Sambia succinica, a crown group tenthredinid from Eocene Baltic amber (Hymenoptera: Tenthredinidae)

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Abstract
Sambia succinica gen. et sp. n. from Eocene Baltic amber is described and illustrated. It is apparently the first amber fossil that can be definitively assigned to Tenthredininae. It displays two diagnostic forewing characters for this subfamily: having a bend distally in vein R and the junctions of veins M and Rs + M with vein R being some distance from each other. The variance and possible transitions between the anal vein configurations among the genera in Tenthredininae is briefly discussed.

Keywords
amber inclusion, sawfly, Tertiary, Eocene, taxonomy

Introduction
Tenthredinidae is the largest family of non-apocritan Hymenoptera by far, comprising more than 5500 described species (Huber 2009; Taeger & Blank 2010). Together with five other families they comprise the Tenthredinoidea or true sawflies. The larvae of the members of the superfamily are all herbivores and most are external feeders on green parts of angiosperms; however, other host plants and feeding modes (e.g., leafrolling, leafmining, or galling in leaves, buds and shoots; see Nyman et al. 1998, 2000) do occur. Recent comprehensive treatments of the phylogeny of the basal hymenopteran lineages, while providing strong support for the Tenthredinoidea, have consistently failed to retrieve the Tenthredinidae as monophyletic (Vilhelmsen 2001; Schulmeister 2003; Ronquist et al. in press). Indeed, a thorough, phylogenetically based revision of this family and its constituent subclades is highly needed. Of the six subfamilies recognized within the Tenthredinidae, Tenthredininae is the largest, with approx. 1700
described species comprising about 30% of the known diversity of the family (Taeger & Blank 2010).

The fossil record of Tenthredinoidea is substantial; putative stemgroup taxa are known from the Late Jurassic and Early Cretaceous, whereas fossils that can be placed in extant families do not occur until the Tertiary (see Grimaldi & Engel 2005 and references therein). The majority of tenthredinoid fossils known are compressions, only Argidae and Diprionidae being represented in Tertiary amber so far (Grimaldi & Engel 2005). Brischke (1886) mentioned _Tenthredo_ in a list of genera occurring in Baltic amber (a record repeated by Handlirsch (1907), Statz (1936), Larsson (1978), and Spahr (1987)), but the whereabouts of the material is unknown as well as the veracity of his identification. In this paper, we describe an adult female from Eocene Baltic amber (Lutetian stage, ca. 45 million years ago), which we tentatively place in the subfamily Tenthredininae. We place the fossil taxon in a new genus, since comparison with a range of representatives of extant members of the Tenthredininae and the descriptions of most of the extinct ones did not reveal any obvious candidate for placement in an existing genus.

**Materials and Methods**

The specimen is included in a piece of middle Eocene (Lutetian) Baltic amber. The age, paleofauna, and origin of Baltic amber have been reviewed by Weitschat and Wichard (2002, 2010) and Grimaldi and Engel (2005).

**Extant material for comparison**

The following extant taxa were all dissected and preserved in glycerol as part of an ongoing project aiming to resolve the phylogeny of the Tenthredinidae, its constituent subgroups and its closest relatives: _Aglaostigma fulvipes_ (Scopoli, 1763); _Corymbas koreana_ Konow, 1903; _Filacus albipes_ (Provancher, 1895); _Lagium atrovioleaceum_ (Norton, 1860); _Leucopelmonus annulicornis_ (Harrington, 1893); _Macrophya duodecimpunctata_ (Linnaeus, 1758); _Pachyprotasis rapae_ (Linnaeus, 1767); _Perineura rubi_ (Panzer, 1805); _Rhogogaster viridis_ (Linnaeus, 1758); _Sciapteryx consobrina_ (Klug, 1816); _Siobla sturmii_ (Klug, 1817); _Tenthredo (T.) scrophulariae_ Linnaeus, 1758; _Tenthredo (Elinora) koehleri_ Klug, 1817; _Tenthredopsis ornata_ (Audinet-Serville, 1823); _Zaschizonyx montana_ (Cresson, 1865). In addition, comparison was made with the following taxa, which were observed from entire pinned specimens deposited in the Natural History Museum, London: _Armitarsus punctifemoratus_ Malaise, 1931; _Conaspidia guttata_ (Matsumura, 1912); _Conaspidia byalina_ Sato, 1933; _Colochelyna magrettii_ Konow, 1898; _Tenthredo (Elinora) amasiensis_ (Kriechbaumer, 1869); _Tenthredo (Tenthredo) arctica_ (Thomson, 1870); _Tenthredo (Propodea) fentoni_ W.F. Kirby, 1882; _Tenthredo (Tenthredina) fortunii_ W.F. Kirby, 1882; _Tenthredo (Peus) priva_ (Konow, 1903); _Tenthredo sibirica_ (Kriechbaumer, 1869); _Tenthredo (Propodea) spinosa_ Cameron, 1899; _Tenthredo (Metallopeus) splendida_ (Konow, 1907); _Tyloceridius dorsatus_ (Mocsáry, 1883).
Fossil taxa considered

Taeger et al. (2010) listed 19 fossil species under Tenthredininae, all of which are compression fossils with little to no relief. These species are summarized in Table 1, noting those of which the descriptions were consulted and compared with the current amber inclusion. Should Piton’s (1940) species be assigned correctly, it would then represent the oldest record of the subfamily, predating the current fossil by 6–10 million years.

Examination and imaging

The amber fossil and the dissected extant taxa were examined under a Leica MZ APO dissecting stereomicroscope. The photographic illustrations were compiled from stacks of images produced with a Leica DFC 420 digital camera and integrated with Leica software. The wings of extant taxa imaged in Figs 5–8 were slide mounted in glycerol prior to imaging.

Systematic Paleontology

Family Tenthredinidae Leach
Subfamily Tenthredininae Leach

Sambia gen.n.

Type species

Sambia succinica sp.n.

Diagnosis

The new genus can be recognized by the following unique combination of characters: clypeus only slightly incurved; inner margins of compound eyes parallel; antenna subclavate, with nine antennomeres; occipital carina present dorsally, absent laterally; inner margins of propleura abutting; metatibial spur one-third length of metabasitarsus; metabasitarsus longer than tarsomeres 2–4; forewing with bend in R distal; junction of M and Rs+M with R some distance from each other; cell 1RS wider than 2RS; insertion of cu-a on Cu1 one-third from base; 1a distal, oblique.

Etymology

The new genus-group name honors the Old Prussian tribe of the Sambians who occupied Sambia (Samland/the Sambian Peninsula) where much Baltic amber historically has originated. The tribe was active in the amber trade prior to being conquered by the Teutonic Knights in the 13th Century (Wikipedia). The name is feminine.
**Sambia succinica** sp.n. (Figs 1–4)

*Specimen provenance*

_Holotype._ The holotype specimen has the label: ‘AMBER: BALTIC / Middle Eocene (Lutetian) / Blau Erde, Northern Europe / KU-NHM-ENT, B-145’. It is deposited in the Division of Entomology, Natural History Museum, University of Kansas, Lawrence, KS, USA.

*Diagnosis*

As for the genus (see above).

*Description*

_Female._ Specimen mostly intact, except dorsally where it is exposed and eroded, removing most of mesonotum (see Fig. 4, right). Amber matrix mostly clear, with few cracks; small occlusions present at mouthparts and ovipositor. Body length as preserved 5.6 mm, forewing length 4.8 mm, head width 1.7 mm. _Head._ Broad; compound eyes slightly converging, cuticular covering missing from central part of compound eyes; ocelli placed in obtuse triangle (Fig. 2); clypeus slightly incurved; occipital carina developed dorsally (Fig. 2), but apparently not laterally; frons, gena, vertex, and occiput covered with short, erect setae. Antenna with nine antennomeres (Fig. 1; left missing apical antennomere); scapus and pedicellus short, latter slightly longer than broad; antennomere 3 longest, approximately 1.5-times longer than antennomere 4; antennomeres 5–9 broader than antennomeres 1–4, tapering slightly towards antennomere 9; antennomeres 6–9 each shorter than antennomere 5, subequal in length. Features of mouthparts hard to observe; maxillary palp with six palpomeres, labial palp with four. _Thorax._ Pronotum flat, closely abutting mesopleuron, with incurvation for anterior thoracic spiracle (Fig. 1); propleura mostly hidden, medioventral margins abutting for some distance; two apical protibial spurs, anterior spur (calcar) longer, slightly curved and bifid at tip. Mesonotum destroyed (see above); postspiracular sclerite present, prepectus not observed; posterior thoracic spiracle not observed, slight swelling/lobe present at usual position of spiracle (Fig. 1); other features, including sculpture, impossible to observe on meso- and metapleuron; two apical mesotibial spurs of subequal length. Metanotum covered by wings; metatibia significantly longer than metafemur (Fig. 1), with two apical spurs of equal length and not much longer than width of metatibia (Fig. 3); metatarsomeres 1–4 with distinct pulvilli, metabasitarsomere approximately equal in length to metatarsomeres 2–4 (Fig. 3); pretarsal claw bifid, rami equal in length. _Wings._ Wings in normal resting position above abdomen, forewings covering hind wings, making it impossible to observe latter and dorsal part of abdomen. Costal cell narrow, veins C and R separated by less than their combined width in middle of costal cell; vein R bent posteriorly at junction of R and M, this junction separate from junction between R and Rs+M (Fig. 4), latter being slightly curved; abscissa of vein Rs well developed; vein 2r insert on Rs close to junction between Rs and 3r-m; angle between veins Cu1 and 1m-cu more than 120°, latter
parallel with vein M (Fig. 4); veins 1m-cu and 2m-cu insert on different cells; both veins 2r-m and 3r-m present; proximal part of posterior anal vein (2A) difficult to observe, apparent crossvein 1a oblique and distal in anal cell (Fig. 4), recurving vein 3A might be present, but distal termination impossible to see. Abdomen. Abdominal tergum 1 could not be observed (see above); lateral parts of abdominal tergites accommodating spiracles continuous with dorsal parts. Ovipositor apparatus partly occluded,
but outline of proximal parts (first and second valvifers, first and second valvulae) visible; cerci not observed.

**Etymology**

The specific epithet is based on “succinite”, the term for Baltic amber, referring to its high succinic acid content and the Latin term *succinum*, meaning “amber”.

**Discussion**

The placement of *S. succinica* in Tenthredininae is based on the combination of two forewing characters: the bend in vein R distally and the junctions of veins M and Rs+M with vein R being some distance from each other (cf., Figs 5–8). Each character in isolation is shared by other members of the Tenthredinoidea, e.g., the bend in vein R by some Argidae and Diprionidae, the M – Rs+M configuration by Nematinae, most Selandriinae, and some Heterarthrinae (Vilhelmsen, unpubl.). However, the two traits are only very rarely observed together outside Tenthredininae. Furthermore, the presence of a dorsal occipital carina is rather uncommon outside this subfamily.

A number of characters were considered when trying to decide whether to place the fossil taxon in an extant genus or describe it as a new genus (condition in fossil in italics): The inner margins of the compound eyes (*parallel/converging*); the configuration of the clypeus (*straight/incurved*); the presence of a dorsal (*present*) and lateral (*absent*) occipital carina; the configuration of the antenna (not clavate/subclavate); the inner margins of the propleura (*separate/abutting*); the relative length of the metatibial spur (*1/3 the length of the metabasitarsus/longer*); the relative length of the metabasitarsus (*shorter/longer than tarsomere 2-4*); the relative widths of cells 1RS and 2RS in the forewing (*1RS wider than 2RS/2RS wider than 1RS*); the insertion of crossvein cu-a on vein Cu1 in the forewing (*1/3 from base/further away*); the configuration of crossvein 1a in the forewing (absent/proximal, straight/distal, oblique (but see above)).

None of the extant genera examined provide a good match for *S. succinica*. *Siobla* resembles *S. succinica* in a number of features, but differs in having the lateral occipital carina present, the apical metatibial spur long, forewing cell 2RS wider than 1RS (compare Figs 4 and 5), and forewing crossvein cu-a inserting on Cu1 very close to M. The only taxon having the occipital carina present dorsally but absent laterally like *S. succinica* is *A. fulvipes*, but it differs from the former in having non-clavate antennae, the medioventral propleural margins being separate, and having crossvein 1a in the forewing proximal and straight. If a recurving vein 3A is present in the forewing of the fossil, this indicates that veins 2A and 3A might be discontinuous for some distance (see Fig. 8), a condition that is most developed in some specimens of *Pachyprotasis*. This genus differs from *S. succinica* in having a deeply incurved clypeus, an occipital carina present laterally as well as dorsally, the antenna elongate and not clavate, and the metatibial apical spur reaching half the length of the metabasitarsus.
Figs 5–8. Forewings of extant Tenthredininae: (5) *Siobla sturmii* (Klug); (6) *Corymbas koreana* Konow; (7) *Tenthredo scrophulariae* Linnaeus; (8) *Macrophya duodecimpunctata* (Linnaeus). Scale bars=1 mm. This figure is published in colour in the online edition of this journal, which can be accessed via http://booksandjournals.brillonline.com/content/1876312x.
There is some variation in the configuration of the forewing anal veins among Tenthredininae. *Corymbas* (Fig. 6) has both veins 1A and 2A developed and has an incurvation in the latter as does *Siobla*, but crossvein 1a is absent, a configuration otherwise only observed in some Selandriinae. In *Aглаostigma*, *Filacus*, *Lagium*, *Lеуocopelmonus*, *Rhogogaster*, *Sciapteryx*, *Tenthredo* and *Tenthredopsis*, 1a is placed more proximally and in the incurvation of 2A and 1a is consequently shortened and straight (Fig. 7). In *Macrophya*, *Pachyprotasis*, *Perineura* and *Zaschizontyx*, vein 2A is discontinuous with 3A and two separate anal cells are formed (Fig. 8). It is possible that the proximal part of the vein delimiting the distal anal cell posteriorly is homologous with 1a. Alternatively, 1a has been obliterated when the midpoint of vein 2A has been displaced anteriorly to merge with the midpoint of 1A. In this case, the transformation from the condition observed in *Siobla* to that in *Macrophya* and others perhaps took place via an intermediate state like the one displayed by *Aглаostigma* and others.

Very little information that can be put in a phylogenetic context can be gleaned from the descriptions of the compression fossils putatively assigned to Tenthredininae (see Table 1). Most of the fossils (e.g., those from the Eocene-Oligocene boundary of Florrisant, CO, USA) were described approximately a century ago, preserve very few reliable characters, and no attempt has been made to put them in any phylogenetic context, assuming they are correctly assigned as to family. Zhelochovtzev and Rasnitsyn (1972) attempted to provide revised interpretations of a small set of Florissant tenthredinids, but again, few details outside of forewing venation are available and all except *Florissantinus angulatus* appear to belong to other subfamilies. Also, as compression fossils they are less likely to yield character information than amber inclusions. In addition, many of the older descriptions are rather cursory and often poorly or not at all illustrated (e.g., those of Brues, Cockerell, and Rohwer).

Of the characters discussed above, the configuration of the anal veins is the only one that most descriptions provide information on. The most common configuration, displayed by *Macrophya adventitia*, *M. pervestuta*, *Nortonella typica*, *Tenthredo infossa*, *T. meunieri*, *T. miocenica*, *T. oblita*, *T. physia*, *T. saxorum*, *T. submersa* and *Tenthredoides plurilocellata*, is the one also present in most extant Tenthredininae, i.e., veins 1A and 2A separate and crossvein 1a short, straight, and placed proximally. Only *Tenthredo petrae* unequivocally has the plesiomorphic condition, oblique and distally placed crossvein 1a. *Tenthredoides plurilocellata* actually has two straight transverse veins in the anal cell of the right forewing: one proximal straight vein like all the taxa just listed, and one distal straight (‘a2’ in fig. 225A of Zhang 1989). The latter is probably best interpreted as a teratological feature as only one, proximal straight vein is observed in the anal cell of the left forewing of this fossil (compare figs 225A and B of Zhang 1989), and two transverse veins in the anal cell are not observed in any other Tenthredinoidea known to us.

It is, thus, not really possible to attempt to place any of the described compression fossils more precisely within the Tenthredininae, or to decide whether they should be placed in the subfamily at all. To achieve that, the fossils themselves will have to be re-examined. However, that is beyond the scope of this contribution. Regardless,
Table 1. Nineteen fossil species putatively assigned to the Tenthredininae in Taeger et al. (2010) and those for which original descriptions were consulted for comparison with the current amber specimen.

<table>
<thead>
<tr>
<th>Fossils considered herein</th>
<th>Age</th>
<th>Locality</th>
</tr>
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<tbody>
<tr>
<td>Macrophya adventitia Lewis, 1969</td>
<td>Miocene</td>
<td>Latah Formation, Washington &amp; Idaho, USA</td>
</tr>
<tr>
<td>Macrophya pervetusta Brues, 1908</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
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<tr>
<td>Nortonella typica Rohwer, 1908</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
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<tr>
<td>Taeniurites foris Cockerell, 1917a</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
</tr>
<tr>
<td>Tenthredininae bifasciata Meunier, 1915</td>
<td>Oligocene</td>
<td>Aix-en-Provence, France</td>
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<tr>
<td>Tenthredo avia Brues, 1908</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
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<tr>
<td>Tenthredo fenestralis Cockerell, 1927</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
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<tr>
<td>Tenthredo infissa Brues, 1908</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
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<tr>
<td>Tenthredo meunieri Taeger et al., 2009</td>
<td>Oligocene</td>
<td>Rott, Germany</td>
</tr>
<tr>
<td>Tenthredo miocenica Zhang &amp; Zhang, 1990</td>
<td>Miocene</td>
<td>Shanwang, Shandong, China</td>
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<tr>
<td>Tenthredo oblieta Cockerell, 1917b</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
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<tr>
<td>Tenthredo petrae Zhang &amp; Zhang, 1990</td>
<td>Miocene</td>
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<td>Tenthredo rhyia Zhang, 1989</td>
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<td>Tenthredo saxorum Rohwer, 1908</td>
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<td>Tenthredo submersa Cockerell, 1907</td>
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<tr>
<td>Tenthredo toddi Cockerell, 1914</td>
<td>Eocene-Oligocene</td>
<td>Florissant, Colorado, USA</td>
</tr>
<tr>
<td>Tenthredoides pluriocellata Zhang, 1989</td>
<td>Miocene</td>
<td>Shanwang, Shandong, China</td>
</tr>
</tbody>
</table>

Fossils not considered herein

| Tenthredo gervaisi Heer, 1861                      | Oligocene         | Aix-en-Provence, France                |
| Tenthredo primordialis Piton, 1940                 | Paleocene         | Menat, France                          |

Note: Tenthredo meunieri Taeger et al., 2009 is a replacement name for *T. fasciata* Meunier, 1922. *Tenthredo miocenica* Zhang & Zhang, 1990 was redescribed and refigured by Zhang et al. (1994).

*S. succinica* in mid-Eocene amber predates any of the aforementioned fossils and therefore represents the oldest, definitive record of the Tenthredininae.

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