. 40
rasps prolaterally to dorsal, patella IV with ca. 10 rasps prolaterally and proximo-dorsally.

**Phylogenetic analyses.** Our phylogenetic analyses places *M. harewoodi* sp. n. as sister taxon to an undescribed *Missulena* species from the Pilbara region in Western Australia, *Missulena ‘DNA02’* (WAM T124777) (Fig. 3; Table 1), although this relationship is not well supported. Sequence divergence between both specimens is 14.5%, which is considerably larger than the current operational sequence divergence of 9.5%, employed to differentiate species in the Actinopodidae (Castalanelli et al. 2014). Morphological comparison between both species is not possible as the Pilbara species is represented by a single juvenile specimen. *Missulena harewoodi* sp.
Figure 3. Topology of *Missulena* species based on COI (fragment of 658 bp). Nodal support is indicated in squares (posterior probabilities). Branches of more than three specimens of the same species are collapsed. Morphotype designations of undescribed species reflects those databased at the Western Australian Museum. See Castalanelli et al. (2013) for details on the ‘MYG’-coding system and museum registration numbers.

Distribution. *Missulena harewoodi* sp. n. is currently only known from the type locality, ca. 20 km East of the City of Kalgoorlie-Boulder in the Goldfields region of Western Australia (Fig. 4).

Habitat. The type specimen was collected alive in a 10-litre, dry bucket pitfall trap targeting vertebrates. The collecting site is described as Goldfields Blackbutt (*Eucalyptus lasiocarpa*) low woodland over open scrub on loamy soil (G. Harewood, personal communication to VWF). Similar to many other *Missulena* species, *M. harewoodi* sp. n. appears to mature in autumn (collected in April), contradicting the assumption that many mygalomorph spiders in arid and semi-arid Australia reproduce in the months with highest rainfall (e.g. January/February in the Goldfields near Kalgoorlie) (BoM 2017; EPA 2016).

Remarks. *Missulena harewoodi* sp. n. is the seventeenth named species of this genus in Australia and within a radi-
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Figure 4. Distribution records of specimens sequenced at the COI gene fragment for this study (see Table 1). Small black dots represent all Missulena records from Western Australia (as downloaded from the Atlas of Living Australia - http://www.ala.org.au; accessed 10 April 2017).

us of about at least 100 km of its type locality, the only described species in the genus with the exception of the widespread *M. occatoria* (Walckenaer, 1805) (based on data of the Atlas of Living Australia; http://ala.org.au; accessed 10 April 2017). The species is yet another example of the extremely diverse invertebrate fauna of the semi-arid Goldfields region of Western Australia that is currently poorly studied in relationship to its invertebrate fauna when compared to other bioregions in Western Australia, e.g. the Pilbara (Durrant et al. 2010; McKenzie et al. 2009; Volschenk et al. 2010). Whilst comprehensive biological studies have been conducted in the Goldfields more than two decades ago, these rarely considered invertebrates (Biological Surveys Committee 1984; Keighery et al. 1995). Recent studies on terrestrial snails in the genus *Bothriembryon* Pilsbry, 1894 (Breure and Whisson 2012), millipedes in the genus *Antichiropus* Attems, 1911 (Car and Harvey 2013, 2014) and trapdoor spiders of the family Idiopidae Simon, 1892 (Rix et al. 2017) have highlighted both extreme diversity and endemism in invertebrates within the vast woodlands of the Goldfields and numerous unpublished reports as part of environmental impact assessments have supported these general findings for other taxonomic groups. The fact that *M. harewoodi* sp. n. has never been sampled before near the regional centre of Kalgoorlie-Boulder and remains only known from a single specimen that was collected as by-catch during a vertebrate trapping survey, highlights the need for a comprehensive invertebrate fauna survey of this region that includes both its woodlands and isolated ranges of banded ironstone.

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Walckenaer CA (1805) Tableau des araneides ou caractères essentiels des tribus, genres, families et races que renferme le genre Aranea de Linné, avec la désignation des espèces comprises dans chacune de ces divisions, Paris.