The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera): a unique character condition in Myrmeleontoidea

Vladimir N. Makarkin¹², Qiang Yang¹⁴, Chaofan Shi¹⁵, Dong Ren¹¹

¹ College of Life Sciences, Capital Normal University, 105 Xisanhuanbeilu, Haidian District, Beijing, 100048, China ² Institute of Biology and Soil Sciences, Far Eastern Branch of the Russian Academy of Sciences, 100 let Vladivostoku Avenue 159, Vladivostok, 690022, Russia

† http://zoobank.org/38B237B6-2E03-40E8-B8EB-A895C91ADDD9 ‡ http://zoobank.org/82D4C606-5047-457D-B7DF-21BB8F9FD1BE § http://zoobank.org/A6F5614C-CBAA-44EA-A9C3-7A03233057E0 | http://zoobank.org/D507ABBD-6BA6-43C8-A1D5-377409BD3049

Corresponding author: Dong Ren (rendong@mail.cnu.edu.cn)

Academic editor: S. Winterton  |  Received 2 May 2013  |  Accepted 5 August 2013  |  Published 20 August 2013

http://zoobank.org/AD2A3F47-26BA-422C-9062-583DFC6764D

Citation: Makarkin VN, Yang Q, Shi C, Ren D (2013) The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera): a unique character condition in Myrmeleontoidea. ZooKeys 325: 1–20. doi: 10.3897/zookeys.325.5453

Abstract

A well-developed recurrent veinlet is found in the forewing of two species of Nymphidae from the Middle Jurassic locality of Daohugou (Inner Mongolia, China), Liminympha makarkini Ren & Engel and Daonymphes bisulca gen. et sp. n. This is the first record of this trait in the clade comprised of the superfamilies Myrmeleontoidea and Chrysopoidea. We interpret the recurrent veinlet in these species as a remnant of the condition present more basally in the psychopsoid + ithonoid + chrysopoid + myrmeleontooid clade (i.e., as a plesiomorphy). Other venational character states of D. bisulca of interest include the configuration of subcosta anterior (ScA), which is very similar to that of extant Nymphidae. We consider the short ScA terminating on ScP to be an autapomorphy of Neuroptera.

Keywords

Neuroptera, Nymphidae, recurrent veinlet, subcosta anterior, Daohugou, Middle Jurassic

Copyright Vladimir N. Makarkin et al. This is an open access article distributed under the terms of the Creative Commons Attribution License 3.0 (CC-BY), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.
Introduction

Understanding the evolution of particular morphological characters during the broad historical development of higher insect taxa is a very interesting and important question. In fact, the phylogeny of a taxon may be realistic only if the evolutionary transformations of all available characters are correctly interpreted (Hennig 1981; Bechly 2000; Kukalová-Peck 2008).

The forewing humeral veinlet (i.e., the proximal-most veinlet of ScP) in all Neuroptera (and generally in all Neoptera) is plesiomorphically a simple, crossvein-like. However, in some families of Neuroptera it is curved, obviously directed towards wing base (i.e., recurrent), and often profusely branched. This condition (the recurrent veinlet for short) is one of the significant informative venational characters found within this order. It is characteristic of several families and subfamilies, the vast majority of which have the costal space basally dilated (see below). However, it was hitherto not found in any of numerous species of the clade comprised the superfamilies Myrmeleontoidea (Nymphidae, Nemopteridae, Ascalaphidae, Myrmeleontidae, Palaeoleontidae and Babinskaiidae) and Chrysopoidea (Chrysopidae, Mesochrysopidae and Ascalochrysidae) (Yang et al. 2012: fig. 32). The costal space in the vast majority of species of this clade is basally narrowed.

Nymphidae are generally accepted to be the most basal family in the superfamilies Myrmeleontoidea, and the only myrmeleontoid family known from the Jurassic. In general, 18 fossil (named) species are known in the family (Krüger 1923; Carpenter 1929; Panfilov 1980; Makarkin 1990a, b; Ponomarenko 1992a; Martins-Neto 2005; Menon et al. 2005; Ren and Engel 2007; Engel and Grimaldi 2008; Archibald et al. 2009; Jepson et al. 2012; Shi et al. in press). Here, we report the occurrence of a recurrent veinlet in the two oldest known specimens of Nymphidae, recovered from the Middle Jurassic Daohugou locality, China: Liminympha makarkini Ren & Engel, 2007; and a new species, described here. Three other nymphid specimens are known from that locality; the recurrent veinlet appears to be absent in one (but its humeral area is poorly preserved), and the basal portion of wings is missing in other two (Shi et al. in press). In this paper, we re-describe L. makarkini, describe a new genus and species, and discuss the occurrence of the recurrent veinlet in neuropteran taxa and its possible phylogenetic implications. Also, we discuss the presence of the distinct subcosta anterior in Daonymphes bisulca gen. et sp. n. We consider ScA terminating on ScP to be an autapomorphy of Neuroptera.

Material and methods

This paper is based on two specimens collected from the Daohugou locality and housed in the Key Laboratory of Insect Evolution and Environmental Changes, College of Life Sciences, Capital Normal University, Beijing, China (CN UB; Dong Ren,
The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)... 3

The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)...

3
curator). Daohugou Village is situated in Shantou Township, Ningcheng County, Inner Mongolia, China. The insect-bearing beds of the Daohugou locality are considered to belong to the Jiulongshan Formation, and are dated as Bathonian, Middle Jurassic (Gao and Ren 2006) or Bathonian to Callovian (Middle to late Middle Jurassic) (Yang et al. 2012).

The specimens were examined under a Leica MZ12.5 dissecting microscope; line drawings were prepared with CorelDraw 12 graphics software and Adobe Illustrator CS5 with the aid of Adobe Photoshop CS3; photographed by a Nikon SMZ1000 stereomicroscope; the whole specimens were photographed by Nikon D100 Digital Camera.

Terminology and abbreviations. Wing venation terminology in general follows Kukalová-Peck and Lawrence (2004), except that we treat the median vein as in Yang et al. (2012). Terminology of wing spaces and details of the venation (e.g., subcostal veinlets) follows Oswald (1993).

Venational abbreviations. AA, analis anterior; AP, analis posterior; Cu, cubitus; CuA, cubitus anterior; CuP, cubitus posterior; M, media; MA, media anterior; MP, media posterior; R, radius; RA, radius anterior; RP, radius posterior; RP1, proximalmost branch of RP; rv, recurrent veinlet; ScP, subcosta posterior.

Systematic paleontology

Order Neuroptera Linnaeus, 1758
Family Nymphidae Rambur, 1842

Genus Daonymphes gen. n.
http://zoobank.org/17B6B9EB-D7C9-46A1-A8F9-0F5631FDA7B7
http://species-id.net/wiki/Daonymphes

Type and only species. Daonymphes bisulca sp. n.

Diagnosis. Forewing broad proximally [strongly narrowed proximally in Liminympha]; vast majority of proximal subcostal veinlets forked [simple in all other Mesozoic genera]; crossveins between branches of MP absent [present in Nymphites Haase, 1890, Mesonymphes Carpenter, 1929]; CuP space broad, nearly two times as wide as intracubital space [nearly as wide as intracubital space in Liminympha].

Etymology. From Daohugou, the locality of the type species, and Nymphes, a genus-group name. Gender: feminine.

Remarks. This genus is most closely related to four other Mesozoic genera (Nymphites, Mesonymphes, Liminympha and Sialium Westwood, 1854) but clearly distinguished from these as indicated in the diagnosis. The latter genus is only known from single hind wing from the early Berriasian of Purbeck, England. In general, Nymphidae occur very rare in the Jurassic, only six specimens are known. The monotypic genera Liminympha (Middle Jurassic [Bathonian/Callovian] of Dao-
hugou, China) and *Mesonymphys* (Late Jurassic [Tithonian] of Solnhofen, Germany) are represented by almost complete specimens possessing both fore- and hind wings. The type species of *Nymphites* from Solnhofen is represented by only two incomplete hind wings. Two species from Daohugou are assigned to this genus: one species (two specimens) have almost complete fore- and hind wings; the other only hind wings (Shi et al. in press).

*Daonymphys bisulca* sp. n.
http://zoobank.org/61D8C637-96AD-4D41-BEF6-270FE5F3A6BA
http://species-id.net/wiki/Daonymphys_bisulca
Figs 1, 2, 6A

**Diagnosis.** As for the genus.

**Description.** Forewing 29 mm long as preserved (estimated complete length about 40 mm), 11.5 mm wide. Costal space relatively broad for entire length, narrowed basally. Humeral veinlet recurrent, with one short branch. Subcostal veinlets somewhat curved, mostly forked once or twice, shallowly, few deeply; several proximal veinlets simple. ScA well developed, terminating on ScP within humeral area. Subcostal space narrow, with several scarce crossveins. RA space relatively narrow, slightly narrowed towards wing apex, with relatively numerous crossveins. RP with nine preserved branches, becoming more closely spaced towards wing apex. Crossveins rather numerous over entire radial space, irregularly spaced. M forked distal of origin of RP. MA incomplete, simple for entire preserved length. MP distally pectinate, with six preserved branches, most of these forked once or twice; no crossveins detected between branches. Cu divided into CuA, CuP rather far from wing base. CuA distally pectinate, with six preserved branches, all forked once or twice; one series of crossveins between branches of CuA, continued CuP (‘pseudo-CuP’). CuP long, strongly pectinately branched with nine long branches; of four proximal-most branches, two forked; all distal branches dichotomously forked; no crossveins detected. Eleven crossveins between CuA, CuP rather irregularly spaced. AA3+4 and its posterior branch deeply forked. Basal crossvein between CuP, AA3+4 very short; distal crossvein strongly oblique. AP1+2 deeply forked. AP3+4 not preserved. Membrane around one crossvein between MA, MP near MP and three crossveins between CuA, CuP near CuA broadly heavily shaded; most crossveins between RA, RP apparently broadly shaded, pale fuscous.

**Material.** Holotype CNU-NEU-NN2011119, deposited in CNUB; an incomplete forewing.

**Type locality and horizon.** Daohugou Village, Shantou Township, Ningcheng County, Inner Mongolia, China. Jiulongshan Formation, Middle Jurassic.

**Etymology.** From the Latin *bisulcus,-a,-um*, forked, divided into two parts, in reference to its forked subcostal veinlets.
The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)...

Figure 1. *Daonymphes bisulca* gen. et sp. n. Holotype CNU-NEU-NN2011119 as preserved. Scale bar = 10 mm.

Figure 2. *Daonymphes bisulca* gen. et sp. n. Forewing venation of the holotype CNU-NEU-NN2011119. Scale bar = 10 mm.

**Liminympha Ren & Engel, 2007**
http://species-id.net/wiki/Liminympha

*Liminympha* Ren & Engel, 2007: 212.

**Type and only species.** *Liminympha makarkini* Ren & Engel, 2007, by original designation.

**Diagnosis (revised).** Forewing strongly narrowed proximally [broad proximally in *Nymphites, Daonymphes* gen. n. and *Mesonymphes*]; all veinlets of ScP simple [at least distal veinlets forked once or twice in *Nymphites, Daonymphes* gen. n.]; CuP space narrow, nearly as wide as intracubital space [nearly two times as wide as intracubital
space in *Nymphites, Daonymphes* gen. n. and *Mesonymphes*. Hind wing MP with four pectinate branches [eight in *Sialium*]; branches of CuA not connected by crossveins [at least proximal branches of CuA connected by crossveins in *Nymphites*]; anterior branch of CuP simple [deeply forked in *Nymphites* and *Mesonymphes*].

**Liminympha makarkini** Ren & Engel, 2007

http://species-id.net/wiki/Liminympha_makarkini

Figs 3–5, 6B

**Liminympha makarkini** Ren & Engel, 2007: 212, figs 1–3; Engel and Grimaldi 2008: 9; Yang et al. 2010: 177.

**Redescription.** Body (metathorax, abdomen) poorly, fragmentarily preserved. First abdominal tergite rather long; distally with distinct transverse suture, probably heavily

**Figure 3.** *Liminympha makarkini* Ren & Engel, 2007. Holotype CNU-NEU-NN1999024 (counter-part) as preserved. Scale bar = 5 mm.
sclerotized; medially with mediolongitudinal short suture; portion of 1st tergite distal to transverse suture (‘Transversalnaht’ of Achtelig 1975) very narrow (Fig. 4). Second tergite nearly as wide as long. Other tergites indistinct.

Forewing elongate, narrowed in proximal portion, most dilated at distal 3/4 length; about 30 mm long, 8 mm wide. Costal space narrow, basally narrowed, distally dilated. All preserved subcostal veinlets simple, strongly oblique distally; veinlets of ScP+RA dichotomously branched. Humeral veinlet rather strongly recurrent, branching not detected (Fig. 6B). Subcostal space very narrow; two crossveins in distal part detected, others possible. RA spaces slightly narrower than costal space, narrowed towards apex; crossveins irregularly spaced for entire preserved portion. RP originates rather near wing base, with about 16 branches; RP1 originates far from origin of RP; at least four proximal-most branches widely spaced, more distal branches closely spaced. In right forewing, RP2 terminated at RP1; RP3, RP4 fused for short distance (probably aberrations). Crossveins between branches of RP very scarce, restricted to area between RP1 to RP5. M appears fused with R basally; forked at nearly equal distance from origin of RP, origin of RP1. MA long, slightly arched, distally few-branched. MP long, its anterior trace (stem of MP) nearly straight before terminal branching; with five long distal branches, quite strongly inclined. Crossveins between R/RP and MA, MA and
MP irregularly spaced, arranged differently in right, left wings. Cu dividing into CuA, CuP rather near to wing base. CuA long, smoothly curved anteriorly, pectinate with four long branches, each dichotomously branched. Between branches of CuA at last three crossveins forming gradate series continued CuP (‘pseudo-CuP’). CuP long, pectinately branched, with seven-eight rather short branches, most simple. AA3+4 rather

Figure 5. *Liminympha makarkini* Ren & Engel, 2007. Wing venation of the holotype CNU-NEUNN1999024 **A** right forewing **B** left forewing **C** right hind wing **D** left hind wing. Scale bar = 5 mm (all to scale).
The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)...

Figure 6. Basal portion of forewings of the Middle Jurassic Nymphidae showing the recurrent veinlet (rv) and the anterior subcosta (ScA) A the holotype of Daonymphes bisulca gen. et sp. n., CNU-NEU-NN2011119 B the holotype of Liminympha makarkini Ren & Engel, 2007, CNU-NEU-NN1999024. Both wetted with ethanol. Scale bar = 1 mm.
short; stem simple, with one or two simple branches. AP1+2 incompletely preserved, probably simple. AP3+4 not preserved. One dark rather big spot in distal portion of radial space might be present (apical half of other wing not preserved).

Hind wing similar in shape to forewing, slightly narrower; about 28 mm long, 7 mm wide. Costal space narrow, dilated distally. Subcostal veinlets simple, becoming more oblique, closely spaced, curved towards apex. Humeral veinlet bent to base, nearly straight, not branched. Subcostal space narrow; three crossveins detected, others possible. ScP, RA fused far from wing apex; preserved veinlets of ScP+RA long, dichotomously forked; crossveins between them not detected. RA space basally as wide as costal space, narrowed towards apex; crossveins rather regularly spaced for entire preserved portion (right wing). RP originated near wing base, with about 15 branches; RP1 originated far from origin of RP (but closer to wing base than in forewing); five proximal-most branches widely spaced, other (distal) branches closely spaced. Crossveins between branches of RP very scarce, restricted to area between RP1 to RP4. Origin of M and its fork not preserved. MA long, nearly straight, distally dichotomously branched. MP long, its anterior trace (stem of MP) slightly incurved, with four long distal branches, quite strongly inclined. Origin of Cu and its dividing into CuA, CuP not present. CuA long, pectinate, with 15 branches, which proximally simple, distally once or twice forked. CuP short, deeply forked. Distal crossvein between CuA, CuP rather short connecting CuA, anterior branch of CuP fork. Anal veins not preserved. Crossveins between R/RP and MA, MA and MP, MP and CuA poorly preserved, irregularly spaced, arranged differently in right and left wings; crossveins between branches of MP, CuA absent.

Material. Holotype CNU-NEU-NN1999024 (part, counterpart), deposited in CNUB; an incomplete specimen.

Type locality and horizon. Daohugou Village, Shantou Township, Ningcheng County, Inner Mongolia, China. Jiulongshan Formation, Middle Jurassic.

Remarks. This redescription is only based on the counterpart of the holotype.

Discussion

Recurrent veinlet

The recurrent veinlet is known in the following extant taxa of Neuroptera: all species of Psychopsidae and Ithonidae (including Polystoechotidae and Rapismatidae), the vast majority of Hemerobiidae; rarely in Berothidae (including Rhachiberothinae) and Mantispidae (only Symphrasinae).

The recurrent veinlet has not been detected in any Permian Neuroptera (Permithonidae, Archeosmylidae) and the majority of Triassic taxa represented by these two families, and ‘Mesoberothidae’; however the basal portion of the forewing in any reported specimen of the latter is poorly preserved (e.g., Riek 1955; Lambkin 1988; Vilesov and Novokshonov 1994; Vilesov 1995; Novokshonov 1996; Jell 2004; VM,
The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)... pers. obs.). In Triassic Neuroptera, the recurrent veinlet is only found in those taxa that have the costal space strongly dilated basally, i.e., the psychopsoids (Psychopsidae, Osmylopsychopidae, and broad-winged taxa whose family affinities are not yet clear). These are: (1) several undescribed Osmylopsychopidae from the Ladinian/Carnian Madygen Formation (Kyrgyzstan) (see Shcherbakov 2008: fig. 6); (2) *Triassopsychops superb*us Tillyard, 1922: fig. 89; (3) *Archesychops triassicus* Tillyard, 1919: fig. 27; (4) possibly in *Osmylopsychops spil*l*erae* Tillyard, 1923: fig. 93 (however see Lambkin 1992: fig. 2), and (5) *Petropsychops superb*us Riek, 1956: pl. 1, fig. 2 (see also Grimaldi and Engel 2005: fig. 9.15) (all from the Carnian Ipswich Series of Australia). Of these taxa, the recurrent veinlet is most clearly visible in one of the Osmylopsychopidae wings from the Madygen Formation, where it is strongly curved towards the wing base and has eight branches (VM, pers. obs.). In the Australian taxa, the recurrent veinlet is poorly preserved.

Of the Early Jurassic Neuroptera, the recurrent veinlet is detected in all psychopsoids whose forewing base is well preserved (e.g., Tillyard 1933: fig. 2; Ansorge 1996: fig. 52) and in Prohemerobiidae. The latter family, however, remains undefined; we treat it as containing only the type genus *Prohemerobius* Handlirsch, 1906 (probably paraphyletic), recorded from the Early Jurassic of Germany and England (Handlirsch 1906–1908, 1939; Bode 1953; Whalley 1988; Ponomarenko 1995) and the Late Jurassic of Mongolia (Khramov 2011). The majority of species in this genus have the basal portion of the forewing costal space missing or poorly preserved. Still, some species (including the type species *Prohemerobius dilaroide*is Handlirsch, 1906) possess the recurrent veinlet, configured similarly to that of the Middle Jurassic Nymphidae, i.e., with few (or one) simple branches (J. Ansorge, VM, pers. obs.). The wings of *Prohemerobius* are not of the psychopsoid type; they are small, relatively narrow with hemerobiid-like venation, and the costal space is narrow compared with those of psychopsoids. All known Early to Middle Jurassic Parakseneuridae possess a very well developed recurrent veinlet, along with up to 15 dichotomously forked branches (Yang et al. 2012: figs 1, 25, 30). All Jurassic to Cretaceous Osmylopsychopidae and Psychopsidae have the recurrent veinlet (e.g., Peng et al. 2010: fig. 3A; VM, pers. obs.). Some species of the Middle Jurassic to Early Cretaceous family Kalligrammatidae possess the well-developed recurrent veinlet (e.g., Yang et al. 2009: fig. 3, Yang et al. 2011: figs 2, 3A, Ponomarenko 2002: fig. 254); however, this condition is obviously absent in other species, even from the Middle Jurassic (VM, QY, pers. obs.).

The costal space in all numerous undescribed Ithonidae from the Middle Jurassic locality of Daohugou is basally narrowed, and therefore, the recurrent veinlet (where well developed) has short simple branches. Sometimes, the humeral veinlet is not branched and crossvein-like (VM, pers. obs.). Younger Ithonidae (Early Cretaceous to Recent) have the costal space usually broader basally and have a well developed recurrent veinlet (e.g., Archibald and Makarkin 2006: figs 5B, 8D, 9A; Makarkin and Archibald 2009: fig. 3; Makarkin et al. 2012: figs 3D, E). All species of the Middle Jurassic to Early Cretaceous subfamily Mesomantispinae possess a rather well developed recurrent veinlet (Makarkin 1996: fig. 1; Jepson et al. 2013: figs 1C, 2D, 4B).
A similar recurred veinlet is present in the extant subfamily Symphrasinae, the single fossil species of which is known from the middle Eocene of Messel, Germany (Wedmann and Makarkin 2007). Other Mantispidae (both fossil and extant) do not possess the recurred veinlet. All fossil Hemerobiidae, including the oldest known specimens from the Late Jurassic and Early Cretaceous have well-developed recurred veinlet (e.g., Henriksen 1922: fig. 4; Panfilov 1980: fig. 91; Makarkin 1991: fig. 2a; Ponomarenko 1992b: fig. 4). Of the Berothidae, the well-developed recurred veinlet is detected in the Late Jurassic to Early Cretaceous Mesithoninae (Panfilov 1980: figs 86, 87, 90; Ren and Guo 1996: fig. 6; Makarkin 1999: figs 2a, b, 4), and few other genera (e.g., Plesiorobius Klimaszewski & Kevan, 1986: fig. 2). In other berothids, the recurred veinlet is poorly developed, only slightly recurrent and bearing few branches, or does not develop at all (e.g., Engel and Grimaldi 2008: figs 19, 21, 24, 25).

In the Mesozoic family Brongniartiellidae, the well-developed recurred veinlet with long branches is detected in two species from the Early Cretaceous of the Baissa locality, Transbaikalia (Makarkin 2010: fig. 3). The taxonomic affinities of several Mesozoic genera which bear a well developed recurrent veinlet are uncertain, e.g., Sibithone Ponomarenko, 1984: fig. 5a (Early/ Middle Jurassic of Novospasskoe, Siberia); Osmylogramma Ponomarenko, 1992b: fig. 5 (Early Cretaceous of Tsagan-Tsab, Mongolia); Meilingius Ren et al., 2002: fig. 2 and Jurapolystoechotes Ren et al., 2002 (both from Daohugou; pers. obs.).

The presence of the recurrent veinlet was long considered a plesiomorphic condition in Neuroptera (e.g., MacLeod 1970; Oswald 1993; Aspöck and Nemeschkal 1998) because the most ancient, ‘primitive’ extant families possess this condition (e.g., Ithonidae, Psychopsidae), and but it is absent in the youngest, more derived families (e.g., Myrmeleontidae, Ascalaphidae). More recently published phylogenetic analyses of the group place these families in more derived positions and thus the presence of the recurrent veinlet is now considered an apomorphic state evolving independently at least three times (Winterton et al. 2010; Winterton and Makarkin 2010; Yang et al. 2012). The origin of the recurrent veinlet throughout Neuroptera most likely occurred during in the late Permian to earliest Triassic (Figure 7). This condition was secondarily lost at least five times in those families of the clade ‘R’ that lack this condition; and was never possessed in families basal to this clade.

The two species described above from the Middle Jurassic locality of Daohugou are the oldest known record of Nymphidae, indeed, of Myrmeleontoidea. Myrmeleontoidea are believed to belong to the suborder Myrmeleontiformia, which also includes Psychopsidae (Aspöck et al. 2001) or all families of Psychopsoidea (Engel and Grimaldi 2008). According to our phylogeny (Yang et al. 2012: fig. 32), however, the clade comprising Myrmeleontoidea and Chrysopoidea is sister to the ithonoid clade, and that combined clade is sister to the psychopsoeid clade. Known psychopsoeid fossils are much older than those of Myrmeleontoidea, and the recurrent veinlet first appears in their Triassic representatives (see above). Therefore, the presence of the recurrent veinlet in the Middle Jurassic Nymphidae may be interpreted as a remnant of ancient condition occurring in the psychopsoeid + ithonoid + chrysopoid + myrmeleontoid
The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)...

clade before separation of the clade Myrmeleontoidea + Chrysopoidea. This character condition was lost in other families of Myrmeleontoidea and Chrysopoidea, and in the younger Nymphidae.

The recurrent veinlet is known mainly in species whose costal space is basally dilated. The strong dilation of the basal portion of the costal space is characteristic of Neuroptera. It never occurs in other Holometabola including other Neuropterida orders, although in some taxa, the costal space is markedly dilated at some distance from wing base (e.g., Raphidioptera: Aspöck 1986; the mecopteran family Dinopanorpidae: Archibald 2005: figs 3A–D). The costal space of earlier Neuroptera (basal to the clade ‘R’) is also narrowed basally. The appearance and development of the recurrent veinlets and the strong dilation of the basal costal space are probably functionally and structur-
ally tied. However, in some Mesozoic Neuroptera which possess the recurrent veinlet the costal space is narrowed basally: the two species of the Middle Jurassic Nymphidae described here, and at least some Early Jurassic Prohemerobiidae and the Middle Jurassic Ithonidae. It seems reasonable that these taxa probably retained the recurrent veinlet after the costal space was secondarily narrowed in those lineages.

Subcosta anterior (ScA)

The ScA in all Neuroptera (when present) has a similar configuration: a short vein structure terminating on ScP before the recurrent veinlet. The structure of ScA of *Daonymphes bisulca* gen. et sp. n. is very similar to that found in extant Nymphidae, especially *Nymphes* Leach, 1814 (VM, pers. obs.) and closely related genera (e.g., *Australonymphes* sp.: see Riek 1967: fig. 3D). A similar ScA occurs in most extant Hemero- biidae (e.g., Makarkin 1993: fig. 13). Other extant Neuroptera have no distinct ScA, but a convex sclerotized bulge presented in the humeral area of most other families is considered as the modified ScA (Kukalová-Peck and Lawrence 2004). Of the fossil taxa, the ScA is present in the Mesozoic Kalligrammatidae (e.g., Yang et al. 2011: fig. 2) and Parakseneuridae (Yang et al. 2012: figs 24, 25, 30; QY, VM, pers. obs.). According to the hypothesis of Kukalová-Peck (1983), the ScA of the pterygote wing venation ground plan is a relatively short vein running to the costal margin with a strong subcostal brace connecting ScA and ScP midway (Kukalová-Peck 1983, fig. 15), or by other interpretation the ScA is divided midway into the anterior branch (ScA1+2) running to the costal margin and the posterior branch (ScA3+4) terminating on ScP (Kukalová-Peck 1991: fig. 6.3C). Such a primitive ScA is believed to present in the gigantic Carboniferous Bojophlebiidae (Riek and Kukalová-Peck 1984: fig. 20; Kukalová-Peck 1991: fig. 6.14A), although it appears to be poorly preserved (Prokop et al. 2010). In general, the configuration of ScA in some Paleozoic Palaeoptera is similar to that of Neuroptera in that ScA is terminating on ScP, i.e., in Dictyoneuridae (Palaeodictyoptera; see Kukalová 1970: figs 55, 59, 60, 73) and some families of Ephemeroptera (e.g., Carpenter 1979: figs 1, 8, 11); however, the ScA is branched in the latter order.

The ground-plan Neuroptera wing venation is hypothesized to have ScA consisting of two separate veins with no common stem, ScA1+2 and ScA3+4; the former runs to the costal margin, the latter is terminated on ScP (Haas and Kukalová-Peck 2001: fig. 1; Kukalová-Peck 2009: fig. 14). Therefore, the ScA of Neuroptera may be interpreted as the homologue of this hypothesized ScA3+4, i.e., as the ground-plan neuropteran ScA lost its anterior branch ScA1+2. ScA in other Neuroptera orders (Megaloptera, Raphidioptera) is configured similarly to most other winged insects, i.e., a vein terminating on the costal margin (Kukalová-Peck 1983: fig. 17A; 1991: fig. 6.16; Liu et al. 2013). Such a configuration of ScA is most developed in Orthoptera (e.g., Béthoux and Nel 2001: fig. 1). This state of ScA may be interpreted as the homologue of the hypothesized ScA1+2, i.e., as the ground-plan Neuroptera ScA lost its posterior branch ScA3+4.
The configuration of ScA characteristic of Neuroptera (i.e., short and terminating on ScP) does not occur in other orders of Neoptera, and therefore may be considered as an autapomorphy. On the other hand, the similarities of ScA in Neuroptera and some Paleozoic Palaeoptera (see above) may indicate their homology, and therefore a symplesiomorphy. The latter, however, appears to be less probable than the former.

Acknowledgements

We thank S. Bruce Archibald (Simon Fraser University, Burnaby, Canada) for correction of English; Shaun Winterton (California State Collection of Arthropods, Sacramento, California, U.S.A.) and André Nel (Museum National d’Histoire Naturelle, Paris, France) for critical reading of the manuscript. This work was supported by the National Basic Research Program of China (973 Program) (grant 2012CB821906), National Science Foundation of China (grants 31230065, 31272352, 41272006), Project of Great Wall Scholar and KEY project of Beijing Municipal Commission of Education Project (grants KZ201310028033), China Geological Survey (grant 1212011120115).

References

Achtelig M (1975) Die Abdomenbasis der Neuropteroidea (Insecta, Holometabola). Eine vergleichend anatomische Untersuchung des Skeletts und der Muskulatur. Zoomorphologie 82: 201–242. doi: 10.1007/BF00993588
Ansorge J (1996) Insekten aus dem oberen Lias von Grimmen (Vorpommern, Norddeutschland). Neue Paläontologische Abhandlungen 2: 1–132.
Archibald SB (2005) New Dinopanorpidae (Insecta: Mecoptera) from the Eocene Okanagan Highlands (British Columbia, Canada and Washington State, USA). In: Archibald SB, Greenwood DR (Eds) The Okanagan Highlands: Eocene biota, environments, and geological setting. Canadian Journal of Earth Sciences 42: 119–136.
Archibald SB, Makarkin VN (2006) Tertiary giant lacewings (Neuroptera: Polystoechotidae): revision and description of new taxa from western North America and Denmark. Journal of Systematic Palaeontology 4: 119–155 [Errata: 4: 307]. doi: 10.1017/S1477201906001817
Archibald SB, Makarkin VN, Ansorge J (2009) New fossil species of Nymphidae (Neuroptera) from the Eocene of North America and Europe. Zootaxa 2157: 59–68.
Aspöck H (1986) The Raphidioptera of the world: a review of present knowledge. In: Gepp J, Aspöck H, Hölzle H (Eds) Recent Research in Neuropterology. Proceedings of the 2nd International Symposium on Neuropterology. Privately printed, Graz, Austria, 15–29.
Aspöck U, Nemeschkal HL (1998) A cladistic analysis of the Berothidae (Neuroptera). Acta Zoologica Fennica 209: 45–63.
Aspöck U, Plant JD, Nemeschkal HL (2001) Cladistic analysis of Neuroptera and their systematic position within Neuroptera (Insecta: Holometabola: Neuroptera: Neuroptera). Systematic Entomology 26: 73–86. doi: 10.1046/j.1365-3113.2001.00136.x
Bechly G (2000) Mainstream cladistics versus Hennigian phylogenetic systematics. Stuttgarter Beiträge zur Naturkunde (A) 613: 1–11.

Béthoux O, Nel A (2001) Venation pattern of Orthoptera. Journal of Orthoptera Research 10: 195–198. doi: 10.1665/1082-6467(2001)010[0195:VPOO]2.0.CO;2

Bode A (1953) Die Insektenfauna des Ostniedersächsischen Oberen Lias. Palaeontographica: Beiträge zur Naturgeschichte der Vorwelt (A) 103: 1–375

Carpenter FM (1929) A Jurassic neuropteran from the lithographic limestone of Bavaria. Psyche 36: 190–194. doi: 10.1155/1929/12134

Carpenter FM (1979) Lower Permian insects from Oklahoma. Part 2. Orders Ephemeroptera and Palaeodictyoptera. Psyche 86: 261–290. doi: 10.1155/1979/96359

Engel MS, Grimaldi DA (2008) Diverse Neuropterida in Cretaceous amber, with particular reference to the paleofauna of Myanmar (Insecta). Nova Supplementa Entomologica 20: 1–86.

Gao KQ, Ren D (2006) Radiometric dating of ignimbrite from Inner Mongolia provides no indication of a post-Middle Jurassic age for the Daohugou Beds. Acta Geologica Sinica 80: 42–45. doi: 10.1111/j.1755-6724.2006.tb00793.x

Grimaldi DA, Engel MS (2005) Evolution of the insects. Cambridge University Press, Cambridge, UK, xv+755 pp.

Haas F, Kukalová-Peck J (2001) Dermaptera hind wing structure and folding: new evidence for familial, ordinal and superordinal relationships within Neoptera (Insecta). European Journal of Entomology 98: 445–509.

Haase E (1890) Bemerkungen zur Palaeontologie der Insecten. Neues Jahrbuch für Mineralogie, Geologie und Paläontologie 2(1): 1–33.

Handlirsch A (1906–1908) Die fossilen Insekten und die Phylogenie der rezenten Formen. Ein Handbuch für Palaeontologen und Zoologen. W. Engelmann, Leipzig, ix + 1430 pp. (issued in 1906, pp. 1–640, 1907, pp. 641–1120, 1908, pp. 1121–1430).

Handlirsch A (1939) Neue Untersuchungen über die fossilen Insekten mit Ergänzungen und Nachträgen sowie Ausblicken auf phylogenetische, palaeogeographische und allgemein biologische Probleme. II. Teil. Annalen des Naturhistorischen Museums in Wien 49: 1–240.

Hennig W (1981) Insect phylogeny. John Wiley & Sons, Chichester, New York, Brisbane & Toronto, 514 pp.

Henriksen KL (1922) Eocene insects from Denmark. Danmarks Geologiske Undersøgelse (II) 37: 1–36.

Jell PA (2004) The fossil insects of Australia. Memoirs of the Queensland Museum 50 (1): 1–124.

Jepson JE, Heads SW, Makarkin VN, Ren D (2013) New fossil mantidflies (Insecta: Neuroptera: Mantispidae) from the Mesozoic of north-eastern China. Palaeontology 56: 603–613. doi: 10.1111/pala.12005

Jepson JE, Makarkin VN, Coram RA (2012) Lacewings (Insecta: Neuroptera) from the Lower Cretaceous Purbeck Limestone Group of southern England. Cretaceous Research 34: 31–47. doi: 10.1016/j.cretres.2011.10.001

Khramov AV (2011) Two new lacewings (Neuroptera) from the Upper Jurassic locality Shar-Teg (Mongolia). Paleontologicheskii Zhurnal 2011 (2): 52–56. [in Rus-
The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)...
Makarkin VN (1991) Miocene Neuroptera from North Caucasus and Sikhote-Alin’. Paleontologicheskii Zhurnal 1991 (1): 57–68. [in Russian; English translation: (1992) Paleontological Journal 25 (1): 55–65]
Makarkin VN (1993) The brown lacewings from Vietnam (Neuroptera, Hemerobiidae). Tropical Zoology 6: 217–226. doi: 10.1080/03946975.1993.10539222
Makarkin VN (1996) [1997] Fossil Neuroptera of the Lower Cretaceous of Baisa, East Siberia. Part 5. Mantispidae. Russian Entomological Journal 5: 91–93.
Makarkin VN (1999) Fossil Neuroptera of the Lower Cretaceous of Baisa, East Siberia. Part 6. Mesithonidae (Insecta). Neues Jahrbuch für Geologie und Paläontologie, Monatshefte 1999 (12): 705–712.
Makarkin VN (2010) New psychopsoid Neuroptera from the Lower Cretaceous of Baissa, Transbaikalia. Annales de la Société Entomologique de France (N.S.) 46: 254–261.
Makarkin VN, Archibald SB (2009) A new genus and first Cenozoic fossil record of moth lacewings (Neuroptera: Ithonidae) from the Early Eocene of North America. Zootaxa 2063: 55–63.
Makarkin VN, Yang Q, Ren D (2012) A comparative overview of the neuropteran assemblage of the Early Cretaceous Yixian Formation (China), with description of a new genus of Psychopsideae (Insecta: Neuroptera). Cretaceous Research 35: 57–68. doi: 10.1016/j.cretres.2011.11.013
Makarkin VN, Yang Q, Ren D (2013) A new Cretaceous family of enigmatic two-winged lacewings (Neuroptera). Fossil Record 16: 67–75. doi: 10.1002/mmng.201300002
Martins-Neto RG (2005) New Neuroptera from Crato Formation, Lower Cretaceous, Araripe Basin, Northeast Brazil. GAEA 1: 5–10.
Menon F, Martins-Neto RG, Martill DM (2005) A new Lower Cretaceous nymphid (Insecta, Neuroptera, Nymphidae) from the Crato Formation of Brazil. GAEA 1: 11–15.
Novokshonov VG (1996) Systematic position of some Upper Permian Myrmeleontida (=Neuroptera; Insecta). Paleontologicheskii Zhurnal 1996 (1): 39–47. [in Russian; English translation: Paleontological Journal 30: 38–45]
Oswald JD (1993) Revision and cladistic analysis of the world genera of the family Hemerobiidae (Insecta: Neuroptera). Journal of the New York Entomological Society 101: 143–299.
Panfilov DV (1980) New representatives of lacewings (Neuroptera) from the Jurassic of Karatau. In: Dolin VG, Panfilov DV, Ponomarenko AG, Pritykina LN. Fossil insects of the Mesozoic. Naukova Dumka, Kiev, 82–111. [In Russian]
Peng YY, Makarkin VN, Yang Q, Ren D (2010) A new silky lacewing (Neuroptera: Psychopsideae) from the Middle Jurassic of Inner Mongolia, China. Zootaxa 2663: 59–67.
Ponomarenko AG (1984) Neuroptera from the Jurassic of eastern Asia. Paleontologicheskii Zhurnal 1984 (3): 64–73 [in Russian; English translation: Paleontological Journal, 1985, 18 (3): 59–69]
Ponomarenko AG (1992a) Neuroptera (Insecta) from the Lower Cretaceous of Transbaikalia. Paleontologicheskii Zhurnal 1992 (3): 43–50. [in Russian; English translation: Paleontological Journal 26 (3): 56–66]
Ponomarenko AG (1992b) New lacewings (Insecta, Neuroptera) from the Mesozoic of Mongolia. In: Grunt TA (Ed) New taxa of fossil invertebrates of Mongolia. Transactions of the Joint Soviet-Mongolian Paleontological Expedition. Vol. 41, 101–111. [in Russian]
The presence of the recurrent veinlet in the Middle Jurassic Nymphidae (Neuroptera)...

Ponomarenko AG (1995) [1996] Upper Liassic neuropterans (Insecta) from Lower Saxony, Germany. Russian Entomological Journal 4: 73–89.
Ponomarenko AG (2002) Superorder Myrmeleontidea Latreille, 1802 (=Neuropteroidea Han- dIlirsch, 1903). In: Rasnitsyn AP, Quicke DLJ (Eds) History of Insects. Kluwer Academic Publishers, Dordrecht, 176–192.
Prokop J, Nel A, Tenny A (2010) On the phylogenetic position of the palaeopteran Syntonopteroidea (Insecta: Ephemeroptera), with a new species from the Upper Carboniferous of England. Organisms Diversity & Evolution 10: 331–340. doi: 10.1007/s13127-010-0022-2
Rambur JP (1842) Histoire Naturelle des Insectes. Névroptères. Fain et thunot, Paris, [xvii] + 534 pp.
Ren D, Engel MS (2007) A split-footed lacewing and two epiosmylines from the Jurassic of China (Neuroptera). Annales Zoologici (Warszawa) 57: 211–219.
Ren D, Engel MS, Lu W (2002) New giant lacewings from the Middle Jurassic of Inner Mongolia, China (Neuroptera: Polystoechotidae). Journal of the Kansas Entomological Society 75: 188–193.
Ren D, Guo ZG (1996) On the new fossil genera and species of Neuroptera (Insecta) from the Late Jurassic of northeast China. Acta Zootaxonomica Sinica 21: 461–479.
Riek EF (1955) Fossil insects from the Triassic beds at Mt. Crosby, Queensland. Australian Journal of Zoology 3: 654–691. doi: 10.1071/ZO9550654
Riek EF (1956) A re-examination of the mecopteroid and orthopteroid fossils (Insecta) from the Triassic beds at Denmark Hill, Queensland, with descriptions of further specimens. Australian Journal of Zoology 4: 98–110. doi: 10.1071/ZO9560098
Riek EF (1967) Structures of unknown, possibly stridulatory, function of the wings and body of Neuroptera; with an appendix on other endopterygote orders. Australian Journal of Zoology 15: 337–348. doi: 10.1071/ZO9670337
Riek EF, Kukalová-Peck J (1984) A new interpretation of dragonfly wing venation based upon Early Upper Carboniferous fossils from Argentina (Insecta: Odonatoidea) and basic character states in pterygote wings. Canadian Journal of Zoology 62: 1150–1166. doi: 10.1139/z84-166
Shcherbakov DE (2008) Madygen, Triassic Lagerstätte number one, before and after Sharov. Alavesia 2: 113–124.
Shi CF, Makarkin VN, Yang Q, Archibald SB, Ren D (in press) New species of Nymphites Haase (Neuroptera: Nymphidae) from the Middle Jurassic of China, with a redescription of the type species of the genus. Zootaxa.
Tillyard RJ (1919) Mesozoic insects of Queensland. No. 5. Mecoptera, the new order Paratrichoptera, and additions to Planipennia. Proceedings of the Linnean Society of New South Wales 44: 194–212.
Tillyard RJ (1922) Mesozoic insects of Queensland. No. 9. Orthoptera, and additions to the Protorthoptera, Odonata, Hemiptera and Planipennia. Proceedings of the Linnean Society of New South Wales 47: 447–470.
Tillyard RJ (1923) Mesozoic insects of Queensland. No. 10. Summary of the Upper Triassic insect fauna of Ipswich, Queensland (with an appendix describing new Hemiptera and Planipennia). Proceedings of the Linnean Society of New South Wales, 48: 481–498.
Tillyard RJ (1933) The panorpoid complex in the British Rhaetic and Lias. In: Fossil Insects 3: 1–79; British Museum (Natural History), London.

Vilesov AP (1995) Permian lacewings (Insecta: Myrmeleontida) from the Chekarda locality (Ural). Paleontologicheskii Zhurnal 1995(2): 95–105. [in Russian; English Translation: Paleontological Journal 29 (2): 115–129.]

Vilesov AP, Novokshonov VG (1994) New fossil insects (Myrmeleontida, Jurinida) from the Upper Permian of eastern Kazakhstan. Paleontologicheskii Zhurnal 1994(2): 66–74. [in Russian; English translation: Paleontological Journal 28(2): 81–92.]

Wedmann S, Makarkin VN (2007) A new genus of Mantispidae (Insecta: Neuroptera) from the Eocene of Germany, with a review of the fossil record and palaeobiogeography of the family. Zoological Journal of the Linnean Society 149: 701–716. doi: 10.1111/j.1096-3642.2007.00273.x

Westwood JO (1854) Contributions to fossil entomology. Quarterly Journal of the Geological Society of London 10: 378–396. doi: 10.1144/GSL.JGS.1854.010.01-02.43

Whalley PES (1988) Mesozoic Neuroptera and Raphidioptera (Insecta) in Britain. Bulletin of the British Museum of Natural History (Geology) 44: 45–63.

Winterton SL, Hardy NB, Wiegmann BM (2010) On wings of lace: phylogeny and Bayesian divergence time estimates of Neuropterida (Insecta) based on morphological and molecular data. Systematic Entomology 25: 349–378. doi: 10.1111/j.1365-3113.2010.00521.x

Winterton SL, Makarkin VN (2010) Phylogeny of moth lacewings and giant lacewings (Neuroptera: Ithonidae, Polystoechotidae) by using DNA sequence data, morphology, and fossils. Annals of the Entomological Society of America 103: 511–522. doi: 10.1603/AN10026

Yang Q, Makarkin VN, Ren D (2011) Two interesting new genus of Kalligrammatidae (Neuroptera) from the Middle Jurassic of Daohugou, China. Zootaxa 2873: 60–68.

Yang Q, Makarkin VN, Ren D (2013) A new genus of the family Panfiloviidae (Insecta, Neuroptera) from the Middle Jurassic of China. Palaeontology 56: 49–59. doi: 10.1111/j.1475-4983.2012.01157.x

Yang Q, Makarkin VN, Winterton SL, Khramov AV, Ren D (2012) A remarkable new family of Jurassic insects (Neuroptera) with primitive wing venation and its phylogenetic position in Neuropteroidea. PLoS ONE 7(9): e44762. doi: 10.1371/journal.pone.0044762

Yang Q, Ren D, Shih CK, Wang YJ, Shi CF, Peng YY, Zhao YY (2010) Neuroptera – Grace with Lace. In: Ren D, Shih CK, Gao TP, Yao YZ, Zhao YY. Silent Stories – Insect Fossil Treasures from Dinosaur Era of the Northeastern China. Sciece Press, Beijing, 165–179.

Yang Q, Zhao YY, Ren D (2009) An exceptionally well-preserved fossil Kalligrammatid from the Jehol Biota. Chinese Science Bulletin 54: 1732–1737. doi: 10.1007/s11434-009-0284-2