# Phylogenetic placement of eight poorly known spiders of Microdipoena (Araneae, Mysmenidae), with descriptions of five new species 

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#### Abstract

Ten species of the spider genus Microdipoena Banks, 1895 are reported from China, Laos, Indonesia, Georgia, and Seychelles. DNA sequences of the eight species are obtained to confirm their correct identification. The molecular phylogenetic analysis based on five gene fragments ( $16 \mathrm{~S}, 18 \mathrm{~S}, 28 \mathrm{~S}, \mathrm{COI}$, and H 3 ) were used to test the relationships and taxonomic placements of eight Microdipoena species, of which five species are documented as new to science: i.e., M. huisun sp. nov. ( $q$, China), M. lisu sp. nov. ( $\ell$, China), M. shenyang sp. nov. ( ${ }^{\circ}+$, China), M. thatitou sp. nov. ( $\uparrow$, Laos), and M. zhulin sp. nov. (̊̊ㅇ, China). Five known species are redescribed: M. elsae Saaristo, 1978 (̊ , Seychelles), M. gongi (Yin, Peng \& Bao, 2004) (ð千口, China), M. menglunensis (Lin \& Li,  China). All but $M$. menglunensis are diagnosed and illustrated. The family Mysmenidae is also the first recorded from Laos and Georgia.


Key words: Description, Georgia, Indonesia, Laos, mysmenids, new record, new species

## Introduction

Since its inception, the genus Microdipoena Banks, 1895 had a very confusing taxonomic history lasting more than a century. Originally considered as a monotypic genus, it was placed in Theridiidae Sundevall, 1833. Its generotype, Microdipoena guttata Banks, 1895, was designated from Long Island in New York (Banks 1895). Bishop and Crosby (1926) synonymized Microdipoena with Mysmena Simon, 1894 that was accepted by Levi (1956) and Gruia (1977). At almost the same time, Mysmena was transferred from Theridiidae to Symphytognathidae Hickman, 1931 by Forster (1959), and subsequently to Mysmenidae Petrunkevitch, 1928 by Forster and Platnick (1977). Saaristo (1978) considered that the male palp and epigyne of Microdipoena guttata are quite different in structure and revalidated this genus that consists of two species: M. guttata and M. elsae. Shortly thereafter, Brignoli (1980) reported some Oriental and Australian mysmenids and transferred Mysmena jobi in Symphytognathidae (Kraus 1967), Mysmena illectrix and Mysmena saltuensis in Theridiidae (Simon, 1895) to his created
genus Mysmenella Brignoli, 1980. Over the next three decades, some Microdipoena species were placed in so-called "Mysmenella" by other arachnologists (e.g., Baert 1984a, b, 1989; Ono 2007; Yin et al. 2004; Lin and Li 2008, 2013). Based on the phylogenetic hypothesis of Mysmenidae, Lopardo and Hormiga (2015) re-diagnosed and circumscribed Microdipoena and proposed it synonymized with Anjouanella Baert, 1986 and Mysmenella Brignoli, 1980. At this point, the placement and circumscription of this genus has been reasonably confirmed.

Microdipoena is distributed almost worldwide except Antarctica, although it currently consists of only 16 valid species, accounting for about 8.7 percent of 183 mysmenid species (WSC 2023). Known congeners are mainly distributed in Eurasia, Africa, Americas, some Oceanic and Pacific islands.

This paper reports our findings on the study of an inventory specimens collected from China, Laos, Indonesia, Georgia, and Seychelles during 2006 through to 2018, which revealed a total of nine Microdipoena species, including five new to science and four previously known species. The purpose of this study are to sequence five genes of these species, to test their phylogenetic positions and relationships within the genus Microdipoena, and to describe and illustrate the five new Microdipoena species from China and Laos. This paper is also the first report of Mysmenidae from Laos and Georgia.

## Materials and methods

## Species sampling and preservation

Specimens were collected by hand or sifting from leaf litter. All of the specimens were preserved in a $95 \%$ ethanol solution at $-20^{\circ} \mathrm{C}$. All examined materials and molecular vouchers involved in this study are stored in the Natural History Museum of Sichuan University in Chengdu (NHMSU), China.

## Molecular data

To test taxonomic position of these novel species in this study within the Mysmenidae, twelve individuals from eight species were picked out from the examined materials for molecular sequencing. Their prosoma and legs were used to extract genomic DNA and sequence five gene fragments: $16 \mathrm{~S}, 18 \mathrm{~S}, 28 \mathrm{~S}, \mathrm{COI}$, and H3. Primer pairs and PCR protocols are given in Table 1. The abdomens and male palps were kept as vouchers. Whole genomic DNA was extracted from tissue samples with the TIANamp MicroDNA Kit (TIANGEN) following the manufacturer's protocol for animal tissue. The five gene fragments were amplified in $25 \mu \mathrm{~L}$ reactions. Raw sequences were edited and assembled using BioEdit v. 7.2.5 (Hall 1999). Newly obtained DNA sequence data has been uploaded to GenBank for preservation (accession numbers given in Table 2).

We used these new sequences and a selection from previously sequenced taxa to assemble a partial phylogeny of mysmenid spiders, which only involved five representative genus. A total of 26 mysmenid species was used for phylogenetic analysis (Table 2). The ingroup includes 11 known, nine undescribed, and four new mysmenid species. Two Maymena species were used as outgroups (see gray region in Table 2). We used the MAFFT v. 7.450 online server (https://mafft. cbrc.jp/alignment/server/) with default parameters to align the sequences of the

Table 1. Primers and amplification conditions used in PCR.

| Locus | Annealing temperature/time | Direction | Primer | Sequence $5^{\prime} \rightarrow \mathbf{3}^{\prime}$ | Reference |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 16 S | $46.45{ }^{\circ} \mathrm{C} / 30 \mathrm{~s}$ | F | 16sb2_12864 | CTCCGGTTTGAACTCAGATCA | Hormiga et al. 2003 |
|  | $43.65{ }^{\circ} \mathrm{C} / 30 \mathrm{~s}$ | R | LR-J-13360 | GTAAGGCCTGCTCAATGA | In this study |
| 18S | $52.1^{\circ} \mathrm{C} / 30 \mathrm{~s}$ | F | 18S-1F | TACCTGGTTGATCCTGCCAGTAG | Giribet et al. 1996 |
|  |  | R | SSU rRNA reverse | GTGGTGCCCTTCCGTCAATT | Balczun et al. 2005 |
| 28S | $54.9{ }^{\circ} \mathrm{C} / 30 \mathrm{~s}$ | F | 28sa | GACCCGTCTTGAAACACGGA | Rix et al. 2008 |
|  |  | R | LSUR | GCTACTACCACCAAGATCTGCA |  |
| COI | $48.95{ }^{\circ} \mathrm{C} / 30 \mathrm{~s}$ | F | LC01490 | GGTCAACAAATCATAAAGATATTGG | Folmer et al. 1994 |
|  | $46^{\circ} \mathrm{C} / 30 \mathrm{~s}$ | R | HCO2198 | TAAACTTCAGGGTGACCAAAAAATCA |  |
| H3 | $50^{\circ} \mathrm{C} / 30 \mathrm{~s}$ | F | H3nf | ATGGCTCGTACCAAGCAGAC | Colgan et al. 1998 |
|  |  | R | H3nr | ATRTCCTTGGGCATGATTGTTAC |  |

Table 2. List of 26 mysmenid species and their DNA data used for molecular phyligenetic analysis (including new sequences data obtained from asterisked species* in this study).

| Species | Identifier | Source | 16S | 18S | 28 S | COI | H3 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Maymena ambita | Maymena ambita | NCBI | GU456746 | GU456765 | GU456824 | GU456876 | GU456921 |
| Maymena mayana | Maymena mayana | NCBI | HM030403 | HM030411 | HM030421 | - | - |
| Trogloneta granulum | Trogloneta granulum | NCBI | HM030409 | HM030418 | HM030429 | - | - |
| Trogloneta yuensis | XX52F | NHMSU | MZ612929 | MZ613003 | MZ613076 | MZ584802 | MZ603614 |
|  | XX52M | NHMSU | MZ612930 | MZ613004 | MZ613077 | MZ584803 | MZ603615 |
| Trogloneta yunnanensis* | GlgMY49F | NHMSU | OQ756232 | OQ756222 | OQ756224 | OQ756244 | OQ753870 |
|  | GlgMY49M | NHMSU | OQ756233 | OQ756223 | OQ756225 | OQ756245 | OQ753871 |
| Yamaneta kehen | GlgMY15F | NHMSU | MK908793 | MK908809 | MK908801 | MK895534 | MK895542 |
|  | GlgMY15M | NHMSU | MK908792 | MK908808 | MK908800 | MK895533 | MK895541 |
| Yamaneta paquini | GlgMY16M | NHMSU | MK908794 | MK908810 | MK908802 | MK895535 | MK895543 |
|  | GlgMY16F | NHMSU | MK908795 | MK908811 | MK908803 | MK895536 | MK895544 |
| Mysmena sp. MYSM006MAD | MYSM-006-MAD | NCBI | - | GU456774 | GU456832 | GU456882 | GU456929 |
| Mysmena sp. MYSM011ARG | MYSM-011-ARG | NCBI | - | GU456779 | GU456837 | - | GU456934 |
| Mysmena sp. MYSM018MAD | MYSM-018-MAD | NCBI | - | GU456785 | GU456843 | - | GU456939 |
| Mysmena sp. MYSM028MAD | MYSM-028-MAD | NCBI | - | GU456795 | GU456851 | GU456893 | - |
| Mysmenidae sp._MD2476 | Mysmenidae sp._MD2476 | NCBI | - | - | MT651634 | - | - |
| Mysmeninae sp._7502_050 | Mysmeninae sp._7502_050 | NCBI | MG947326 | - | - | - | - |
| Mysmeninae sp._Fuzhou | Fuzhou-Dahuxiang-32 | NCBI | - | - | - | MF467659 | - |
| Microdipoena nyungwe | Microdipoena nyungwe | NCBI | GU456748 | GU456767 | GU456826 | GU456878 | GU456923 |
| Microdipoena guttata | Microdipoena guttata | NCBI | GU456747 | GU456766 | GU456825 | GU456877 | GU456922 |
| Microdipoena sp. AtoL_ARAGH000003 | ARAGH000003 | NCBI | HM030404 | HM030412 | HM030422 | HM030432 | HM030439 |
| Microdipoena sp. MYSM-030-MAD | MYSM-030-MAD | NCBI | - | GU456797 | - | GU456895 | GU456948 |
| Microdipoena elsae* | SEY02F | NHMSU | OQ780298 | OQ780273 | OQ780288 | OQ739554 | 0Q753867 |
| Microdipoena gongi* | XX53F | NHMSU | OQ780302 | 0Q780277 | OQ780292 | OQ739558 | 0Q753861 |
|  | XX53M | NHMSU | - | OQ780278 | OQ780293 | OQ739559 | 0Q753862 |
| Microdipoena huisun sp. nov.* | TW03F | NHMSU | 0Q780299 | 0Q780274 | OQ780289 | OQ739555 | - |
| Microdipoena lisu sp. nov.* | GlgMY83F | NHMSU | OQ780295 | OQ780268 | OQ780283 | OP462192 | 0Q753864 |
| Microdipoena menglunensis* | XSBN20F | NHMSU | OQ780300 | OQ780275 | OQ780290 | 0Q739556 | 0Q753868 |
|  | XSBN20J | NHMSU | OQ780301 | OQ780276 | OQ780291 | OQ739557 | 0Q753869 |
| Microdipoena shengyang sp. nov* | LN01F | NHMSU | - | OQ780269 | OQ780284 | - | - |
|  | LN01M | NHMSU | - | OQ780270 | OQ780285 | - | - |
| Microdipoena yinae* | SC11F | NHMSU | OQ780296 | 0Q780271 | OQ780286 | OQ739552 | OQ753866 |
|  | SC11M | NHMSU | OQ780297 | OQ780272 | OQ780287 | OQ739553 | OQ753865 |
| Microdipoena zhulin sp. nov.* | GX02F | NHMSU | OQ780294 | OQ780267 | OQ780282 | OQ739551 | OQ753863 |

eight Microdipoena species. All sequences were concatenated in sequences Matrix v. 1.7.8 (Vaidya et al. 2011). We used PartitionFinder2 (Lanfear et al. 2017) to identify the best-fit models of molecular evolution for each locus. GTR+I+G was selected for $\mathrm{COI}, \mathrm{H} 3,18 \mathrm{~S}$, and 28S, and GTR+G was selected for 16 S .

Topology The maximum likelihood (ML) tree was constructed using Phylosuite v. 1.2.2 (Zhang et al. 2020) with TBR (Tree-Bisection-Reconnection) branch swapping and 2000 bootstrap replicates with default parameters. Bayesian phylogenetic inference (BI) was performed using MrBayes v. 3.2.7 (Ronquist et al. 2012) through the Cipres Science Gateway (Miller et al. 2010) using four Markov Chain Monte Carlo (MCMCs) chains with default heating parameters for 50,000,000 generations or until the average standard deviation of split frequencies was less than 0.01 . The Markov chains were sampled every 1000 generations, and the first $25 \%$ of sampled trees were burn-in. The program Tracer v. 1.7.1 (Rambaut et al. 2018) was used to analyze the performance of our BI analyses.

## Morphological data

Specimens were examined and measured using a Leica M205 C stereomicroscope. Further details were studied with an Olympus BX 43 compound microscope. Male palps and epigynes dissected from the bodies were photographed with a Canon EOS 60D wide zoom digital camera ( 8.5 megapixels) mounted on an Olympus BX 43 compound microscope. The individual spider was photographed directly under the compound microscope after being reshaped to its natural status. To show more detailed features, epigynes and each disassembled parts of male palps were treated with lactic acid before being embedded in Hoyer`s gum to take photos of the vulvae. The images were montaged using Helicon Focus 3.10.3 (Khmelik et al. 2006) image stacking software.

All measurements are in millimeters. Leg measurements are given as follows: total length (femur, patella, tibia, metatarsus, and tarsus). References to figures in the cited papers are in lowercase (fig. or figs), figures in this paper are noted with an initial capital (Fig. or Figs). Nomenclature of the genital structures was mainly based on Lopardo et al. (2011) for Microdipoena. Abbreviations of terms or institutions used in the text or figures are as follow:

## Male palp

CT cymbial tooth;
Cy cymbium;
CyC cymbial conductor;
CyF cymbial fold;
CyFs setae on cymbial fold;
CyP cymbial process;
E embolus;
Pa patella;
PC paracymbium;
SD spermatic duct;
T tegulum;
TC tegular conductor;
Ti tibia.

## Epigyne

CD copulatory duct;
FD fertilization ducts;
S spermathecae;
Sp scape

## Somatic morphology

AP abdominal protuberance;
FS femoral spot;
MC Metatarsal clasping spine;
TS tibial macrosetae on male leg I.

Institutions acronyms

HNU College of Life Science, Hunan Normal University, Changsha, China;
IZCAS Institute of Zoology, Chinese Academy of Sciences, Beijing, China;
NHMSU Natural History Museum of Sichuan University, Chengdu, China;
SMF Senckenberg Research Institute, Frankfurt, Germany;
ZMUTU Zoological Museum, University of Turku, Turku, Finland.

## Results

## Phylogenetic analysis

The topology inferred by the two different phylogenetic analyses based on the combined sequence dataset of five gene fragments performed (Figs 1, 2) show high consistencies in several mysmenid groupings. High support values are common at each end clade. Except for two Maymena species designated as outgroup, the remaining 24 mysmenid species are divided into four major clades, each of which represents a different genus. Both ML and BI trees analyses recovered Microdipoena as monophyletic and a sister group of Mysmena + Trogloneta + Yamaneta.

In the ML tree, the clade of Yamaneta represented by two known species ( Y . kehen and K . paquini) is near the base of ML and BI trees, with high support (indicated by yellow box in Fig. 1). The clade of Trogloneta containing three known species (T. granulum, T. yuensis, and T. yunnanensis) is also monophyletic with high support (indicated by pink box in Fig. 1). Four undescribed Mysmena (MYSM-011-ARG, MYSM-018-MAD, and MYSM-028-MAD) and Mysmenidae species (Mysmenidae sp._MD2476) composed the clade of Mysmena shown in the blue box in Fig. 1, which is located in the middle of the topological structure trees, between the clades of Trogloneta and of Microdipoena, and also has relatively high support. A clade composed of ten Microdipoena species (including eight species involved in this study, indicated by red font in the pale box in Fig. 1) and five undescribed species (Fuzhou-Dahuxiang-32, MYSM-006-MAD, MYSM-030-MAD, Mysmeninae sp._7502_050, and AtoL_ARAGH000003) were monophyletic, but with low support. These results support our taxonomic classification.


Figure 1. Tree topology obtained by maximum likelihood in IQ-TREE v. 2.0 using combined genes of 18 mysmenid species from NCBI plus eight Microdipoena species (red font). Numbers at nodes indicate bootstrap values. Note fifteen species representing the genus Microdipoena are clustered into a monophyly but with low support; the high support of eight species (red font) in the genus Microdipoena (pale box). Mysmena (including four undescribed species in blue box), Trogloneta (pink box) and Yamaneta (yellow box) are also monophyletic and with high support respectively.

The result of BI is consistent with ML for all major clades (Fig. 2). In the BI topology, seven Chinese and one Seychelles species involved in our study (indicated by red font in Fig. 2), together with two known species (Microdipoena nyungwe, M. guttata), two undescribed species (MYSM-030-MAD, AtoL_ARAGH000003), and three undescribed Mysmeninae species (Fuzhou-Dahuxiang-32, MYSM-006-MAD, and Mysmeninae sp._7502_050) form a separate monophyletic, lower supported clade compared to Mysmena, Trogloneta, and Yamaneta. However, each species has high support at each end clade of the BI tree respectively. The available molecular evidence seems sufficient to justify the taxonomic placement of four new and four known Microdipoena species in this study.


Figure 2. Tree topology from Bayesian analysis in MrBayes v. 3.2.7. Numerical values at nodes indicate posterior probabilities; other comments as in Fig. 1. Note the high support of eight species (red font) in the clade of Microdipoena (pale box), which is monophyletic but with low support. Other three clades, Mysmena (blue box), Trogloneta (pink box), and Yamaneta (yellow box), are also monophyletic respectively and with high support.

## Taxonomy

Family Mysmenidae Petrunkevitch, 1928

## Genus Microdipoena Banks, 1895

Mysmena Simon, 1895: 149.
Microdipoena Banks, 1895: 85 (synonymized by Bishop and Crosby 1926: 127). Mysmena Bishop \& Crosby, 1926: 177.
Microdipoena Saaristo, 1978: 124.
Mysmenella Brignoli, 1980: 731 (synonymized by Lopardo and Hormiga 2015: 783). Anjouanella Baert, 1986: 265 (synonymized by Lopardo and Hormiga 2015: 783). Microdipoena Lopardo \& Hormiga, 2015: 783.

Type species. Microdipoena guttata Banks, 1895 by original designation; type locality Long Island, New York, USA.

Diagnosis. The male can be distinguished from other mysmenids by there are two or three distal-prolateral macrosetae on the tibia I (Figs 3A, 8A, 11A, $15 \mathrm{~A}, 21 \mathrm{~B}$ ). The palpal bulb is very large (at least $4-5 \times$ tibia in size, $2-3 x$ in other mysmenids). The cymbium has an apical part with complex structure, which specialized as a cymbial conductor and a cymbial process (Figs 6H, 9C, 12B, 16B, 22D). A lobe-shaped paracymbium bears several long setae along its edge (Figs 4C, 6H, 9B, 12B, F, 16B, 20D, 22D). The thick embolus folds into a twisted complex structure at distal end, wrapped by a membranous structure on the apex of cymbium (Figs 9A-C, 12D-G, 16C-G, 20D, E, 22A, F-H) (except in M. comorensis, M. elsae, M. guttata, M. mihindi, M. nyungwe, and M. vanstallei, without complex structure, but with either a distal apophysis or irregular membrane). The female differed from other mysmenids by the abdomen with a whitish ventral ring around the spinnerets (Figs 3F, 11E, 19D, 21F) (except M. comorensis, with all ventral abdominal area paler). The shape of spermathecae are mostly round, oval, or semicircular, wrapped by membranous copulatory ducts from posterior or around (Figs 7C, 9F, 10F, 13C, 14F, 17C, 18F, 23D).

Composition. Twenty one species: Microdipoena comorensis (Baert, 1986)
 M. guttata Banks, 1895 ( ${ }^{1}$ q), M. huisun sp. nov. ( ( $)$, M. illectrix (Simon, 1895)

 tai (Ono, 2007) (ठ̊) , M. papuana (Baert, 1984) (ठ), M. pseudojobi (Lin \& Li, 2008) ( ${ }^{\top}$ q), M. saltuensis (Simon, 1895) ( q ), M. samoensis (Marples, 1955) (§?), M. shenyang sp. nov. (§ㅇ), M. thatitou sp. nov. ( ( ) , M. vanstallei Baert,


Distribution. From Europe to the Caucasus, from East Asia to Southeast Asia and South Asia to the Middle East, from central Africa to Madagascar and Seychelles, from USA to Paraguay, from New Guinea to Samoa and Hawaii.

## Microdipoena elsae Saaristo, 1978

Figs 3-7

Microdipoena elsae Saaristo, 1978: 124, figs 255-265 (ð̛o); Saaristo 2010: 92, fig. 17.1-17.8 (ð) ${ }^{\text {® }}$ ); Lopardo et al. 2011: 287, fig. 9a (ô); Lopardo and Hormiga 2015: 783, figs 17A-D, 129D, 132A, 141M, N (đ̊)
Mysmena elsae: Roberts 1978: 932, figs 65-73 (ôq); Logunov 2022: 89, fig. 18 ( ${ }^{\top}$ ).

Type material. Holotype $q$ (ZMUTU), allotype $1 \uparrow 4 \widehat{ }$ (ZMUTU), and paratypes 8 \& (ZMUTU) Seychelles: Mahé near La Misére, sieving leaf litter, 600 m elev., 30.V.1975, M. Saaristo leg. Not examined.

Other material examined. $2 \uparrow 3 q 3$ juvs (NHMSU-SEY02), Seychelles: Mane, at half of Morne Blanc, a pile of chopped wood ( $4^{\circ} 39.553 '$ S, $55^{\circ} 26.199^{\prime} \mathrm{E} ; 461 \mathrm{~m}$ elev.), 30.VI.2013, H. Zhao leg.; 2 q 1 juv (NHMSU-SEY01), La Digue, Belle-Vue Mountain ( $4^{\circ} 21.611^{\prime} \mathrm{S}, 55^{\circ} 50.470^{\prime} \mathrm{E} ; 213 \mathrm{~m}$ elev.), 5.VII.2013, H. Zhao leg.


Figure 3. Microdipoena elsae Saaristo, 1978, male (A-C) and female (D-F) from Seychelles A habitus, dorsal B habitus, ventral $\mathbf{C}$ habitus, lateral $\mathbf{D}$ habitus, lateral $\mathbf{E}$ habitus, dorsal $\mathbf{F}$ habitus, ventral. Abbrrviations: $\mathrm{FS}=$ femoral spot, MC = Metatarsal clasping spine, TS = tibial spine on male leg I. Scale bars: 0.50 mm .


Figure 4. Microdipoena elsae Saaristo, 1978, from Seychelles A male palp, apical B male palp, dorsal C male palp, prolateral D male palp, retrolateral. Abbreviations: $C y=$ cymbium, $C y C=$ cymbial conductor, $C y F s=$ setae on cymbial fold, $\mathrm{E}=$ embolus, $\mathrm{Pa}=$ patella, $\mathrm{PC}=$ paracymbium, $\mathrm{SD}=$ spermatic duct, $\mathrm{T}=$ tegulum, $\mathrm{Ti}=$ tibia. Scale bars: 0.10 mm .

Diagnosis. Male of Microdipoena elsae differs from other congeners except for M. comorensis (Baert, 1986), M. guttata Banks, 1895, M. nyungwe Baert, 1989, M. vanstallei Baert, 1985 by the filiform embolus without a distal twisted complex structure. Its male seems to be most similar to $M$. nyungwe, but can be distinguished by the embolus having a small membranous hook at the intermediate constriction and the absence of cymbial groove (cf. Figs 5D, 6C and Lopardo and Hormiga 2015: 554, fig. 22C, F, G). Its female is similar to M. nyungwe and $M$. guttata but distinguished by the ovate spermathecae (round in $M$. nyungwe and $M$. guttata), the fertilization duct is slightly sclerotized and longer (cf. Fig. 7B, C vs Lopardo and Hormiga 2015: 554, 672, figs 18G, 22B, 129A, B).

Description. Male: Total length 1.04 . Carapace 0.44 long, 0.42 wide, 0.44 high. Clypeus 0.10 high. Sternum 0.33 long, 0.30 wide. Abdomen 0.60 long, 0.63 wide, 0.88 high. Length of legs: I $1.17(0.44,0.14,0.25,0.16,0.18)$; 11.06 ( $0.42,0.14,0.22$, $0.12,0.16)$; III 0.78 ( $0.22,0.12,0.14,0.14,0.16$ ); IV 1.08 ( $0.25,0.12,0.27,0.20,0.24$ ).

Somatic characters (Fig. 3A-C). Coloration: carapace dark brown centrally, yellow brown marginally. Ocular base black. Chelicera, endites, and labium yellow. Sternum yellow with two brown stripes. Legs yellow and black. Abdomen dark brown with white spots dorsally, yellow with brown spots ventrally. Prosoma: carapace nearly pear-shaped in dorsal view and peak-shaped in lateral view. Cephalic area upheaved. Sternum triangular, slightly plump, covered with sparse, short setae. Legs: covered with setae. Mating clasper on metatarsus I, two macrosetae on tibia I, femur I with femoral spot. Abdomen: nearly ovoid.

Palp (Figs 4-6): large, ca as big as $1 / 2$ size of the carapace. Cymbium translucent, distal end specialized as a broad, collared cymbial conductor, and a small cymbial process, modified by weakly sclerotized folds and a row of stiff short setae (Figs 4D, $5 \mathrm{C}, 6 \mathrm{~F}-\mathrm{H}$ ). Paracymbium smooth, with long setae at the edge. Conductor wide, sclerotized, with two upper and a lower processes (Fig. 6A, B). Tegulum sclerotized, with two upper (a wide, a narrow) and one lower (a narrow) processes (Fig. 6D, E). Embolus filiform, with a membranous hook at the constriction near the middle (Figs 5D, 6C), its distal part coiled into 2.5 loops around cymbial conductor (Figs 4, $5 C$ ). Spermatic ducts faintly visible through the surface of palpal bulb and cymbium.

Female. Total length 0.92 . Carapace 0.40 long, 0.46 wide, 0.58 high. Clypeus 0.08 high. Sternum 0.28 long, 0.30 wide. Abdomen 0.52 long, 0.46 wide, 0.62 high. Length of legs: I $1.77(0.52,0.16,0.40,0.34,0.35)$; II $1.38(0.46,0.14,0.34,0.20$, 0.24 ); III 0.98 ( $0.30,0.12,0.20,0.16,0.20$ ); IV 1.08 ( $0.30,0.10,0.24,0.20,0.24$ ).

Somatic characters (Fig. 3D-F). Coloration: carapace dark brown centrally, yellow-brown marginally. Ocular base black. Chelicera, endites, and labium yellow. Sternum yellow with two brown stripes. Legs yellow and black. Abdomen dark brown with white spots dorsally, yellow with brown spots ventrally. Prosoma: carapace nearly pear-shaped in dorsal view. Cephalic part slightly elevated. Sternum triangular, covered with sparse short setae. Legs: covered with setae and bristles. Femurs I and II with femoral spot. Abdomen: nearly globose.

Epigyne (Fig. 7A-C): spermathecae heavily sclerotized, nearly vertically ovoid, spaced by ca $3 \times$ their width. Copulatory duct almost all membranous cystic structure with irregular folds, surround the entirely spermathecae, which enters the spermathecae from posteromedially after gradually harden at the posterior area of spermathecae. Weakly sclerotized fertilization duct starts at the posterolateral side of spermatheca, and then folds back toward the center of vulva (Fig. 7C).

Distribution. Seychelles, Congo, and Comoros.


Figure 5. Microdipoena elsae Saaristo, 1978, from Seychelles A conductor, from behind $\mathbf{B}$ tegulum, from behind $\mathbf{C}$ male palp, ventral $\mathbf{D}$ bulbus with conductor and tegulum removed, dextrolateral. Abbreviations: $\mathrm{E}=$ embolus, $\mathrm{T}=$ tegulum. Scale bars: 0.10 mm .


Figure 6. Microdipoena elsae Saaristo, 1978, from Seychelles A conductor, dorsal B conductor, ventral C embolic distal end, dextrolateral $\mathbf{D}$ tegulum, dorsal $\mathbf{E}$ tegulum, ventral $\mathbf{F}$ cymbium, dorsal $\mathbf{G}$ cymbium, apical-lateral $\mathbf{H}$ cymbium, prolateral. Abbreviations: $\mathrm{Cy}=$ cymbium, $\mathrm{CyC}=$ cymbial conductor, $\mathrm{CyF}=$ cymbial fold, $\mathrm{CyFs}=$ setae on cymbial fold, $\mathrm{CyP}=$ cymbial process, PC = paracymbium. Scale bars: 0.10 mm .


Figure 7. Microdipoena elsae Saaristo, 1978, from Seychelles A epigyne, ventral B vulva, ventral C vulva, dorsal. Abbreviations: $C D=$ copulatory duct, $F D=$ fertilization duct, $S=$ spermathecal, $S p=$ scape. Scale bars: 0.10 mm .

## Microdipoena gongi (Yin, Peng \& Bao, 2004)

Figs 8, 9

Mysmenella gongi Yin, Peng \& Bao, 2004: 80, figs 1-8 (ð̊ㅇ). Microdipoena gongi: Lopardo and Hormiga 2015: 783.

Type material. Holotype $q$ (HNU) and paratypes $1 \delta 3 q$ (HNU), China: Hunan Province, Daoxian County, Timber Mill ( $25^{\circ} 31.000^{\prime} \mathrm{N}, 111^{\circ} 33.000^{\prime} \mathrm{E}$ ), 8.IV.1988; 1q7§, China: Hunan Province, Daoxian County, Shuangqiao Town, 1.VI.1987, L. Gong leg. Examined in 2008.

Other material examined. $1 \delta 2$ (NHMSU-XX53), China: Hunan Province, Changsha City, Yuelu District, Yuelu Mountain scenic spot, leaf litter ( $28^{\circ} 10.502^{\prime} \mathrm{N}, 112^{\circ} 56.391^{\prime} \mathrm{E} ; 121 \mathrm{~m}$ elev.), 20.IV.2018, G. Zhou leg.

Diagnosis. Male of Microdipoena gong can be distinguished from other congeners except for M. illectrix, M. jobi, M. menglunensis, M. mihindi, M. ogatai, M. papuana, M. pseudojobi, M. samoensis, M. shenyang sp. nov., M. yinae, and
M. zhulin sp. nov. by the embolic end twisted into a complex structure (Figs 9A, $B, 12 D, 16 E, 20 B, 22 A)$. It can be distinguished from these aforementioned species by the cymbium wraps the bulb prolaterally or lacking a cymbial tooth (cf. Figs 9A, B, 12B, 16B, 22D; Lin and Li 2013: CP shown in fig. 21F; Lopardo and Hormiga 2015: CyP shown in fig. 132C-F). Female of Microdipoena gong differs from other congeners except for $M$. huisun sp. nov., M. ogatai, and $M$. yinae in lacking a long, soft, membranous scape, but having a small, weakly sclerotized, small scape (cf. Figs 9F, 10F, 19H, and Ono 2007: figs 69, 70; Figs 13C, 14F, 17C, 18F, 23D and Lopardo and Hormiga 2015: fig. 129E, F), but can be distinguished from these aforementioned species by the horizontal ovate spermathecae, which is almost as large as the posterior sclerotized area of copulatory ducts (globose in M. huisun sp. nov., smaller spermatheca in M. ogatai, inclined hemispheric spermatheca in M. yinae) (cf. Figs 9F, 10F, 19H and Ono 2007: fig. 70).

Description. Male: Total length 0.98 . Carapace 0.44 long, 0.48 wide, 0.52 high. Clypeus 0.08 high. Sternum 0.30 long, 0.30 wide. Abdomen 0.54 long, 0.56 wide, 0.60 high. Length of legs: I $1.33(0.48,0.12,0.27,0.22,0.24)$; II 1.20 ( $0.42,0.12,0.24,0.18,0.24$ ); III 0.81 ( $0.25,0.10,0.16,0.14,0.16$ ); IV 1.04 ( 0.28 , $0.12,0.24,0.18,0.22)$.

Somatic characters (Fig. 8A-C). Coloration: carapace dark brown. Ocular base black. Chelicera, endites, labium and sternum dark brown. Legs brownblack. Abdomen dark with three white spots dorsally. Prosoma: carapace nearly round in dorsal view and peak-shaped in lateral view. Cephalic part flat, slightly elevated. Sternum scutiform, slightly plump. Legs: covered with setae. Mating clasper on metatarsus I, two strong spines on tibia I, femur I with sclerotized femoral spot. Abdomen: nearly globose.

Palp (Fig. 9A-C): The palp $45^{\circ}$ inclined to the surface of the tibia. Cymbium translucent, originating prolaterally, with a large cymbial conductor. Paracymbium large, finger-like, with long setae. Tegulum translucent, surface swollen. Embolus thin and relatively short, coiled into one loop over the cymbium, tip with complex structure. Spermatic ducts can be seen through translucent tegulum.

Female. Total length 1.37. Carapace 0.45 long, 0.52 wide, 0.40 high. Clypeus 0.10 high. Sternum 0.38 long, 0.32 wide. Abdomen 0.92 long, 0.88 wide, 0.86 high. Length of legs: I $1.32(0.34,0.14,0.32,0.28,0.24)$; II $1.06(0.22,0.12$, $0.30,0.24,0.18)$; III $0.82(0.20,0.12,0.20,0.16,0.14)$; IV $0.92(0.22,0.12,0.22$, $0.20,0.16$ ).

Somatic characters (Fig. 8D-F). Coloration: carapace nearly black. Ocular base black. Chelicera, endites, labium black, sternum dark brown. Legs brown-black. Abdomen black with four white spots dorsally. Prosoma: carapace nearly round in dorsal view. Cephalic part unraised. Sternum scutiform, slightly plump, covered with sparse setae. Legs: covered with setae and bristles. Femurs I and II with sclerotized femoral spot. Abdomen: nearly spherical, covered with short setae.

Epigyne (Fig. 9D-F): sclerotized, length twice the width, structure can be seen slightly through the cuticle. Scape nodular shape, very small. Copulatory duct membranous, coiled under the spermathecae. Paired spermathecae transverse oval, separated by nearly double diameter. Fertilization ducts membranous, inconspicuous.

Distribution. China (Hunan).


Figure 8. Microdipoena gongi Yin, Peng \& Bao, 2004, male (A-C) and female (D-F), from Hunan of China A habitus, dorsal B habitus, ventral $\mathbf{C}$ habitus, lateral $\mathbf{D}$ habitus, lateral $\mathbf{E}$ habitus, dorsal $\mathbf{F}$ habitus, ventral. Abbreviations: $F S=$ femoral spot, MC = Metatarsal clasping spine, TS = tibial spine on male leg I. Scale bars: 0.50 mm .


Figure 9. Microdipoena gongi Yin, Peng \& Bao, 2004, from Hunan of China A male palp, prolateral B male palp, retrolateral $\mathbf{C}$ male palp, apical $\mathbf{D}$ epigyne, ventral $\mathbf{E}$ vulva, ventral $\mathbf{F}$ vulva, dorsal. Abbreviations: $\mathrm{CD}=$ copulatory duct, $\mathrm{Cy}=$ cymbium, $\mathrm{CyC}=$ cymbial conductor, $\mathrm{E}=$ embolus, $\mathrm{Pa}=$ patella, $\mathrm{PC}=$ paracymbium, $\mathrm{S}=$ spermathecal, $\mathrm{SD}=$ spermatic duct, $\mathrm{Sp}=$ Scape, $\mathrm{T}=$ tegulum, $\mathrm{Ti}=$ tibia. Scale bars: 0.10 mm .

## Microdipoena huisun sp. nov.

https://zoobank.org/36513469-CD9E-483A-B226-E13FBD5FC82B
Fig. 10

Type material. Holotype $q$ and paratype 4 $q$ (NHMSU-TW03), China: Taiwan Province, Nantou County, Huisun Forest Farm ( $24^{\circ} 05.279^{\prime} \mathrm{N}, 121^{\circ} 02.078^{\prime} \mathrm{E}$; 788 m elev.), 1.VII.2013, G. Zheng leg.

Etymology. The specific name is derived from the type locality; noun in apposition.

Diagnosis. This new species differs from other species except for M. gongi, $M$. ogatai, and $M$. yinae by the nearly spherical spermathecae and having a small scape rather than a long, soft, membranous one (cf. Figs 10E, F, 13C, $14 \mathrm{~F}, 17 \mathrm{C}$ ). It can be distinguished from M. gongi, $M$. ogatai, and $M$. yinae by the nearly spherical spermathecae and the narrower posterior sclerotized area of copulatory ducts (cf. Figs 10F, 9F, 19H and Ono 2007: fig. 70).

Description. Female (holotype): Total length 0.76. Carapace 0.28 long, 0.29 wide, 0.28 high. Clypeus 0.07 high. Sternum 0.24 long, 0.19 wide. Abdomen 0.48 long, 0.48 wide, 0.50 high. Length of legs: I 0.94 ( $0.30,0.08,0.22,0.16$, 0.18 ); II 0.92 ( $0.32,0.08,0.22,0.14,0.16$ ); III $0.70(0.22,0.08,0.14,0.12,0.14)$; IV 0.94 ( $0.24,0.08,0.24,0.18,0.20$ ).

Somatic characters (Fig. 10A-C). Coloration: carapace nearly silver-brown centrally, black marginally. Ocular base black. Chelicera, endites yellow, labium silver-brown, sternum pale yellow with four symmetrical black spots. Legs brownblack. Abdomen black with multiple symmetrical white spots dorsally, black with symmetrical white and yellow spots ventrally. Prosoma: carapace nearly pearshaped in dorsal view. Cephalic part unraised. Sternum triangular, covered with sparse setae. Legs: covered with setae and bristles. Femurs I and II with sclerotized femoral spot. Abdomen: nearly oval in dorsal view, covered with sparse setae.

Epigyne (Fig. 10D-F): scape nodular shape, very small. Copulatory duct membranous, coiled under the spermathecae. Paired spermathecae nearly globose, separated by nearly $3 \times$ its diameter. Fertilization ducts membranous, inconspicuous.

Male. Unknown.
Distribution. China (Taiwan).

## Microdipoena jobi (Kraus, 1967)

Figs 11-13

Mysmena jobi Kraus, 1967: 392, figs 12-18 (§̊).
Mysmenella jobi: Brignoli 1980: 731; Namkung 2002: 146, fig. 16a, b (ôq).
Microdipoena jobi: Lopardo and Hormiga 2015: 783 (§) $\uparrow$ ); Naumova et al. 2017:
478, figs 12-14 (ð).
 (Mus. Paris) France: Paris, Mainz-Gonsenheim, Gonsenheimer forest, forests trap, 20.IV.1967, W. Job leg. Examined in 2008.

Other material examined. $1 \$^{\top} \uparrow$ (NHMSU-No.57), Georgia: Adjara, road along Acharistsqali River, dry stone debris near bridge ( $41^{\circ} 34.720^{\prime} \mathrm{N}, 41^{\circ} 51.760^{\prime} \mathrm{E} ; 113$ m elev.), 21.VII.2012, S. Li leg.

Diagnosis. This species is similar to M. gongi, M. illectrix, M. menglunensis, M. mihindi, M. ogatai, M. papuana, M. pseudojobi, M. samoensis, M. shenyang sp. nov., $M$. yinae, and $M$. zhulin sp. nov. in having a twisted complex structure at the embolus end, but can be distinguished by the peculiar shape of conductor with a huge lower lateral process, the cymbial tooth near the ventral center of cymbium, the spherical spermathecae spaced by ca $2.3 \times$ their diameter, and the long, soft scape (cf. Figs 12B, C, F, G, 13B, C, 14F, 16A, B, 17C, 20C, 22A, B, D, 23D, and Lin and Li 2013: fig. 21F; Lin and Li 2008: figs 11F, G, 12F; Lopardo and Hormiga 2015: figs 129E, F, 132C-E).

Description. Male (holotype): Total length 0.96 . Carapace 0.46 long, 0.50 wide, 0.50 high. Clypeus 0.06 high. Sternum 0.32 long, 0.26 wide. Abdomen 0.5 long, 0.46 wide, 0.50 high. Length of legs: I $1.42(0.52,0.08,0.40,0.20,0.22)$; II 1.22 ( $0.40,0.08,0.32,0.20,0.22$ ); III 0.72 ( $0.20,0.08,0.18,0.12,0.14$ ); IV 1.98 ( $0.20,0.08,0.26,0.20,0.24$ ).

Somatic characters (Fig. 11A-C). Coloration: carapace silver-brown. Ocular base black. Chelicera, endites, labium yellow; sternum yellow with two longitudinal brown stripes. Legs yellow-brown. Abdomen black with symmetrical white spots dorsally, yellow with two symmetrical white stripes ventrally. Prosoma: carapace nearly oval in dorsal view and peak-shaped in lateral view. Cephalic part elevated. Sternum scutiform, plump, covered with sparse setae. Legs: covered with setae. Mating clasper on metatarsus I, two strong spines on tibia I, femur I with sclerotized femoral spot. Abdomen: nearly oval in dorsal view.

Palp (Fig. 12A-G): The palp $45^{\circ}$ inclined to the surface of the tibia. Cymbium translucent, cymbial tooth on the prolateral, sclerotized; the tip specialized as cymbial conductor; a large cymbial process at the contralateral of the cymbial conductor. Paracymbium small, with long setae. Conductor 7-shaped, sclerotized, with two large apophyses apically and two pointed apophyses basally. Tegulum translucent, swollen surface. Embolus wide and Iong, the tip with complex structure, most structure coiled into two loops. Spermatic ducts can be seen through translucent tegulum.

Female. Total length 1.38 . Carapace 0.40 long, 0.40 wide, 0.33 high. Clypeus 0.06 high. Sternum 0.32 long, 0.30 wide. Abdomen 0.98 long, 0.92 wide, 0.66 high. Length of legs: I $1.32(0.36,0.16,0.30,0.24,0.26)$ : II 1.12 ( $0.32,0.16$, $0.26,0.18,0.20$ ); III $0.70(0.16,0.12,0.14,0.12,0.16)$; IV 0.84 ( $0.32,0.12,0.16$, $0.12,0.12$ ).

Somatic characters (Fig. 11D-F). Coloration: carapace dark brown. Ocular base black. Chelicera, endites, labium yellow; sternum yellow with two longitudinal brown stripes. Legs brown-black. Abdomen silvery brown with multiple white spots dorsally, brown with two symmetrical white stripes and multiple little white spots ventrally. Prosoma: carapace nearly pear-shaped in dorsal view. Cephalic part unelevated. Sternum scutiform, slightly plump, covered with sparse setae. Legs: covered with setae and bristles. Femurs I and II with sclerotized femoral spot. Abdomen: nearly spherical in dorsal view, covered with sparse setae.

Epigyne (Fig. 13A-C): slightly sclerotized. Scape long, base wide, with thin folds. Copulatory duct long, membranous, with sclerotized S-shaped, coiled under the spermathecae. Paired spermathecae transverse ovoid, separated by more than $3 \times$ diameter. Fertilization ducts membranous, inconspicuous.

Distribution. France, Georgia (Adjara), Caucasus, Iran, China, Korea, and Japan.


Figure 10. Microdipoena huisun sp. nov., female from Taiwan of China $\mathbf{A}$ habitus, dorsal $\mathbf{B}$ habitus, ventral $\mathbf{C}$ habitus, lateral $\mathbf{D}$ epigyne, ventral $\mathbf{E}$ vulva, ventral $\mathbf{F}$ vulva, dorsal. Abbreviations: $C D=$ copulatory duct, $\mathrm{S}=$ spermathecal, $\mathrm{Sp}=\mathrm{Scape}$. Scale bars: $0.50 \mathrm{~mm}(\mathbf{A}-\mathbf{C}), 0.10 \mathrm{~mm}(\mathbf{D}-\mathbf{F})$.


Figure 11. Microdipoena jobi Kraus, 1967, male (A-C) and female (D-F) from Georgia A habitus, dorsal B habitus, ventral $\mathbf{C}$ habitus, lateral $\mathbf{D}$ habitus, lateral $\mathbf{E}$ habitus, dorsal $\mathbf{F}$ habitus, ventral. Abbreviations: $F S=$ femoral spot, MC $=$ Metatarsal clasping spine, TS = tibial spine on male leg I. Scale bars: 0.50 mm .


Figure 12. Microdipoena jobi Kraus, 1967 from Georgia A bulbus with conductor removed, from behind B cymbium, prolateral C conductor, dorsal D embolic end, retrolateral E male palp, apical F male palp, prolateral G male palp, retrolateral. Abbreviations: $\mathrm{CT}=$ cymbial tooth, $\mathrm{Cy}=$ cymbium, $\mathrm{CyC}=$ cymbial conductor, $\mathrm{CyP}=$ cymbial process, $\mathrm{E}=\mathrm{embolus}$, $\mathrm{Pa}=$ patella, $\mathrm{PC}=$ paracymbium, $\mathrm{T}=$ tegulum, $\mathrm{Ti}=$ tibia. Scale bars: 0.10 mm .


Figure 13. Microdipoena jobi Kraus, 1967 from Georgia A epigyne, ventral B vulva, ventral C vulva, dorsal. Abbreviations: $C D=$ copulatory duct, $F D=$ fertilization duct, $S=$ spermathecal, $S p=$ Scape. Scale bars: 0.10 mm .

## Microdipoena lisu sp. nov.

https://zoobank.org/9BFA1810-CC60-4F33-8555-12EC6BB10603
Fig. 14

Type material. Holotype $q$ and paratype $1 q$ (NHMSU-Glg83), China: Yunnan Prov., Gongshan Co., Bingzhongluo Town, halfway from Bingzhongluo to Puhuasi Temple, leaf litter in roadside native forest ( $28^{\circ} 01.420^{\prime} \mathrm{N}, 098^{\circ} 36.133^{\prime} \mathrm{E}$, 1867 m elev.), 15.VIII.2018, Y. Lin et al. leg.

Etymology. The new species is named after the Lisu people, an ethnic minority mainly living in the Nujiang River basin in Yunnan Prov.; noun in apposition.

Diagnosis. The new species seems similar to $M$. jobi and $M$. pseudojobi in the configuration of the vulva and having a long, soft, membranous scape, but can be distinguished by the subtle difference in the shape of the spermathecae, the ca $3 \times$ spacing of the spermathecae, and the copulatory duct enters spermatheca from the posterolateral position (cf. Figs 14E, F, 13B, C and Lin and Li 2008: fig. 12F).

Description. Female (holotype): Total length 1.16. Carapace 0.48 long, 0.44 wide, 0.42 high. Clypeus 0.08 high. Sternum 0.24 long, 0.20 wide. Abdomen 0.68 long, 0.54 wide, 0.60 high. Length of legs: I $1.30(0.46,0.18,0.21,0.17$, $0.28)$; II 1.26 ( $0.32,0.18,0.24,0.23,0.29$ ); III $0.80(0.20,0.15,0.18,0.16,0.21)$; IV 1.24 ( $0.29,0.15,0.28,0.22,0.30$ ).

Somatic features (Fig. 14A-C): Body fulvous. Abdomen with irregular paler patches of all sizes. A bright yellow longitudinal stripe at middle line of sternum. Legs pale yellow, with darkish pigment. Carapace pear-shaped. Ocular base black. Eight eyes in two rows. AME black, the rest white. AER procurved, PER straight. Lateral eyes adjacent. Mouthparts pale brown. Labium fused to sternum. Sternum heart-shaped, longer than wide. Femoral spot on legs I and II. Abdomen dorsally short, ovate, posterior integument slightly ridged.

Epigyne (Fig. 14D-F): epigynal area dark, inner structures not visible. Scape long, membranous, slender, ridged, with tiny notch at tip. Most of copulatory ducts membranous, rugose, with weakly sclerotized duct distally, entering from ventral side of spermathecae. Sclerotized, ovoid spermathecae separated by ca $3 \times$ their width. Fertilization ducts translucent, membranous, arising from posteromedial side of spermathecae, distally intertwined with membranous parts of copulatory ducts.

Male. Unknown.
Distribution. China (Gongshan Co., Yunnan).

## Microdipoena menglunensis (Lin \& Li, 2008)

Mysmenella menglunensis Lin \& Li, 2008: 506, fig. 13A-I ( ${ }^{\text {® }}$ ).
Microdipoena menglunensis: Zhang et al. 2022: 71, figs 7A-F, 8A-F, 9A-D (ơq).

Material examined. 3才 $5 \nrightarrow$ (MNHSU-BN110), China: Yunnan, Mengla, Menglun, XTBG, Rubber-Tea plantation (about 20 yr.) ( $21^{\circ} 54.684^{\prime} \mathrm{N}, 101^{\circ} 16.319^{\prime} \mathrm{E}$; $569 \pm 11 \mathrm{~m}$ elev.), by pitfall trapping, 16-31.V.2007, G. Zheng leg.; $3{ }^{\wedge} 3 \nrightarrow(\mathrm{MN}-$ HSU-BN111), China: Yunnan, Mengla, Menglun, XTBG, Rubber plantation (about


Figure 14. Microdipoena lisu sp. nov., female from Yunnan of China $\mathbf{A}$ habitus, dorsal $\mathbf{B}$ habitus, ventral $\mathbf{C}$ habitus, lateral D epigyne, ventral $\mathbf{E}$ vulva, ventral $\mathbf{F}$ vulva, dorsal. Abbreviations: $C D=$ copulatory duct, $\mathrm{FD}=$ fertilization duct, $\mathrm{S}=$ spermathecal, $\mathrm{Sp}=$ Scape. Scale bars: 0.50 mm (A-C); 0.10 mm (D-F).

20 yr .) ( $21^{\circ} 54.483^{\prime} \mathrm{N}, 101^{\circ} 15.978^{\prime} \mathrm{E} ; 586 \pm 9 \mathrm{~m}$ elev.), by searching, $4-11 . I V .2007$, G. Zheng leg.; $2{ }^{\AA}$ juv. $5 \ell$ (NHMSU-XSBN20), China: Yunnan, Mengla, Menglun, XTBG, tropical rainforest ( $21^{\circ} 55.020^{\prime} \mathrm{N}, 101^{\circ} 16.500^{\prime} \mathrm{E} ; 558 \mathrm{~m}$ elev.), 5.X.2017, Y. Lin and Y. Li leg.

Diagnosis and description. See Zhang et al. 2022: 71-75.
Distribution. China (Xishuangbanna of Yunnan).

## Microdipoena shenyang sp. nov.

https://zoobank.org/6EB3FEB4-7886-4EBA-B8F2-021EF7883C05
Figs 15-17

Type material. Holotype $q$ and paratype $7 \widehat{\AA} 1 \not \subset$ (NHMSU-LN01); 3§ (NHM-SU-No.63); $7 \widehat{6} 6$ (NHMSU-No.64) China: Liaoning Province, Shenyang City, Huanggu District, Beiling Forest Park ( $42^{\circ} 01.503^{\prime} \mathrm{N}, 123^{\circ} 43.823^{\prime} \mathrm{E} ; 352 \mathrm{~m}$ elev.), 15-25.X.2010, X. Sun leg.

Etymology. The specific name is derived from the type locality; noun in apposition.

Diagnosis. Male of this new species differs from other congeners by the embolic end twisted into a complex structure, having a cymbial tooth, which is located on the dorsal edge of cymbium, the peculiar shape conductor with a small membranous process and a large, thickened process on lower sides (cf. Figs 12B, C, F, 16A, B, F, 20C, 22B, D, and Lopardo and Hormiga 2015: fig. 132A-F). Female seems similar to M. jobi and M. lisu sp. nov. in the configuration of vulva and having a long scape with wide at basally and weakly sclerotized at distally, but can be distinguished by the smaller spermathecae spaced by ca $3.5 \times$ their diameter and the more visible fertilization ducts (cf. Figs 17C, 13C, 14F).

Description. Male: Total length 1.02. Carapace 0.40 long, 0.48 wide, 0.48 high. Clypeus 0.08 high. Sternum 0.28 long, 0.34 wide. Abdomen 0.62 long, 0.62 wide, 0.70 high. Length of legs: I $1.24(0.30,0.12,0.28,0.26,0.28)$; II 1.14 ( $0.26,0.12,0.26,0.24,0.26$ ); III 1.03 ( $0.26,0.12,0.22,0.19,0.24$ ); IV 1.06 ( 0.28 , $0.12,0.20,0.22,0.24)$.

Somatic characters (Fig. 15A-C). Coloration: carapace dark brown. Chelicera, endites, labium dark yellow, sternum yellow with two small brown spots. Legs yellow-brown. Abdomen black with symmetrical white spots dorsally and ventrally. Prosoma: carapace nearly round in dorsal view and peak-shaped in lateral view. Cephalic part elevated and flat. Sternum scutiform, plump, covered with sparse setae. Legs: covered with setae. Mating clasper on metatarsus I, two strong spines on tibia I. Abdomen: nearly globose in dorsal view, cover pale setae.

Palp (Fig. 16A-G): The cymbium long and translucent, covering $3 / 4$ of the bulb horizontally, cymbial tooth on the distal cymbium, the tip specialized as cymbial conductor, and the other tip forming a large cymbial process. Paracymbium large, with long setae. Conductor 7-shaped, sclerotized, with two large apophyses apically and a tooth-shaped apophysis basally. Tegulum translucent, slightly swollen. Embolus filiform, the tip with complex structure, the other structure coiled into two circles. Spermatic ducts can be seen through translucent tegulum.


Figure 15. Microdipoena shenyang sp. nov., male (A-C) and female (D-F) from Liaoning of China $\mathbf{A}$ habitus, dorsal $\mathbf{B}$ habitus, ventral $\mathbf{C}$ habitus, lateral $\mathbf{D}$ habitus, lateral $\mathbf{E}$ habitus, dorsal $\mathbf{F}$ habitus, ventral. Abbreviations: $F S=$ femoral spot, TS = tibial spine on male leg I. Scale bars: 0.50 mm .


Figure 16. Microdipoena shenyang sp. nov. from Liaoning of China A conductor, dorsal B cymbium, ventral C bulbus with conductor removed, prolateral $\mathbf{D}$ bulbus with conductor removed, retrolateral $\mathbf{E}$ embolic end, apical-lateral $\mathbf{F}$ male palp, prolateral $\mathbf{G}$ male palp retrolateral. Abbreviations: $\mathrm{CT}=$ cymbial tooth, $\mathrm{Cy}=$ cymbium, $\mathrm{CyC}=$ cymbial conductor, CyP = cymbial process, $\mathrm{E}=$ embolus, $\mathrm{Pa}=$ patella, $\mathrm{PC}=$ paracymbium, $\mathrm{SD}=$ spermatic duct, $\mathrm{T}=$ tegulum, $\mathrm{Ti}=$ tibia. Scale bars: $0.10 \mathrm{~mm}(\mathbf{A}, \mathbf{B}, \mathbf{E}-\mathbf{G}) ; 0.20 \mathrm{~mm}(\mathbf{C}, \mathbf{D})$.


Figure 17. Microdipoena shenyang sp. nov. from Liaoning of China A epigyne, ventral B vulva, ventral C vulva, dorsal Abbreviations: $C D=$ copulatory duct, $F D=$ fertilization duct, $S=$ spermathecal, $S p=$ scape. Scale bars: 0.10 mm

Female (holotype). Total length 1.25. Carapace 0.42 long, 0.46 wide, 0.42 high. Clypeus 0.08 high. Sternum 0.32 long, 0.28 wide. Abdomen 0.83 long, 0.83 wide, 0.96 high. Length of legs: I $1.20(0.28,0.12,0.24,0.26,0.30)$ II 1.08 ( $0.26,0.12,0.20,0.24,0.26$ ); III 1.04 ( $0.28,0.12,0.18,0.22,0.24$ ); IV 1.00 ( 0.28 , $0.12,0.16,0.20,0.24)$.

Somatic characters (Fig. 15D-F). Coloration: carapace pale brown. Ocular base black. Chelicera, endites, labium yellow; sternum yellow with two brown stripes. Legs yellow-brown. Abdomen silvery brown with multiple symmetrical yellow spots dorsally, black with multiple arched yellow stripes and spots ventrally. Prosoma: carapace nearly pear-shaped in dorsal view. Cephalic part slightly elevated. Sternum scutiform, slightly plump, covered in sparse setae. Legs: covered with setae and bristles. Femurs I and II with sclerotized femoral spot. Abdomen: nearly globose in dorsal view, covered with pale setae.

Epigyne (Fig. 17A-C): scape long, with thin folds, tip sclerotized. Copulatory duct short, the sclerotized part coiled in two circles, membranous part coiled under the spermathecae. Paired spermathecae semicircular, separated by $4 \times$ diameter. Fertilization ducts slightly sclerotized, originated from the lower edge of the spermathecae and bent anteriorly.

Distribution. China (Liaoning)

## Microdipoena thatitou sp. nov.

https://zoobank.org/2FDF9219-4F93-4C86-9D58-672A52EEDC1D
Fig. 18

Type material. Holotype $q$ (NHMSU-No.83) Laos: Champasak Province, Muang Bachieng, Ban Lak 35, That Itou ( $15^{\circ} 11.628^{\prime} \mathrm{N}, 106^{\circ} 06.105^{\prime} \mathrm{E} ; 810 \mathrm{~m}$ elev.), 26.VI.2009, PW. Jäger and S. Bayer leg.

Etymology. The specific name is derived from the type locality; noun in apposition.

Diagnosis. This new species can be distinguished from other congeners by the thickened, long S-shaped fertilization ducts and the entire membranous scape including distal end (Fig. 18D-F).

Description. Female (holotype): Total length 0.82. Carapace 0.24 long, 0.32 wide, 0.24 high. Clypeus 0.06 high. Sternum 0.23 long, 0.18 wide. Abdomen 0.58 long, 0.58 wide, 0.51 high. Length of legs: I $1.13(0.32,0.12,0.23,0.22$, 0.24 ); II 1.22 ( $0.24,0.12,0.20,0.18,0.20$ ); III 0.62 ( $0.20,0.08,0.14,0.10,0.10$ ); IV 0.84 ( $0.24,0.12,0.18,0.14,0.16$ ).

Somatic characters (Fig. 18A-C). Coloration: carapace yellow centrally, yellow-brown marginally. Ocular base black. Chelicera, endites yellow, labium yellow, sternum pale yellow with two symmetrical black stripes. Legs yel-low-brown. Abdomen silvery yellow dorsally, black with symmetrical white and yellow spots, silvery brown with two symmetrical silvery yellow stripes ventrally. Prosoma: carapace nearly pear-shaped in dorsal view. Cephalic part unelevated. Sternum scutiform, covered with sparse setae. Legs: covered with setae and bristles. Femurs I and II with sclerotized femoral spot. Abdomen: nearly spherical, covered with sparse pale setae.


Figure 18. Microdipoena thatitou sp. nov., female from Laos A habitus, dorsal B habitus, ventral C habitus, lateral Depigyne, ventral E vulva, ventral F vulva, dorsal. Abbreviations: $C D=$ copulatory duct, $F D=$ fertilization duct, $S=$ spermathecal, Sp = Scape. Scale bars: $0.50 \mathrm{~mm}(\mathbf{A}-\mathbf{C}), 0.10 \mathrm{~mm}(\mathbf{D}-F)$.

Epigyne (Fig. 18D-F): the structure can be seen through the cuticle. Scape membranous, soft, without weakly sclerotized distal end. Copulatory duct membranous, coiled under the spermathecae. Paired spermathecae oval. Fertilization ducts long, thick, and sclerotized, originated from the lower edge of the spermathecae, the composite structure of fertilization ducts and spermathecae S-shaped.

Male. Unknown.
Distribution. Laos (Champasak)

## Microdipoena yinae (Lin \& Li, 2013)

Figs 19, 20

Mysmenella yinae Lin \& Li, 2013: 470.
Microdipoena yinae: Lopardo and Hormiga 2015: 783.

Type material. Holotype: đ (NHMSU) and Paratypes 14§ 65 ${ }^{\wedge}$ (NHMSU), China: Sichuan Province, Jiuzhaigou County, Dalu Town, the moss under the forest shrub in the side of Heishui River ( $33^{\circ} 33.966^{\prime} \mathrm{N}, 103^{\circ} 40.243^{\prime} \mathrm{E} ; 2495 \mathrm{~m}$ elev.), 28.VI.2011, Y. Lin leg.; 6 31 Q (NHMSU), China: Sichuan Province, Jiuzhaigou County, Dalu Town, the forest shrub, at a fork in the road of Dalu Town and Zoige County ( $33^{\circ} 34.237^{\prime} \mathrm{N}, 103^{\circ} 40.166^{\prime} \mathrm{E}$; 2462 m elev.), 28.VI.2011, Y. Lin leg. Examined.

Diagnosis. See diagnosis for $M$. gongi and $M$. huisun sp. nov.
Description. See Figs 19, 20 and Lin et al. 2013: 470.
Distribution. China (Sichuan).

## Microdipoena zhulin sp. nov.

https://zoobank.org/93F9A4B5-4500-4522-A02C-27433E8D3F84
Figs 21-23

Type material. Holotype $q$ and paratype $1 \circlearrowleft 3 q$ (NHMSU-GX02) China: Guangxi Zhuang Autonomous Region, Guilin City, Lingui County, Ertang Township, Yanmendi Village, bamboo forest ( $25^{\circ} 12.892^{\prime} \mathrm{N}, 100^{\circ} 12.204^{\prime} \mathrm{E}$; 165 m elev.), 19.VII.2013, H. Zhao leg.

Etymology. The specific name is derived from the Chinese pinyin for bamboo forest (zhú lín), refers to this species living in this habitats; noun in apposition.

Diagnosis. This new species can be distinguished from other congeners by a combination of the following features of the copulatory organ: having a cymbial tooth near the distal edge of cymbium, the conductor with a small, thumbshaped, upper process and a right, broad, lower process, the spherical spermathecae separated by ca $2.6 \times$ their diameter, the scape of uniform width from base to end, the vulva with two smooth, transparent membranes (probably part of copulatory ducts) (Figs 22B, D, 23D).

Description. Male: Total length 1.27. Carapace 0.31 long, 0.44 wide, 0.40 high. Clypeus 0.06 high. Sternum 0.31 long, 0.24 wide. Abdomen 0.96 long, 0.87 wide, 0.93 high. Length of legs: I $0.87(0.26,0.10,0.23,0.15,0.13)$; II $0.64(0.10,0.08,0.18$, $0.13,0.15)$; III 0.49 ( $0.18,0.05,0.10,0.08,0.08$ ); IV 0.64 ( $0.18,0.08,0.18,0.1,0.10$ ).


Figure 19. Microdipoena yinae Lin \& Li, 2013, from Sichuan of China A male habitus, dorsal B male habitus, ventral C male habitus, lateral $\mathbf{D}$ female habitus, dorsal $\mathbf{E}$ female habitus, ventral $\mathbf{F}$ female habitus, lateral $\mathbf{G}$ epigyne, ventral $\mathbf{H}$ vulva, dorsal. $\mathrm{CD}=$ copulatory duct, $\mathrm{FD}=$ fertilization duct, $\mathrm{S}=$ spermathecal, $\mathrm{Sp}=$ scape. Scale bars: 0.20 mm .

Somatic characters (Fig. 21A-C). Coloration: carapace silvery yellow centrally, black marginally. Chelicera, endites, labium yellow; sternum yellow with two orange stripes. Legs yellow-black. Abdomen black with large white spots dorsally, yellow with black and white spots ventrally. Prosoma: carapace nearly hexagonal in dorsal view and peak-shaped in lateral view. Cephalic part elevated and flat. Sternum scutiform, plump, covered with sparse setae. Legs: covered with setae. Mating clasper on metatarsus I, two strong spines on tibia I. Abdomen: nearly globose in dorsal view, covered in black setae.


Figure 20. Microdipoena yinae Lin \& Li, 2013, from Sichuan of China A male palp, apical B bulbus with conductor removed, dorsolateral $\mathbf{C}$ conductor, dorsal $\mathbf{D}$ male palp, prolateral $\mathbf{E}$ male palp, retrolateral. Abbreviations: $\mathrm{Cy}=$ cymbium, $\mathrm{CyC}=$ cymbial conductor, $\mathrm{CyP}=$ cymbial process, $\mathrm{E}=$ embolus, $\mathrm{Pa}=$ patella, $\mathrm{PC}=$ paracymbium, $\mathrm{SD}=$ spermatic duct, $\mathrm{T}=$ tegulum, $\mathrm{Ti}=$ tibia. Scale bars: 0.10 mm .


Figure 21. Microdipoena zhulin sp. nov., male (A, B) and female (D-F) from Guangxi of China $\mathbf{A}$ habitus, dorsal $\mathbf{B}$ habitus, ventral $\mathbf{C}$ habitus, lateral $\mathbf{D}$ habitus, lateral $\mathbf{E}$ habitus, dorsal $\mathbf{F}$ habitus, ventral. Abbr.: $\mathrm{FS}=$ femoral spot. Scale bars: 0.50 mm .


Figure 22. Microdipoena zhulin sp. nov. from Guangxi of China A palp, apical B conductor, dorsal C cymbium, apical D cymbium, prolateral E left tibia I and metatarsus I, prolateral $\mathbf{F}$ palp, prolateral $\mathbf{G}$ palp, ventral $\mathbf{H}$ palp, retrolateral. Abbreviations: $\mathrm{CT}=$ cymbial tooth, $\mathrm{Cy}=$ cymbium, $\mathrm{CyC}=$ cymbial conductor, $\mathrm{CyP}=$ cymbial process, $\mathrm{E}=$ embolus, $\mathrm{MC}=$ mating clasper, $\mathrm{Pa}=$ patella, $\mathrm{PC}=$ paracymbium, $\mathrm{T}=$ tegulum, $\mathrm{Ti}=$ tibia, $\mathrm{TS}=$ tibial spine. Scale bars: 0.10 mm .


Figure 23. Microdipoena zhulin sp. nov. from Guangxi of China A epigyne, ventral B epigyne, lateral C vulva, ventral D vulva, dorsal. Abbreviations: $C D=$ copulatory duct, $F D=$ fertilization duct, $S=$ spermathecal, $S p=$ scape. Scale bars: 0.10 mm .

Palp (Fig. 22A, F-H): Cymbium translucent, the tip end of specialized as cymbial conductor, and the other end forming a large cymbial process, a small, sclerotized, cymbial tooth on the outward side of cymbial conductor. Paracymbium finger-shaped, with long setae. Conductor slightly sclerotized, with three large apophyses apically and an arched apophysis basally. Tegulum translucent, slightly swollen. Embolus long, coiled into two circles, the tip coiled and folded into a complex structure. Spermatic ducts can be seen through translucent tegulum.

Female (holotype). Total length 1.58. Carapace 0.65 long, 0.75 wide, 0.62 high. Clypeus 0.12 high. Sternum 0.47 long, 0.47 wide. Abdomen 0.93 long, 0.84 wide, 0.92 high. Length of legs: I $1.61(0.64,0.18,0.26,0.20,0.33)$; II $1.39(0.49,0.18,0.26$, $0.18,0.28$ ); III $0.86(0.28,0.10,0.18,0.14,0.16)$; IV $1.04(0.36,0.10,0.18,0.18,0.22)$.

Somatic characters (Fig. 21D-F). Coloration: carapace pale yellow centrally, brown marginally. Ocular base black. Chelicera, endites, labium, and sternum yellow. Legs yellow-brown. Abdomen nearly white dorsally, black with multiple white and yellow spots ventrally. Prosoma: carapace nearly pear-shaped in dorsal view. Cephalic part slightly elevated. Sternum scutiform, slightly plump, covered in sparse setae. Legs: covered with setae and bristles. Femurs I and II with sclerotized femoral spot. Abdomen: nearly globose in dorsal view, covered with black setae.

Epigyne (Fig. 23A-D): scape long, with wide folds, tip sclerotized. Copulatory duct membranous, coiled under the spermathecae. Fertilization ducts slightly sclerotized, originating from the ventral side of the epigyne and bent anteriorly. Paired spermathecae nearly round, separated by nearly double their diameter.

Distribution. China (Guangxi).

## Discussion

In this paper, we describe a group of species of the genus Microdipoena that are mainly native to Eurasia. The morphological characteristics of copulatory organs were compared between multiple congeneric species. Some of the diagnostic features they shared were verified, and can be distinguished from those of other genera (cf. male with two or three tibial spines on the leg I; male palp with a paracymbium, distal part of the embolus coiled and distorted into a complex structure in most species; Lopardo and Hormiga 2015). To test whether our taxonomic decisions and their classification status are correct, we also conducted phylogenetic analyses based on molecular evidence for ten named and five undescribed Microdipoena species. Our phylogenetic analysis shows that the monophyly of this genus is valid and these taxonomic judgments proposed by us in this study are correct. However, the male characters of copulatory organs of three species are unknown due to inadequate sampling ( $M$. huisun sp. nov., M. lisu sp. nov., and $M$. thatitou sp. nov.).

According to the reported distribution records, the genus is mainly distributed in the continents of Asia and Africa and nearby islands. Most species of the genus are endemic, some of which have multiple distribution sites (M. elsae, $M$. nyungwe, $M$. samoensis), and a few may have expanded distribution ranges as a result of introduction ( $M$. guttata and $M$. jobi). The origin and diffusion history of Microdipoena are questions worthy of further discussion. Faunal surveys and diversity studies of this genus are a prerequisite for answering these questions, but much work remains to be done.

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## Additional information

## Conflict of interest

The authors have declared that no competing interests exist.

## Ethical statement

No ethical statement was reported.

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## Author contributions

Conceptualization: YL. Writing - original draft: QZ.

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## Data availability

All of the data that support the findings of this study are available in the main text.

## References

Baert L (1984a) Mysmenidae and Hadrotarsidae from the Neotropical Guaraní zoogeographical province (Paraguay and south Brasil) (Araneae). Revue Suisse de Zoologie 91: 603-616. https://doi.org/10.5962/bhl.part. 81569
Baert L (1984b) Spiders (Araneae) from Papua New Guinea. IV. Ochyroceratidae, Telemidae, Hadrotarsidae and Mysmenidae. Indo-Malayan Zoology 2: 225-244.
Baert L (1985) Telemidae, Mysmenidae and Ochyroceratidae from Cameroon (Araneae): Scientific report of the Belgian Mount Cameroon Expeditions 1981 and 1983 (no. 13). Biologisch Jaarboek Dodonaea 53: 44-57.

Baert L (1986) Mysmenidae from the Comoro Islands (Araneae). Revue Zoologique Africaine 100: 265-267.

Baert L (1989) Mysmenidae from Rwanda (Araneae). Revue Zoologique Africaine 103: 29-33.
Balczun C, Bunse A, Hahn D, Bennoun P, Nickelsen J, Kück U (2005) Two adjacent nuclear genes are required for functional complementation of a chloroplast trans-splicing mutant from Chlamydomonas reinhardtii. The Plant Journal 43(5): 636-648. https://doi.org/10.1111/j.1365-313X.2005.02478.x
Banks N (1895) A list of the spiders of Long Island; with descriptions of new species. Journal of the New York Entomological Society 3: 76-93.
Bishop SC, Crosby CR (1926) Notes on the spiders of the southeastern United States with descriptions of new species. Journal of the Elisha Mitchell Scientific Society 41: 163-212. [pl. 20-25]

Brignoli PM (1980) On few Mysmenidae from the Oriental and Australian regions (Araneae). Revue Suisse de Zoologie 87: 727-738. https://doi.org/10.5962/bhl.part. 85542
Colgan DJ, McLauchlan A, Wilson GDF, Livingston SP, Edgecombe GD, Macaranas J, Cassis G, Gray MR (1998) Histone H3 and U2 snRNA DNA sequences and arthropod molecular evolution. Australian Journal of Zoology 46(5): 419-437. https://doi. org/10.1071/ZO98048
Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R (1994) DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. Molecular Marine Biology and Biotechnology 3: 294-299.
Forster RR (1959) The spiders of the family Symphytognathidae. Transactions and Proceedings of the Royal Society of New Zealand 86: 269-329.
Forster RR, Platnick NI (1977) A review of the spider family Symphytognathidae (Arachnida, Araneae). American Museum Novitates 2619: 1-29.
Giribet G, Carranza S, Baguñà J, Riutort M, Ribera C (1996) First molecular evidence for the existence of a Tardigrada + Arthropoda clade. Molecular Biology and Evolution 13(1): 76-84. https://doi.org/10.1093/oxfordjournals.molbev.a025573
Gruia M (1977) Sur quelques Theridiidae et Symphytognathidae (Aranea) recueillis par la deuxième expédition biospéologique cubano-roumaine à Cuba. Résultats des Expéditions Biospéologiques Cubano-Roumaines à Cuba 2: 159-163.
Hall TA (1999) BioEdit: A user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symposium Series 41: 95-98.
Hormiga G, Arnedo M, Gillespie RG (2003) Speciation on a conveyor belt: Sequential colonization of the Hawaiian islands by Orsonwelles spiders (Araneae, Linyphiidae). Systematic Biology 52(1): 70-88. https://doi.org/10.1080/10635150390132786
Khmelik VV, Kozub D, Glazunov A (2006) Helicon Focus 3.10.3. http://www.heliconsoft. com/heliconfocus.html [Accessed on 10 Sept 2018]
Kraus O (1967) Zur Spinnenfauna Deutschlands, II. Mysmena jobi n. sp, eine Symphytognathide in Mitteleuropa (Arachnida: Araneae: Symphytognathidae). Senckenbergiana Biologica 48: 387-399.
Lanfear R, Frandsen PB, Wright AM, Senfeld T, Calcott B (2016) PartitionFinder 2: New Methods for Selecting Partitioned Models of Evolution for Molecular and Morphological Phylogenetic Analyses. Molecular Biology and Evolution 34(3): 772-773. https://doi. org/10.1093/molbev/msw260
Levi HW (1956) The spider genus Mysmena in the Americas (Araneae, Theridiidae). American Museum Novitates 1801: 1-13.
Lin Y, Li S (2008) Mysmenid spiders of China (Araneae: Mysmenidae). Annales Zoologici, Warszawa 58(3): 487-520. https://doi.org/10.3161/000345408X364337
Lin Y, Li S (2013) Five new minute orb-weaving spiders of the family Mysmenidae from China (Araneae). Zootaxa 3670(4): 449-481. https://doi.org/10.11646/zootaxa.3670.4.3
Logunov DV (2022) John Alan Murphy (1922-2021) and his contribution to arachnology. Arachnology 19(Special Issue): 77-103. https://doi.org/10.13156/arac.2022.19. sp1.77
Lopardo L, Hormiga G (2015) Out of the twilight zone: Phylogeny and evolutionary morphology of the orb-weaving spider family Mysmenidae, with a focus on spinneret spigot morphology in symphytognathoids (Araneae, Araneoidea). Zoological Journal of the Linnean Society 173(3): 527-786. https://doi.org/10.1111/zoj. 12199
Lopardo L, Giribet G, Hormiga G (2011) Morphology to the rescue: molecular data and the signal of morphological characters in combined phylogenetic analyses - a case
study from mysmenid spiders (Araneae, Mysmenidae), with comments on the evolution of web architecture. Cladistics 27(3): 278-330. [\& supplementary material] https://doi.org/10.1111/j.1096-0031.2010.00332.x

Marples BJ(1955)Spiders from WesternSamoa. Journal of the LinneanSociety of London, Zoology 42(287): 453-504. [pls 56-59] https://doi.org/10.1111/j.1096-3642.1955. tb02217.x

Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for inference of large phylogenetic trees. 2010 Gateway Computing Environments Workshop, GCE 2010, 8 pp. https://doi.org/10.1109/GCE.2010.5676129
Naumova M, Blagoev G, Dimitrov D, Lazarov S, Deltshev C (2017) New data on the spider fauna (Arachnida: Araneae) of Bulgaria. Acta Zoologica Bulgarica 69(4): 477-481.
Ono H (2007) Eight new species of the families Hahniidae, Theridiidae, Linyphiidae and Anapidae (Arachnida, Araneae) from Japan. Bulletin of the National Museum of Nature and Science Tokyo (A) 33(4): 153-173.
Rambaut A, Drummond AJ, Xie D, Baele G, Suchard MA (2018) Posterior summarization in Bayesian phylogenetics using Tracer 1.7. Systematic Biology 67(5): 901-904. https://doi.org/10.1093/sysbio/syy032
Rix MG, Harvey MS, Roberts D (2008) Molecular phylogenetics of the spider family Micropholcommatidae (Arachnida: Araneae) using nuclear rRNA genes (18S and 28S). Molecular Phylogenetics and Evolution 46(3): 1031-1048. https://doi.org/10.1016/j. ympev.2007.11.001
Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA, Huelsenbeck JP (2012) MrBayes 3.2: Efficient Bayesian phylogenetic inference and model choice across a large model space. Systematic Biology 61(3): 539-542. https://doi.org/10.1093/sysbio/sys029
Saaristo MI (1978) Spiders (Arachnida, Araneae) from the Seychelle islands, with notes on taxonomy. Annales Zoologici Fennici 15: 99-126.
Simon E (1895) Etudes arachnologiques. 26e. XLI. Descriptions d'espèces et de genres nouveaux de l'ordre des Araneae. Annales de la Société Entomologique de France 64: 131-160.

Vaidya G, Lohman DJ, Meier R (2011) SequenceMatrix: Concatenation software for the fast assembly of multi-gene datasets with character set and codon information. Cladistics 27(2): 171-180. https://doi.org/10.1111/j.1096-0031.2010.00329.x
WSC (2023) World Spider Catalog. Version 24.5. Natural History Museum Bern. http://wsc.nmbe.ch [Accessed 8 August 2023]

Yin C, Peng X, Bao Y (2004) A new species of the genus Mysmenella from China (Araneae, Mysmenidae). Acta Zootaxonomica Sinica 29: 80-82.
Zhang D, Gao F, Jakovlić I, Zou H, Zhang J, Li WX, Wang GT (2020) PhyloSuite: An integrated and scalable desktop platform for streamlined molecular sequence data management and evolutionary phylogenetics studies. Molecular Ecology Resources 20(1): 348-355. https://doi.org/10.1111/1755-0998.13096
Zhang Q, Li S, Lin Y (2022) Taxonomic study on Mysmenidae spiders (Mysmenidae, Araneae) from Xishuangbanna of Yunnan, China. ZooKeys 1124: 59-108. https://doi. org/10.3897/zookeys.1124.85952

