A unique, large-sized stem Odonata (Insecta) found in the early Pennsylvanian of New Brunswick (Canada)

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Abstract. A stem relative of dragon- and damselflies, Brunellopteron norradi Béthoux, Deregnaucourt and Norrad gen. et sp. nov., is documented based on a specimen found at Robertson Point (Grand Lake, New Brunswick, Canada; Sunbury Creek Formation; early Moscovian, Pennsylvanian) and preserving the basal half of a hindwing. A comparative analysis of the evolution of wing venation in early odonates demonstrates that it belongs to a still poorly documented subset of species. Specifically, it displays a MP + CuA fusion, a CuA + CuP fusion, and a CuP + AA fusion, but it lacks the “extended” MP + Cu / CuA fusion and the “extended” (CuP / CuA + CuP) + AA fusion, the occurrence of which is typical of most Odonata, including Meganeura-like species. The occurrence of intercalary veins suggests that its closest relative might be Galloptypus oudardi Nel, Garroute and Roques, 2008, from the Moscovian of northern France.

1 Introduction

Conjecture of primary homology in wing venation of extant dragon- and damselflies (Odonata) remained a matter of debate for most of the 20th century. Despite the ancient Meganeura-like insects being known, attempts to relate the wing venation of stem and crown Odonata to other insect groups abutted a number of issues. Among them was the propensity, in odonate wings, of cross-veins to convert into main-vein-like elements, including so-called “intercalary veins”, which can prove difficult to distinguish from surrounding genuine main veins. Confusion also arose from persisting gaps in the documentation of venation patterns which had experienced intense modifications since the Pennsylvanian.

A landmark in this debate was the proposal made by Tillyard (1925b), who, at that time, had adopted the view that the insect wing venation ground plan is composed of a number of primary veins sharing the same branching pattern, specifically a convex, anterior branch and a concave, posterior one, also known as the serial ground plan (Lameere, 1922, 1923; contrast with Tillyard, 1925a, in which the author expressed doubts on the adequacy of the Comstock–Needham terminol-
ogy, he used previously). It is relevant to first emphasize that even though Tillyard (1925b, p. 42) stressed that Protodonata (including, in his opinion, Meganeuridae and a few other species) and Odonata were two clearly distinct taxa, he also stated that MP and CuA (his Cu\textsubscript{1}) were not only present as free veins beyond the wing base in Meganeuridae but also in species he regarded as stem Zygoptera, i.e. crown Odonata. It follows that Tillyard, in some respect, regarded a subset of Protodonata as stem Odonata (and see Tillyard and Fraser, 1938).

The key point in Tillyard’s thesis from 1925 (Tillyard, 1925b) is the presumed loss of MP and CuA as free veins, beyond the wing base. He derived this view from the study of a remarkable specimen of a *Meganeura*-like species composed of a negative imprint of a hindwing recovered from Elmo (Kansas, USA; Wellington Formation; Artinskian, early Permian; Fig. 1a, b, d, e). Unlike previously known species, vein elevations were preserved in a pristine manner in this material. Tillyard recognized a fusion of M with R, owing to the occurrence, after ScP, of two closely adjoined veins. The resulting vein further splitting into a convex vein (certainly RA), a concave vein (certainly RP) and a convex vein (likely MA), the concave MP had to have vanished. He observed what he believed was a remnant of it (besides R and MA?), basal to the split of RA and RP + MA (red-filled broad arrow on Fig. 1d), but we could not corroborate his observation (only two veins occur). Tillyard also noted a loop, near the wing base, he regarded as formed by CuA and CuP diverging and fusing shortly afterwards. The resulting vein being concave, and therefore presumably composed of CuP, the convex CuA had to have vanished. Finally, a strong oblique structure, his “anal crossing”, located between CuP (his Cu\textsubscript{2}) and AA (his 1A), was not regarded as part of the main veins scheme, despite its strength. This proposal was further formalized in Tillyard and Fraser (1938) and was followed for decades, including prominent textbooks (e.g., Carpenter, 1992; Rohden- dorf, 1962).

This paradigm was challenged by Riek and Kukalová-Peck (1984; and see Riek, 1976), who described two critical, early Pennsylvanian species, *Eugeropteron lunatum* Riek in Riek and Kukalová-Peck, 1984 and *Geroperon arcuatum* Riek in Riek and Kukalová-Peck, 1984, displaying (i) at the wing base, a set four well-delimited veins between ScP and AA (there are three in *Meganeura*-like species, two in extant Odonata), interpreted as RA, RP, M and Cu, successively, and (ii) in the distal area, a complete set of MA, MP, CuA, CuP and AA veins. These insects could be related to Odonata owing to a sequence of vein fusions where they compose the earliest configurations. Under this scheme, a MP + CuA fusion was acquired in early stem Odonata, then further extended into a MP + Cu/CuP fusion. This made Tillyard’s CuP and AA available for being interpreted as the supposedly missing MP and CuA, respectively, and his “anal crossing” interpretable as secondarily fused CuA and CuP, an unusual configuration among *Meganeura*-like species (this stem then fuses with AA; Fig. 1c, f). This evolutionary scheme definitely reconciled the wing venation of Odonata with the serial insect wing venation ground plan: at the wing base, all extant Odonata possess a R + MA stem and a MP + Cu (or MP + Cu + AA) stem. This proposal was applied at the scale of the entire order by Bechly (1996), who also provided an extensive systematic framework derived from the new conjectures of primary homology. These conjectures were corroborated by detailed analysis of wing base structures in extant Odonata (Jacquelin et al., 2018) and are widely accepted nowadays. A recent proposal made by Trueman and Rowe (2019a; and see associated debate; Nel et al., 2019; Trueman and Rowe, 2019b) lacks relevance as a consequence of ignoring *Eugeropteron*-like species, arguably the most important ones to address the question of wing venation homologies in Odonata. 

Aspects of comparatively minor importance remained unclear, notably the actual course of M and MA in *Eugeropteron*-like species. Béthoux (2015) proposed an alternative to Riek and Kukalová-Peck’s interpretation of the four veins located between ScP and AA at the wing base in these odonates, as follows: RA (as opposed to R), MA (as opposed to RP), MP (as opposed to M), and Cu (unchanged). In other words, instead of assuming an R system splitting into RA and RP at the very wing base, as proposed by Riek and Kukalová-Peck, Béthoux assumed that the M system, instead, is split at the very wing base. This is more consistent with the prediction by Riek and Kukalová-Peck that, in extant Odonata, MA is fused with R, and MP with Cu, from the wing base (which implies a very early split of the M system indeed). Another alternative was proposed by Petrulevičius and Gutiérrez (2016), with RA, RP + MA, MP, and Cu composing the four veins of interest, but this implies that both R and M systems are split at the wing base and is therefore a less parsimonious conjecture.

Despite the abundance of Late Palaeozoic strata in Atlantic Canada (Gibling et al., 2019), evidence of fossil insects is rare, with examples preserved in the Pennsylvanian strata at the UNESCO World Heritage site at Joggins, Nova Scotia (Calder et al., 2006; Prokop et al., 2017; Faulkner et al., 2017); Sydney Mines, Nova Scotia (Dawson, 1878; Copeland, 1957); and Fern Ledges, New Brunswick (Matthew, 1909). A single unpublished specimen has been collected from the early Permian strata of Brule, Nova Scotia (Van Allen et al., 2005). Herein we report the discovery of a hindwing from the Sunbury Formation (New Brunswick) displaying a wing venation pattern known to occur in a subset of yet poorly documented stem Odonata. Owing to its combination of character states and its large size, it composes a remarkable addition to the group.
2 Material and methods

The specimen NBMG 21589, which is the focus of this contribution, is deposited at the New Brunswick Museum (palaeontology collection; Saint John, New Brunswick, Canada). The specimen YPM IP 001021a, also illustrated, is housed at the Yale Peabody Museum (New Haven, MA, USA). Specimen numbers associated with the abbreviations MB and WMNM are housed at the Museum für Naturkunde Berlin (Berlin, Germany) and the Westfälisches Museum für Naturkunde Münster (Münster, Germany), respectively.
Photographs of fossil material reproduced in this account were taken using a digital camera Canon EOS 5D Mark III (Canon, Tokyo, Japan) coupled to a Canon 50 mm macro lens (equipped with polarizing filter). The resulting photographs were optimized using Adobe Photoshop CS6 (Adobe Systems, San Jose, CA, USA). The photograph reproduced in Fig. 1 was generated by manually stitching five individual photographs of different areas of the specimen in dry conditions. The photograph reproduced in Fig. 6b is a combination of photographs of the specimen dry and immersed in ethanol (composite photograph).

Two reflectance transformation imaging (RTI) files were generated to further document the specimen NBMG 21589 (for details of this technique as applied to fossil insect imprints see Béthoux et al., 2016). For each file, 42 photographs were taken using a ca. 30 cm diameter automated light dome driving a digital camera Canon EOS 5D Mark III coupled to a Canon MP-E 65 mm macro lens. Original photographs were optimized using Adobe Photoshop CS6 before RTI-processing, itself achieved using the RTIbuilder software (Cultural Heritage Imaging). We provide an online Dryad dataset (Béthoux et al., 2021) containing these two RTI files.

Items composing Fig. 7 were extracted from them using the normals visualisation mode, which assigns a colour code to each pixel according to the orientation of the vector perpendicular to the tangent plane of the object at the corresponding point. These extracts were optimized for contrast.

Hand-drawn sketches were prepared for both YPM IP 001021a and NBMG 21589 specimens with the aid of a Zeiss SteREO Discovery V8 stereomicroscope, equipped with a pair of W-PL 10×/23 eye pieces, a Plan Apo S 1.0× objective, and a drawing tube (all Jena, Germany). Then final vector drawings were prepared using Adobe Illustrator CS6 (Adobe Systems) using the scanned hand-drawn sketches and photographs, as well as RTI extracts for the specimen NBMG 21589. The final drawing of the specimen YPM IP 001021a involved manual adjustments to correct for areas that are creased on the specimen.

We use wing venation homology conjectures proposed by Riek and Kukalová-Peck (1984) and Béthoux (2015) for Odonata. Corresponding abbreviations are as follows: ScP, posterior subcosta; R, radius; RA, anterior radius; RP, posterior radius; M, media; MA, anterior media; MP, posterior media; Cu, cubitus; CuA, anterior cubitus; CuP, posterior cubitus; AA, anterior analis. Colour-coding is as follows: blue, radial system; red, median system; green, cubital system (dark green, CuA; light green, CuP); orange, anal system; purple, concave intercalary vein; yellow, convex intercalary vein.

Before proceeding, it is necessary to define the term “fusion”, as applying to wing veins, as unambiguously as possible. First, “vein fusion” can be understood veins being simply adjoined for some distance or veins sharing their lumen; and, in the latter case, keeping their respective tracheae distinct or fused into a single one. Only the first condition can be clearly inferred in the rock imprint record of insects, and it is the focus of the following. The states where two veins are connected by a cross-vein (Fig. 2a), briefly connected (Fig. 2b), connected (Fig. 2c), or fused (Fig. 2d) can be described by objective values derived from four points (A–D), placed along the median axes of each main vein, as follows: A and B are located along the anterior vein, and C and D along the posterior one; A and C are located at the first point of inflexion of each vein, and B and D at the second point of inflexion of the each vein. The AC (and BD) distance is a, while the AB (and CD) distance is b. In practice, the distance b is negligible or null in cases represented in Fig. 2a–c. However, it sometimes occurs that a vein forms a brief plateau when connected to a particularly thick cross-vein, in which case two inflexion points occur indeed, at the beginning and end of the plateau. Similarly, the distance a is negligible or null in cases represented in Fig. 2c–d. It has a value when the two veins do not fully overlap (i.e. when the fused section is thicker than the individual veins). Fusion is qualified when

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a < b
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3 Geological context

In New Brunswick, the Devonian–Carboniferous strata were deposited within a triangular region that covers roughly 25,900 km² within the approximate 250,000 km² area of the Maritimes Basin (Williams, 1974; Ball et al., 1981; Lavoie et al., 2009; St. Peter and Johnson, 2009; Dietrich et al., 2011; Gibling et al., 2019). Seven geological groups are preserved within the New Brunswick portion of the Maritimes Basin (Gibling et al., 2019), the youngest of these being the Pictou Group. Within this group, the

![Figure 2. Characterization of vein fusion: (a) two veins are connected by a cross-vein, the points A and B overlap, and C and D overlap, distance a is superior to distance b; (b) two veins are briefly connected, the points A and B overlap, and C and D overlap, distance a is slightly superior to distance b; (c) two veins are connected, the points A, B, C and D overlap, distance a equates distance b; (d) two veins are fused, the points A and C overlap, and B and D overlap, distance a is inferior to distance b.](image-url)
oldest strata have been assigned to formations as early as the Bashkirian–Moscovian boundary, which sits conformably above the underlying Bashkirian-aged Cumberland Group, unconformably on older Mississippian sedimentary basins, or unconformably on crystalline basement rocks. The Minto Formation sits at the base of the Pictou Group, within the New Brunswick Platform in the Marysville Subbasin (Dyer, 1926; Muller, 1951; St. Peter and Johnson, 2009). Muller (1951) renamed the Grand Lake Formation of Dyer (1926) to the Pictou Group of Bell (1944) and upgraded the Minto, Hurley Creek, and Sunbury Creek members of the Grand Lake Formation to formation status.

The Minto Formation is defined as a 200–230 m thick section of grey to reddish brown and locally maroon sediments that are exposed in central New Brunswick (Dyer, 1926; Muller, 1951). Palynomorph assemblages analysed by Haquebard and Bars (1970) and Haquebard (1972) indicated that the Minto Formation belongs to the Vestispora Palynomorph Zone, which is of Westphalian C (Moscovian) age, this corroborating the palaeobotany work of Bell (1962).

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The Hurley Creek Formation was defined as a 45–60 m thick stratigraphic package containing abundant red beds that have a coarsening upwards nature. Lithologies are dominantly red and grey mudstones with fine-grained sandstones in the lower strata and red (locally green-grey) coarse feldspathic and lithic sandstones with polymictic conglomerates in the upper strata (Muller, 1951). Finally, the overlying Sunbury Creek Formation was estimated to have a maximum thickness of approximately 90 m and has similar lithologies to the basal Minto Formation (Muller, 1951). This similarity resulted in St. Peter and Johnson (2009, and earlier references therein) informally incorporating the Sunbury Creek into the Minto Formation and downgrading the Hurley Creek Formation to member status. The stratigraphic descriptions of Muller (1951) are followed here.

The specimen NBMG 21589, which is the focus of the current account, was discovered at Robertson Point (Grand Lake, New Brunswick), where a coastal outcrop extends for a distance of 860 m, between 45°52′12.52″ N and 45°52′4.76″ W (Figs. 3, 4). The lower strata are exposed along the east end of the outcrop. A thickness of up to ~6 m of maroon to buff coloured, fine-grained sandstones are deposited in backfilled channel thalwegs that locally incise, or are lateral to, red to maroon mudstones and siltstones that represent interdistributary floodplain deposits with localized palaeosols. Detailed palynology studies have been attempted on these red-bed strata with little success (Duncan McLean and David Bodman, personal communication to Matthew R. Stimson, 2021) and volcanic ash beds are absent. Macrofloral elements of Cordaites principalis, Calamites sp. and cf. Pecopteris suggest a Pennsylvanian age and are consistent with previous workers descriptions of the Hurley Creek Formation fossil content (Muller, 1951). These strata are thus interpreted and mapped as belonging to the Hurley Creek Formation.

This basal red-bed unit is disconformably overlain by ~12 m of dominantly grey to buff unit that is in erosive contact with the underlaying strata. The contact between the two units is exposed for ~500 m along much of the shoreline outcrop due to a very shallow strike and dip of 150/5 SW. The disconformable overlying strata are interpreted as belonging to the Sunbury Creek Formation (Muller, 1951; Allen et al., 2020) and is the stratigraphic equivalent to the uppermost strata of the Minto Formation according to St. Peter and Johnson (2009, and earlier references therein). At Robertson Point, the Sunbury Creek Formation strata are composed of a basal ~6 m thick extraformational polymictic conglomeratic unit with localized coarse sandstone lenses. These strata locally incise the underlaying red bed Hurley Creek Formation up to 2 m deep. The conglomerates are overlain by 2.5 m of buff-coloured coarse to very fine sandstones that locally exhibit trough cross-bedding and sheet sandstone beds. The basal portion of this sandstone locally contains reworked pebbles derived from the underlaying conglomerate. This unit locally incises into the underlaying conglomerates up

Figure 4. Stratigraphic section of uppermost Hurley Creek Formation (6.5 m) and lowermost Sunbury Creek Formation (11.5 m) strata exposed at the Robertson Point outcrop along the shoreline of Grand Lake (New Brunswick).
to an additional 2.5 m that can be traced along the wave cut platform from the main fossil locality, past the odonate wing locality to the end of the outcrop to the west. The specimen NBGM 21589 was discovered as a float on the beach adjacent to its originating outcrop in this sandstone unit at roughly 45°52′9.86″N, 66°5′43.14″W, exposed as a shallow coastal outcrop on the lake’s wave cut platform. The sandstone unit is locally incised by a 2 m heterolithic and interbedded, very fine, trough cross-beded sandstone and siltstone/mudstone shale. A fossil-rich unit is located between 12.0 and 15.0 m from the base of the measured section. It is composed of a 3 m of grey (laterally grades to maroon coloured) siltstone and shale and was found to contain abundant well-preserved plant fossils, abundant vertebrate and invertebrate ichnofossils, and a yet undescribed forewing of an Archimylacridae (i.e. a stem relative of cockroaches and mantids). These strata are interpreted to represent interdistributary floodplains.

Further above, the fossil bearing shales are incised by 4.5 m of channel bodies composed of coarse-grained buff to grey extraformational polymictic conglomerates and coarse-grained sandstones. These channels preserve abundant carbonized plant stems. Incising channels are composed of sandstone and conglomerate which are resistant to erosion compared to the laterally deposited shales. These channel deposits form localized coastal headlands where the shales have been eroded deeper along the shoreline and expose the sandstone channel margins in the cliff face. Incising and overlaying channel deposits measure up to 6.5 m in thickness (including 3 m incisions into underlying strata). A localized 1.5 m thick maroon-coloured mudstone-filled channel with 5–10 cm very fine sandstone lenses incises the uppermost sandstone units.

The Pictou Group spans much of the latest Pennsylvanian and earliest Permian periods (St. Peter and Johnson, 2009, and references therein). The Minto Formation is thought to be of latest Bashkirian age, while the overlying Hurley Creek Formation straddles the Bashkirian–Moscovian boundary. The overlying Sunbury Creek Formation has traditionally been considered to be allostratigraphically of Moscovian age (St. Peter and Johnson, 2009).

4 Comparative analysis

To ease the systematic placement of the new species described below, we carried out a survey of the configurations involving MP, the cubital system, and AA in early stem Odonata (Fig. 5), inspired from Riekk and Kukalová-Peck (1984) and Brauckmann and Zessin (1989), and considering recent discoveries. The wing morphology of *Eugeropteron lunatum* is widely regarded as a ground plan for the entire Odonata. As far as MP, the cubital system, and AA are concerned, it is characterized by the absence of fusion between MP and CuA, between CuA and CuP (after both veins had diverged), and between CuP and AA (Fig. 5a). Instead, in each instance, the involved veins are connected by a short cross-vein. At the point of connection with these cross veins, veins commonly form a sharp angle. The morphology of *Erasipteron larisch* Pruvost, 1933 (based on original description, Brauckmann and Zessin, 1989, and a new photograph of the holotype (bearing the specimen number MB.I.455, Museum für Naturkunde Berlin), Andreas Abele, personal communication to Olivier Béthoux, 2020), known from a single forewing, is schematized in Fig. 5c. With respect to the previous configuration, it is the outcome of multiple fusions (Fig. 5b), whose sequence of acquisition is not evident. The case is straightforward if considering the holotype of *Argentinaria cristinae* Petrulevičius and Gutiérrez, 2016, in which both wing pairs lack the CuP + AA fusion, suggesting that it was acquired last. This specimen also suggests that a delayed acquisition of derived conditions in the hindwing, with respect to the forewing, is a plausible phenomenon: while MP and CuA are fused for a long distance in the forewing, these two veins are only connected (or briefly connected) in the hindwing, and CuA and CuP are briefly fused in the forewing, but remain distinct in the hindwing. This configuration suggests that the first fusion to occur was the MP + CuA one, as it is the only fusion shared by both wing pairs. It follows that the CuA + CuP fusion came second. This acquisition sequence (MP + CuA, then CuA + CuP, then CuP + AA) is supported by the configuration in both *Kirchnerala treintamil* Petrulevičius and Gutiérrez, 2016 and *Aulertupus tenbrocki* Zessin and Braunckmann, 2010, each known from a forewing in which the CuP + AA fusion is lacking, while the two others are present. However, a notable exception is *Geropteron arcuatum*, in which the only fusion to lack is the MP + CuA one. The scheme represented in Fig. 5c corresponds to a taxon that can be characterized as displaying the derived character state “occurrence of a MP + CuA fusion, of a CuA + CuP fusion (after both
veins had first diverged) and of a CuP + AA fusion” (as opposed to “in both wing pairs, MP and CuA distinct, CuA and CuP distinct, and/or CuP and AA distinct”), applying to both wing pairs.

In both wing pairs, Namurotypus sippeli Brauckmann and Zessin, 1989 displays a MP + CuA fusion extending more basally than in the species considered above. This is best documented based on the holotype (specimen P28957 WMNM – formerly N 1000), in which MP fuses with Cu at the point where CuA and CuP diverge (as represented in Fig. 5e) in the forewing, and shortly after it in the hindwing (based on original description and new photographs, Lothar Schöllmann, personal communication to Olivier Béthoux, 2021). As a consequence, the basal free portion of CuA (dark-green-filled arrow on Fig. 5c) no longer occurs (Fig. 5e), at least in the forewing. Concurrently, this species retains, in both wing pairs, the free portion of CuP occurring distal to the CuP–AA split and basal to its fusion with CuA (hereafter “median free portion of CuP”; light-green-filled arrows on Fig. 5c, e). As pointed out by Nel et al. (2009), the further completion of the fusion of MP with Cu is apparently highly homoplastic among Palaeozoic and Triassic stem Odonata. The scheme represented in Fig. 5e corresponds to a taxon that can be characterized as displaying the derived character state “MP fused with the cubital system at the point where CuA and CuP diverge, or basal to it” (as opposed to “MP fused with the cubital system distal to the point where CuA and CuP diverge”), applying to both wing pairs.

The configuration in Erasipteroides valentini Brauckmann in Brauckmann et al. (1985) differs only slightly from that of Namurotypus sippeli. In the specimen P26133 WMNM (based on Bechly et al., 2001, and new photographs, Lothar Schöllmann, personal communication to Olivier Béthoux, 2021), presumably belonging to this species, both fore- and hindwings conform to the configuration represented in Fig. 5e. Note that in the holotype (specimen P30361 WMNM) the CuP + AA fusion extends slightly beyond the point where CuA fuses with CuP, with the consequence that CuP has no free median portion (based on new photographs, Lothar Schöllmann, personal communication to Olivier Béthoux, 2021).

The configuration in Erasipterella jini (Zhang, Hong and Su in Su et al., 2012) (see original description and Li et al., 2013), known from both fore- and hindwings, suggests that the extension of the MP + CuA fusion (into a MP + Cu/CuA fusion) was acquired before that of the CuP + AA fusion (into a (CuP/CuA + CuP) + AA fusion; Fig. 5f). While no traces of the free basal portion of MP are present in this species (i.e. MP is fully fused with Cu from the wing base or close to it), several of the known hindwings retain a free median portion of CuP. In other words, as far as AA and CuP / CuA + CuP are concerned, it conforms to the scheme represented in Fig. 5e. Concurrently, the forewing of Erasipterella jini displays the “extended version” of the AA + CuP fusion, resulting in a loss of the free median portion of CuP, as represented in Fig. 5g (with the possible exception of the holotype). Note that, besides information on the sequence of character state acquisitions, this species composes another example suggesting that derived traits were acquired first in the forewing and later in the hindwing in the early evolution of Odonata.

Most Meganeura-like species display the configuration represented in Fig. 5g (Nel et al., 2009; Li et al., 2013; Zessin, 1983; among others). Megatypus schucherti (Fig. 1c, f) is peculiar in that the median free portion of CuA and the basal free portion of CuP (dark- and light-green-bordered arrows, respectively, in Fig. 5c, e, g) are fused. The scheme represented in Fig. 5g corresponds to a taxon that can be characterized as displaying the derived character state “AA fused with the cubital system until the point where the free median portions of CuA and CuP fuse, or distal to it” (as opposed

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to “AA diverging from CuP + AA basal to the point where the free median portions of CuA and CuP fuse”), applying to both wing pairs.

Relevant character states having been identified, a systematic framework based on these could have been elaborated by resorting to cladotypic nomenclature or PhyloCode apomorphy-based definitions. However, it proved difficult to find appropriate reference specimens (cladotypes or internal specifiers). Most importantly, the apparent heterobathmy of character state acquisition between fore- and hindwings would have required character state labels to mention the particular organ(s) of concern (viz. “in both wings pairs” or “at least in forewing”). However, this option was difficult to implement because the morphologies of both fore- and hindwing bases are known in very few species of early stem Odonata. Another consequence of this lack of data is that the actuality of the “forewing antecedence” phenomenon is yet to be further verified. Additionally, some degree of intra-specific variability is to be expected, making it hazardous to designate species known from a very limited material as reference. Finally, the basal free portions of CuA and CuP (green-bordered arrows on Fig. 5c, e, g) can prove difficult to locate in Meganeura-like species, where they tend to resemble cross-veins. In turn, it makes it difficult to assess the position of AA with respect to them, necessary to qualify the presence of the configuration represented in Fig. 5g.

5 Systematic description

Order Odonata Fabricius, 1793
Taxon Neodonatoptera Bechly, 1996
Family incertae sedis
Brunellopteron Béthoux, Deregnaucourt and Norrad gen. nov.
(type species: Brunellopteron norradi Béthoux, Deregnaucourt and Norrad sp. nov.)

Type species
Brunellopteron norradi Béthoux, Deregnaucourt and Norrad sp. nov.

Gender
Masculine.

Etymology
The genus name honours Paul-Michael Brunelle (1952–2020), an odonatologist who contributed greatly to our knowledge on odonate biodiversity in the Atlantic Canadian region (see McAlpine, 2020).

Diagnosis
By monotypy, as for the type species.

Discussion
Based on delimitations given by Bechly et al. (2001) the type species of the new genus can be assigned to the taxon Neodonatoptera. Within this taxon, the lack of an archae-dictyon, one of the proposed apomorphies of the taxon Euodonatoptera and present in the new material, would require a reappraisal. Notably, (i) the lack of an archae-dictyon is regarded as homoplastic, and (ii) the occurrence of an archae-dictyon among stem Odonata, in particular in Erasip teron larisch i, is not evident. Moreover, according to Bechly et al. (2001), the monophyly of the taxon regarded as a sister group of Euodonatoptera, namely Eomeganisoptera (including the Erasipteridae), remains to be verified. The length of ScP has also been regarded as a relevant character by Bechly et al. (2001), but (i) it is possibly homoplastic and (ii) it is difficult to evaluate in the available material of the type species of the new genus. Considering that all Euodonatoptera conform to one of the conditions represented in Fig. 5e and g, and that the type species of the new genus does not (it conforms to the condition represented in Fig. 5c), we propose to exclude the new genus from this taxon.

Brunellopteron norradi Béthoux, Deregnaucourt and Norrad, sp. nov.
(urn:lsid:zoobank.org:act:F29F7AC2-5624-4786-B842-7CD22C0E4B3E)
(Figs. 6, 7)

Material
Holotype, specimen NBMG 21589, a positive imprint of a left hindwing, deposited at the New Brunswick Museum (palaeontology collection; Saint John, New Brunswick, Canada).

Etymology
The species epithet composes a dedication to Donald B. Norrad, co-discoverer of the holotype (and father of the second author).

Diagnosis
Hindwing: area between anterior wing margin and ScP broad; MP branched, or area between MP and CuA filled with alternating convex and concave intercalaries; several cross-veins connecting (i) MP and the median free portion of CuA, (ii) the basal free portion of CuA, MP + CuA and the median free portion of CuA on one hand, and the basal free portion of CuP on the other, and (iii) the basal free portion of CuP and AA; area between AA and the posterior wing

Figure 6. Brunellopteron norradi Béthoux, Deregnaucourt and Norrad gen. et sp. nov., holotype (NBMG 21589, New Brunswick Museum; positive imprint of a left hindwing): (a) drawing of visible structures, with indication of convex, weathered portions; (b) photograph (dry-ethanol composite, flipped horizontally), with indication of the location of items in Fig. 7; (c) same as in (a) but with portions reconstructed based on convex, weathered portions and the morphology of known related species; (d) same as in (c) with colour-coding and legend (◦ indicates a particular concave intercalary vein; see text; ∗ indicates an intercalary of second order).
margin well developed, filled with numerous concave intercalary veins; occurrence of intercalaries of second order (i.e. having the same elevation as that of the main vein delimiting the corresponding area).

**Description**

Holotype specimen (NBMG 21589): basal half of a hind-wing, positive imprint, most convex veins eroded; length about 93 mm as preserved, 145–155 mm as estimated; maximum width 36.9 mm as preserved, 38.6 mm as estimated; area between anterior wing margin and ScP broad in basal area; R (or R + MA) not preserved but position of basal portion inferable based on slope elevation; course of basal portion of MA not preserved, either fused with R from the wing base or more distally (dotted lines in Fig. 6d); origin of RP + MA and split of this vein (into RP and MA) not preserved; RP rectilinear, with a fork about 66 mm distal to wing base; MA (distal to the RP–MA split) not preserved but a short portion of its course, opposite the posterior bending of MP, inferable based on slope elevation; MP fusing with CuA for 2.3 mm, then diverging from MP + CuA somewhat obliquely, directed anteriorly; then, MP with a marked posterior arching, inflexion point 27.5 mm distal of the MP–CuA split; first MP fork (or, first concave intercalary arising) 54.3 mm distal of the MP–CuA split; area between the anterior stems of MP and CuA broad; Cu with a distinct basal stem splitting into CuA and CuP; CuA 3.5 mm long before it fuses with MP; CuA portion distal to the split of MP + CuA (into MP and CuA) about 7.4 mm long before it fuses with CuP (itself emerging from CuP + AA); CuA + CuP fusion about 3.0 mm long; distal to the CuA–CuP split, courses of the anterior stem of CuA and of its first posterior branch inferable based on slopes elevations; anterior-most branch of CuA with a posterior arching, more or less opposite to that of MP; basal portion of CuP 9.7 mm long before it fuses with AA; CuP presumably with a short free portion distal to the CuP-AA split and basal to its fusion with CuA; CuP posteriorly pectinate, with 5 terminal branches; course of the anterior stem of AA inferable, to some extent, based on slopes elevation; AA fused with CuP for about 2.0–2.5 mm; in the area between AA and the posterior wing margin, course of actual AA branches barely preserved; cross-venation generally scalariform, forming intercalary veins of elevation opposed to that of surrounding main veins, at least within the CuA, CuP and AA areas; MP and median free portion of CuA, median free portion of CuA and basal free portion of CuP, and basal free portion of CuP and AA connected by several cross-veins; at least in the CuP area, occurrence of an intercalary vein of second order (bordered by two intercalary veins; * in Fig. 6d).

**Locality and horizon**

Robertson Point, Grand Lake, New Brunswick, Canada; Sunbury Creek Formation.

**Discussion**

The nearly complete lack of preserved convex elements (main veins or intercalaries), a consequence of weathering, could be partly circumvented by considering the elevation of the preserved slopes. For example, the course of the basal portion of AA can be easily traced (Fig. 7a). The reconstruction of the course of AA distal to its fusion with CuP was elaborated considering that the concave element indicated by ◦ on Fig. 6d is separated from CuP by an elevated section (Fig. 7a), and therefore it cannot be a branch of CuP. It is therefore a concave intercalary branch located within the AA area. It follows that the elevated section must be AA. Then, given the orientation of the basal and inferred distal portions of AA, the occurrence of a short, free portion of CuP (emerging from CuP + AA and basal to its fusion with CuA) was in-
ferred, but its reconstructed length is speculative. The course of several other portions of convex elements remains more or less speculative. Notably, whether MA fuses with R at the wing base, or more distally, cannot be appreciated. In some cases, conditions known in related species allowed plausible reconstruction, such as the length of the RP + MA stem. Finally, portions of a concave element were observed between genuine CuP branches (\(*\) on Fig. 6d; purple-filled broad arrows on Fig. 7b). It is delimited by two elevated areas, interpreted as remains of convex intercalaries. It follows that this concave structure is an intercalary sharing the same elevation as that of the main vein delimiting the corresponding area (here, CuP), this qualifying an intercalary of second order. The occurrence of a comparatively broad area between AA and the posterior wing margin makes it patent that it is a hindwing.

Based on the above, the new specimen displays the configuration schematized in Fig. 5c. Specifically, it displays the derived character state “occurrence of a MP + CuA fusion, of a CuA + CuP fusion (after both veins had first diverged) and of a CuP + AA fusion” (as opposed to “in both wing pairs, MP and CuA distinct, CuA and CuP distinct, and/or CuP and AA distinct”). As a consequence, close relationships with the following species can be excluded: *Eugeropteron lunatum*, *Geropteron arcuatum*, *Tapacala niuymenae Petrulevičius and Gutiérrez, 2016, Kirchnerala treintamil*, and *Aulertus tembrocki*. Conversely, the new species clearly displays a basal free portion of CuA (dark-green-filled arrow on Fig. 5c), a plesiomorphy indicating that close relationships with *Erasipteroides valentini* and *Euodonatoptera* (as delimited by Bechly et al., 2001) can be excluded.

The remaining species are *Erasipteron larischi*, known from a single forewing whose apex is missing, and *Zessinella siope* Brauckmann, 1988, known from body remains associated with fore- and hindwings whose venation could be deciphered at their bases only. *Piesbergtupus hielscheri* Zessin, 2006, known from the basal half of a single forewing, is also considered as it likely displays a free median portion of CuP and, possibly, a free basal portion of CuA (Fig. 8). The same applies to *Gallotupus oudardi* Nel, Garrouste and Roques, 2008, known from a single, almost complete forewing, creased in several areas.

*Erasipteron larischi* and *Zessinella siope* are herein regarded as essentially similar. Both species have a similar size (forewing about 15 mm wide in the former, about 9 mm in the latter), comparatively large cells, and a very short CuP + AA fusion (at least in the forewing). The main difference is, in the forewing, the number of rows of cells in the area between AA and the posterior margin (two in the former, one in the latter), a trait which could be a mere consequence of the size difference. Interestingly, while the three typical fusions (MP + CuA, CuA + CuP and CuP + AA) occur in the forewing of *Zessinella siope*, it is not evident that it is the case in the hindwing, for which a possible interpretation is that CuP and AA are connected by a cross-vein. This should be further investigated, but if this were the case, the new specimen would then strongly differ from *Zessinella siope* and, by extension, from *Erasipteron larischi*. Additionally, intercalary veins are absent in the area between AA and the posterior wing margin in the forewing of these two species (condition unknown in hindwing). Even though comparison with the new specimen is hindered by the fact that it is a hindwing, given the important development of intercalary veins in this area, it can be reasonably assumed that such structures occurred in the corresponding forewing. Notably, such intercalary veins occur in the known forewing of *Gallotupus oudardi*.

The occurrence of a seemingly branched MP in the new specimen is a very unusual feature among Odonata. This vein is simple in *Erasipteron larischi* (the distal half of MP is unknown in *Zessinella siope* and *Piesbergtupus hielscheri*) and in virtually all Odonata indeed. Among Euodonatoptera, the Permian and Triassic Zygophlebiida possess a wide area between MP and the anterior stem of CuA that is filled with alternating convex and concave vein-like structures. However, the latter, which could be regarded as MP branches, are believed to be intercalaries instead (see Nel et al., 2001, and references therein). Such organization can also be found in some extant forms, for example some Polythoridae (see Garrison et al., 2010). Then, it cannot be completely ruled out that it is also the case in the new specimen (i.e. that it possesses a simple MP and alternating convex and concave intercalaries between MP and CuA). This condition might also be present in *Gallotupus oudardi*. Yet, the character state “area between MP and CuA broad, filled with intercalary veins” would allow differentiating the species the new specimen belongs to from *Erasipteron larischi* (and, by extension, *Zessinella siope*).

One of the peculiarities of *Piesbergtupus hielscheri* is the straight to slightly concave shape of its posterior wing margin in the basal third of the forewing (Fig. 8). In Odonata, but also in many Megasecopteromorpha, this trait is part of a petiolation syndrome that very generally occurs in both fore- and hindwing. Judging from the development of the AA area in the new specimen, the species it belongs to most likely...
had forewings with a broad base or a very short petiole at best. This excludes close relationships with *Piesbergtupus hielscheri*. The new specimen is also larger than this species (estimated lengths 145–155 mm vs. 100 mm).

It occurs that *Gallotupus oudardi* is possibly the closest relative of the species to which the new specimen belongs. Further comparison is made difficult because several critical areas are creased in the only known specimen of the former. In particular, the point where MP fuses with the cubital system (either with CuA or Cu) is unknown. Judging from the distance between the point of fusion of CuA and CuP on one hand, and the first fork of RP on the other, preserved in both specimens, the new one is about 1.6 times as long as the known wing of *Gallotupus oudardi*, this precluding an assignment of the new specimen to this species.

Finally, a trait absent in the species considered above is the occurrence of several cross-veins connecting (i) MP and the median free portion of CuA, (ii) the basal free portion of CuA, MP + CuA and the median free portion of CuA on one hand, and the basal free portion of CuP on the other, and (iii) the basal free portion of CuP and AA (Figs. 6, 7a). This trait is present in *Aulertupus tembrocki*, presumably remotely related to the species the new specimen belongs to, as it lacks the CuP + AA fusion (see above).

Observed differences between the new specimen and known species legitimate the erection of a new species and of a new genus.

6 Conclusions

*Brunellopteron norradi* belongs to a subset of early stem Odonata known from very few species, themselves known from few and mostly incomplete specimens. The corresponding material was recovered from western European localities of Moscovian age (Westphalian C/D), which corroborates the age of the Robertson Point outcrop as deduced from palynomorph content. Despite its incompleteness, the new species remains a useful addition to our knowledge of the early evolution of Odonata wing venation, owing to its unique combination of character states. In particular, it is the only *Erasipteron*-like species to possess such an important development of intercalary veins, more typical of the large *Megaeneura*-like griffinflies, most of which occur in strata younger than the Moscovian.

**Data availability.** The new material included in the paper is accessible and deposited at the New Brunswick Museum (palaeontology collection; Saint John, New Brunswick, Canada), and all data are included in the description. RTI files from which images reproduced in Fig. 7 were extracted are available from https://doi.org/10.5061/dryad.ftdz08s0 (Béthoux et al., 2021).

**Author contributions.** OB and MRS elaborated the project. All authors contributed to redacting the current account: OB as main contributor of the comparative analysis section; OB and ID as main contributors of the systematic section (including specimen documentation and preparation of illustrations); MRS, OAK, LFA, REN, and SJH as main contributors of the geological context section. LFA and REN contributed to collecting the new material.

**Competing interests.** The authors declare that they have no conflict of interest.

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