Morphology of the oldest fossil subfamily of Limoniidae (Diptera, Architipulinae) in the light of exceptionally preserved Mesozoic material

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Based on known fossil evidence the extinct subfamily Architipulinae is considered to be the oldest evolutionary group of the Limoniidae, the largest family within the infraorder Tipulomorpha. The morphology of this subfamily, which includes 11 genera, has so far been based mainly on wing venation. New well-preserved representatives of the genus Cretolimonia Kalugina, 1986 were recovered from the Jurassic/Cretaceous boundary of Shevia and Daya, Transbaikalia, as well as from mid-Cretaceous amber from Kachin, Myanmar. This new material enriches our knowledge of the subfamily Architipulinae and of the genus Cretolimonia, and allows us to ascertain the detailed morphological structure of the female copulatory apparatus with spermathecae and the structure of the male hypopygium. The combination of detailed impression fossils with a specimen preserved three-dimensionally in resin has permitted study of the morphology of this Mesozoic fly genus almost to the level of modern genera. The paper includes descriptions of four new species of Cretolimonia: C. lukashevichae sp. nov., C. pseudojurassica sp. nov., C. dayana sp. nov. from sedimentary rocks, and C. mikolajczyki sp. nov. from Myanmar amber, supported with a key to all known species.

The Limoniidae (limoniid crane flies), with over 10,000 described extant species, is the largest family in the dipteran infraorder Tipulomorpha, and one of the largest among all Nematocera. It is currently divided into seven subfamilies, three of which are extinct (Architipulinae, Eotipulinae, Drinosinae), and four extant: Limnophilinae, Chioneinae, Dactylolabinae, and Limoniinae (Fig. 1). Based on the fossil data known to date Architipulinae is considered the oldest group of Limoniidae1–7, with the oldest representative Architipula youngi Krzemiński, 1992 dating from the Late Triassic of North America3. This species was used to calibrate the age of the Tipulomorpha clade in the phylogenetic tree of Diptera8. During the Jurassic, the family Limoniidae, including Architipulinae, underwent rapid radiation expressed in an abundant and diverse assemblage of genera and species3,7,9–16. In the Cretaceous, Architipulinae gradually became extinct, while other limoniid families had appeared in that time.

The subfamily Architipulinae includes 11 fossil genera: Architipula Handlirsch, 19061, Protopulula Handlirsch, 19066, Mesotipula Handlirsch, 192015, Paratipula Bode, 195318, Haplotipula Bode, 195318, Leptotipula Bode, 195318, Ozotipula Bode, 195318, Microtipula Bode, 195318, Cretolimonia Kalugina, 19862, Grimmenia Krzemiński and Zessin, 199019 and Metarchilimonia Blagoderov and Grimaldi, 20079. Up until now, knowledge of the morphology of the subfamily has been based mainly on wing impressions in sedimentary rocks. No representatives of Architipulinae were known from amber and no male genitalia had been found. Similarly, the second fossil subfamily, Eotipulinae, is known mainly from wing impressions in Jurassic and Cretaceous sedimentary rocks2,7, although a female has been described from Lower Cretaceous Spanish amber20. In contrast, in the third fossil subfamily, Drinosinae, the morphology of both sexes has been elucidated through material preserved in mid-Cretaceous Myanmar amber21.
The Architipulinae material described here shows an exceptional level of preservation, allowing us to augment the diagnosis of the genus *Cretolimonia* and enriching our knowledge of the whole subfamily Architipulinae. The hypopygium (a modified abdominal segment associated with the genitalia and having a clasping function in the males) is described for the first time on the basis of a perfectly preserved imprint in the rock, as well as the first representative of the subfamily from a fossil resin. Four new species of *Cretolimonia* are described, and a key to all known species is provided.

**Results**

**Systematic palaeontology.**

Order Diptera Linnaeus, 1758  
Infraorder Tipulomorpha Rodendorf, 1961  
Family Limoniidae Speiser, 1909  
Subfamily Architipulinae Handlirsch, 1906  
Genus *Cretolimonia* Kalugina, 1986

**Type species** *Cretolimonia popovi* Kalugina, 1986: figs. 87a, b: Gurvan-Ereny-Nuru (West Mongolia), Early Cretaceous. Genus description based on a wing fragment.

**Species included** Table 1.

**Amended diagnosis** The genus is distinguished from all Limoniidae by the characteristic wing venation pattern: vein Sc ends near Rs bifurcation, four radial veins present (R₁, R₂, R₃, and R₄), cross-vein r-r (R₅) atrophied, and vein R₅ very short, slightly curved; basal medial vein (Mb) well visible, long, all four medial veins present, d-cell closed, cross-vein m-cu located at the distal part of the d-cell base or sometimes in the middle (*C. excelsa*).
ovipositor short, strongly curved dorsally; three small spermathecae present; male hypopygium with outer gonostylus strongly hooked; inner gonostylus lobed; aedeagus narrow and slightly curved, parameres large, triangular, very dilated at base.

**Remarks** The genus *Cretolimonia* was included in the subfamily Architipulinae based on complex features which define this subfamily, i.e.: four radial and four medial veins reaching the wing margin, closed d-cell; cross vein m-cu located at the end of d-cell. The absence of cross-vein r-r (R2) distinguishes the genus *Cretolimonia* among other genera of the subfamily Architipulinae. So far, four species from the Middle Jurassic to mid-Cretaceous have been included in the genus (Table 1). For the purpose of the key, their wings are shown in Fig. 2A-D.

### Key to species in genus *Cretolimonia*:

1. wing narrow, at least 3x longer than width .......... 2
   - wing wide, about 2.5x longer than width .......... 7
2. Sc ends far beyond the Rs fork ........... 3
   - Sc ends before or opposite the Rs fork ........... 4
3. M1 twice as long as upper edge of d-cell (Fig. 2A) ........... *C. popovi* Kalugini, 19862
   - M1 only slightly longer than upper edge of d-cell (Fig. 2B) ........... *C. excelsa* Gao et al., 2015 23
4. Sc ends opposite Rs fork ........... *C. dayana* sp. nov.
   - Sc ends in front of the Rs fork ........... 5
5. d-cell very small, no more than 1/10 of the wing length (Fig. 2C) ........... *C. jurassica* Lukashevich, 20097
   - d-cell large, no more than 1/7 of the wing length ........... 6
6. R5 almost equal to the length of R2+3+4 ........... *C. pseudojurassica* sp. nov.
   - R2+3+4 about 1/3 longer than R5 (Fig. 2D) ........... *C. pygmaea* Lukashevich, 2009 7
7. petiola longer than M1; m-cu almost 2/3 along lower part of d-cell ........... *C. lukashevichae* sp. nov.
   - petiola only about 1/3 as long as M1; m-cu just before the M3+4 bifurcation ........... *C. mikolajczyki* sp. nov.

### Table 1. List of species of *Cretolimonia* known from the fossil record.

| Species | Time scale | Type of material | Locality |
|---------|------------|-----------------|----------|
| *Cretolimonia popovi* Kalugina, 19862 | Early Cretaceous | Imprint | West Mongolia |
| *Cretolimonia jurassica* Lukashevich, 20097 | Late Jurassic | Imprint | Mongolia, Shar Teg |
| *Cretolimonia pygmaea* Lukashevich, 20097 | Late Jurassic | Imprint | Mongolia, Shar Teg |
| *Cretolimonia excelsa* Gao et al., 2015 23 | Middle Jurassic | Imprint | Inner Mongolia, Daohugou |
| *Cretolimonia lukashevichae* sp. nov. | Early Cretaceous | Imprint | Transbaikalia, (Shevia) |
| *Cretolimonia pseudojurassica* sp. nov. | Jurassic/Cretaceous boundary | Imprint | Transbaikalia, (Shevia) |
| *Cretolimonia dayana* sp. nov. | Jurassic/Cretaceous boundary | Imprint | Transbaikalia, (Daya) |
| *Cretolimonia mikolajczyki* sp. nov. | mid-Cretaceous | Amber | Northern Myanmar |

### Figure 2. Wing venation of previously described species from genus *Cretolimonia* redrawn from original papers: (A) *Cretolimonia popovi* Kalugini, 1986; (B) *Cretolimonia excelsa* Gao et al., 2015; (C) *Cretolimonia jurassica* Lukashevich, 2009; (D) *Cretolimonia pygmaea* Lukashevich, 2009.
Cretolimonia dayana Kopeć n. sp.
(Fig. 3-4)

Etymology  The name was established from the site where this species was discovered.

Material examined  Holotype No. 3063/1208, female (Fig. 3A,B). Additional material: 3063/399, female; 3063/1072, female; 3063/1079, male (Fig. 3C,D). Specimens come from Daya (Transbaikalia, Russia), Jurassic/Cretaceous boundary, housed in the Borissiak Palaeontological Institute, Russian Academy of Sciences, Moscow, Russia (PIN).

Diagnosis  Wing narrow, about 3.3 times longer than its width; venation differs in the proportions of the individual veins from all species in this genus. Sc ends opposite the fork of Rs, d-cell long, narrow, expanded at distal part, no more than 1/9 of the wing length; m-cell lies at 2/3 length of the base of the d-cell; gonocoxites short, broad; outer gonostylus short, strongly chitinized, significantly hooked at end; inner gonostylus delicate, lobed; aedeagus long, narrow, slightly curved; parameres shorter than penis, broad at base.

Description  Wing length ca. 4.6 mm, width 1.4 mm (Fig. 4A); Head. Antennae with 16 segments, scapus tubular, half as long as its width, pedicel barrel-shaped, only slightly wider than scapus; flagellum with 14 flagellomeres, flagellomeres ovoid, gradually shortening, basal segment of flagellum expanded in lower part, last segment small and round, on all flagellomeres there are bristles shorter than width of segment on which they are located, on last segment 3 or 4 short bristles; clypeus 4-segmented, short, and the last segment almost equal in length to the penultimate one (Fig. 4B). Thorax. Wing narrow, about 3.3 times as long as its width; vein Sc ends opposite the Rs fork; cross-vein sc-r is about twice its length before the end of Sc; R1 ends opposite the R2+3+4 bifurcation at R3 and R4, and R2 (r-r) completely disappears; Rs about 1/4 longer than R3+3+4; R3 short, strongly sickle-shaped, slightly inclined, less than 1/3 length of R3+3+4; R4 about 1/4 shorter than Rs and equal in length to R3+3+4; four medial veins present, petiolar ca. 1/4 longer than M4 and 1/3 longer than upper edge of d-cell; d-cell long, ca. 1/9 of wing length; cross-vein m-cu lies at 2/3 length of d-cell base; A2 vein not visible; legs long and delicate, tibial spurs absent. Abdomen. Male. Gonocoxites short, broad; outer gonostylus short, strongly chitinized, strongly

Figure 3. Cretolimonia dayana sp. nov.: (A) habitus of female, holotype No. 3063/1208; (B) female abdomen, No. 3063/399; (C) habitus of male, No. 3063/1079; (D) male hypopygium, No. 3063/1079.
hooked at the end; inner gonostylus delicate, lobed; penis long, narrow, slightly sigmoidally curved; parameres broad at base, shorter than aedeagus (3D,4C).

**Female.** Ovipositor short, slightly curved dorsally; three small spermathecae (Fig. 3B, 4D).

**Remarks** The male hypopygium is very well preserved in specimen No. 3063/1079 (Fig. 3D), being only slightly deformed by elongation of the left gonocoxite during fossilization. In previously described species of this genus the copulatory apparatus has not been preserved. The female ovipositor (Figs. 3B, 4D) is preserved almost perfectly in specimen No. 3063/399 with three small spermathecae present and well visible. The spermathecae are identical to those of most species in the subfamily Limnophilinae.

*Cretolimonia lukashevichae* Kopeć and Krzemiński n. sp. (Fig. 5A,C 6A)

**Etymology** The species name is dedicated to Dr. Elena D. Lukashevich, who is involved in the study of fossil Diptera.

**Material examined** Holotype No. 3795/637 sex unknown (Fig. 5A); paratype No. 3795/611 (sex unknown, hind wing preserved), Shevia (Transbaikalia, Russia), Jurassic/Cretaceous boundary; housed in the Borissiak Palaeontological Institute, Russian Academy of Sciences, Moscow, Russia (PIN).

**Diagnosis** Wing broad, 2.5 times as long as its width; Sc ends distinctly beyond fork of Rs; petiola more than half as long as M₁ and 1/4 as long as upper margin of d-cell; m-cu lies nearly at 2/3 length of lower part of d-cell.

**Description** Wing length 5.0 mm (right wing), width 2.0 mm, wing 2.5 times longer than its width (Fig. 5C, 6A); Sc ends distinctly beyond the Rs fork; cross-vein sc-r twice its length before the end of Sc; R₁ ends before the R₃₄,₅ bifurcation at R₃ and R₅; R₃ completely atrophied; Rs ca. 1/4 longer than vein R₃₄,₅; R₅ short, almost vertical, constitutes ca. 1/3 length of R₃₄,₅; R₆ ca. 1/4 shorter than R₃₄,₅; four medial veins present, petiola half longer than M₁ and 1/4 longer than upper edge of d-cell; d-cell trapezoidal, relatively large, constituting 1/7 of wing length, cross-vein m-cu lies at 2/3 length of base of d-cell, far before bifurcation of M₃₄ into M₅₆ and M₇₈; A₁ vein almost straight.
Figure 5. Holotypes of new *Cretolimonia* species: (A, C) *Cretolimonia lukashevichae* sp. nov. No. 3795/637, (A) habitus of holotype, (C) wing; (B, D) *Cretolimonia pseudojurassica* sp. nov. No. 3795/628, (B) habitus of holotype, (D) wing.

Figure 6. Holotypes of new *Cretolimonia* species: (A) *Cretolimonia lukashevichae* sp. nov., No. 3795/637; (B) *Cretolimonia pseudojurassica* sp. nov., No. 3795/623—wing venation.
Remarks The holotype specimen shows two well-preserved wings and a fragment of the thorax. The left wing is longer than the right one, most probably deformed (elongated) during the fossilization process. Such a phenomenon has been observed in different groups of insects\(^2\)\(^3\)\(^4\). The right wing retained its normal structure.

*Cretolimonia pseudojurassica* Krzemiński sp. nov.
(Fig. 5B,D 6B)

**Etymology** The specific name emphasizes the similarity to another species in the genus, *Cretolimonia jurassica* Lukashevich, 2009\(^7\).

**Material examined** Holotype No. 3795/628 (Fig. 5B), sex unknown, only a single wing preserved. Additional material: 3795/633; 3795/638. Shevya (Transbaikalia, Russia), Jurassic/Cretaceous boundary. Housed in the Boris-siak Palaeontological Institute, Russian Academy of Sciences, Moscow, Russia (PIN).

**Diagnosis** Wing narrow, nearly 3.3 times as long as its width; vein Sc ends distinctly before fork of Rs; R\(_4\) equal in length to R\(_2+3+4\); four medial veins present, and vein M\(_1\) 2.5 times longer than petiola and equal in length to upper margin of d-cell.

**Description** Wing well-preserved with clearly visible veins, 5.1 mm long and 1.6 mm wide, almost 3.3 times as long as its width (Figs. 5D, 6B); Sc ends distinctly before bifurcation of Rs; cross-vein sc-r about its length before end of Sc; R\(_4\) ends before bifurcation of R\(_2+3+4\); R\(_1\) ends before bifurcation of R\(_2+3+4\) into R\(_3\) and R\(_4\); Rs about 1/4 longer than R\(_2+3+4\); R\(_2\) short, almost parallel to R\(_4\); ca. 1/3 length of R\(_2+3+4\); R\(_4\) equal in length to R\(_2+3+4\); four medial veins present, petiola half longer than M\(_1\) and equal in length to upper edge of d-cell, d-cell trapezoidal, relatively large, constitutes ca. 1/6 length of wing; m-cu lies at 2/3 length of base of d-cell, far before bifurcation of M\(_3+4\) into M\(_3\) and M\(_4\); vein A\(_2\) slightly wavy.

**Remarks** The wing venation resembles that of *C. jurassica* but significantly differs from that species in the structure of the d-cell and the proportions of the medial veins.

*Cretolimonia mikolajczyki* Kopeć, Krzemiński, Soszyńska-Maj sp. nov.
(Figs. 7-8)

**Etymology** We dedicate the species to the memory of a great and regrettably late colleague, the well-known Polish dipterologist Dr Waldemar Mikolajczyk.

**Material examined** Holotype No. MP/4082, male (Fig. 7A); Kachin amber (Northern Myanmar); mid-Creta-ceous, earliest Cenomanian. Housed in the Institute of Systematics and Evolution of Animals, Polish Academy of Sciences (ISEA PAS).

**Diagnosis** Wing broad, about 2.3 times as long as its width; wing venation differs in proportions of individual veins from all species in this genus; vein Sc ends opposite the fork of Rs, d-cell trapezoidal and distinctly expanded at the tip, constituting 1/10 of the wing length; petiola constitutes only about 1/3 of M\(_1\); cross-vein m-cu just before the fork M\(_3+4\) on M\(_3\) and M\(_4\); gonocoxites long, narrow, with numerous bristles, outer gonostylus long, narrow, strongly hooked at the tip; inner gonostylus delicate, lobed, elongated at the tip.

**Description** Well-preserved specimen. Wing length 2.6 mm; width 1.2 mm; body length 2.8 mm. *Head*. Considerably wider than its length; antennae 16 segmented, scapus large, tubular half as long as its width, pedicel round, almost half as wide as scapus; flagellum 14 segmented, the first basal segment expanded in the lower part, the last small and rounded; bristles are present on all the flagellomeres, on the three basal segments bristles 2.5-3 times as long as the width of the segment, on the other flagellomeres bristles are shorter, 1.5-2 times as long as the width of the segment, on the last segment 3 or 4 short bristles (Fig. 7B, 8A). Palpi invisible. *Thorax*. Wing about 2.3 times as long as its width (Fig. 7C, 8B), additionally expanded in the anal field, vein Sc ends opposite bifurcation of Rs; cross-vein sc-r is about its length opposite the end of Sc; Sc ends opposite bifurcation of R\(_2+3+4\) into R\(_1\) and R\(_3\); R\(_2\) fully atrophied; Rs about 1/4 longer than R\(_2+3+4\); R\(_3\) short, slanting, less than 1/3 length of R\(_2+3+4\); R\(_4\) only slightly shorter than R\(_2+3+4\); four medial veins present, and vein M\(_1\) 2.5 times longer than petiola and twice as long as upper edge of d-cell; d-cell trapezoidal, small, constituting only 1/10 of wing length; cross-vein m-cu at end of d-cell, just before M\(_3+4\) bifurcates into M\(_3\) and M\(_4\); A\(_2\) vein strongly waved. Legs long, delicate, tibial spurs absent. *Abdomen*. Hypopygium (Fig. 7D, 8C): gonocoxites long, narrow, with numerous bristles, at end bearing a bunch of long bristles; outer gonostylus strongly chitinized, long, narrow, strongly hooked at end; inner gonostylus delicate, lobed, elongated at the end; penis long, narrow, slightly curved; parameres shorter than penis, broad at base, similar to parameres in *Cretolimonia dayana* sp. nov..

**Remarks** *Cretolimonia mikolajczyki* sp. nov. is the first representative of the genus, and of the Architipulinae, found in fossil resin. Its excellent preservation permits detailed examination of morphological characters, especially the structure of the antennae and copulatory apparatus.
Representatives of the infraorder Tipulomorpha were already present among the oldest fossil Diptera specimens from the early Middle Triassic (Anisian), ca. 245 Ma, from Arzviller (Vosges Mts., France)\(^3\),\(^5\),\(^15\),\(^27\),\(^28\). *Archilimonia vogesiana* Krzemiński and Krzemińska, 2003\(^3\) and *A. krzeminski* Lukashevich and Ribeiro, 2019\(^16\) belong to the fossil family Archilimoniidae. Archilimoniidae was included as a subfamily within the family Limoniidae, which we consider erroneous since the wing venation more closely resembles that of Pediciidae, especially in

**Figure 7.** *Cretolimonia mikolajczyki* sp. nov., holotype No. MP/4082: (A) habitus of male; (B) antennae; (C) wing; (D) male hypopygium.

**Figure 8.** *Cretolimonia mikolajczyki* sp. nov. Holotype No. MP/4082: (A) antenna; (B) wing venation; (C) male hypopygium. Abbreviation: *aed* aedeagus, *gx* gonocoxite, *ing* inner gonostylus, *oug* outer gonostylus, *par* parameres, *ped* pedicel, *scp* scapus.
the radial field. This being the case, the oldest known representative of the Limoniidae would be *Architipula youngi* Krzeminski, 1992 from the Late Triassic of North America (ca. 220 Ma) used for age calibration of Tipulomorpha. It should be noted that the first 100 million years of dipteran, and hence tipulomorph, evolution is documented only by impression fossils, mainly wings. The earliest examples of Limoniidae revealing three-dimensional structure have been recovered from Lower Cretaceous Lebanese amber, but such inclusions are very few in number.

The specimens described here are of paramount importance for understanding the morphology of Architipulinae (Fig. 7A), and thus shed new light on the early evolution of flies of the suborder Tipulomorpha. The male genitalia of these flies are usually so severely deformed during fossilization that their precise structure and spatial arrangement cannot be determined. Although the preservation of *Cretolimonia dayana* sp. nov. is very good, being an impression fossil, it is only possible to obtain information in two dimensions. Therefore, finding a male of the same genus (*C. mikolajczyki* sp. nov.) in mid-Cretaceous amber presented a unique opportunity to fully reconstruct the genital anatomy, and thus to verify the structure of the hypopygium of *C. dayana* sp. nov. (Fig. 3B,4D). In sedimentary rocks, the genitalia of females are much better preserved, as they are usually strongly chitinized; in favourable circumstances even spermathecae are visible, and sometimes also genital plates. However, so far, the number of spermathecae has not been known for the oldest known Limoniidae, the subfamilies Architipulinae or Eotipulinae. The well-preserved material from the Daya site allowed us to reconstruct the genitalia of the female *C. dayana* sp. nov. and to determine the number of spermathecae. The spermathecae are identical to those of most species in the family Limoniidae.

Most genera of Limoniidae have a full set of five radial veins, including the R2 vein, which takes the form of a cross-vein r-r. However, as far back as the Early Jurassic, a number of genera had appeared in which the cross-vein r-r (R2) had disappeared (i.e., atrophied) and one of these is the genus *Cretolimonia*. The disappearance of this cross-vein is observed also in some other Limoniidae, for example, in the subfamily Chioneinae, in the modern genera *Gonomyia* Meigen, 1818 and *Rhabdomastix* Skuse, 1890. There are, however, significant differences in other sectors of the wing in these genera; only three medial veins are always present, and the cross-vein m-cu is located in the anterior half of the d-cell, usually near the bifurcation of M3+4 into M3 and M4.

New material described here enabled us to characterize the morphology of the oldest (at least in the geological sense), extinct group of the Limoniidae; these features will be of great importance when introduced to the phylogenetical analysis of this large dipteran family.

**Materials and methods**

**Geological context.** This study was based on impression fossils and an amber inclusion (Fig. 9). The rock material comes from two sites, Shevia and Daya, in the Shelopuginsky District of the Chita region of Transbaikalia (Russia). Shevia (Dain Formation), dated Early Cretaceous, is located on the right bank of the Shevia River, 3 km below Shevia village, 2 km above the confluence of the Shevia and Shiviinsky Bumulei Rivers. The Daya site is located on the left bank of the Daya River above the Shevia Valley. It exposes sediments of the Glushkovo Formation, which are imprecisely dated but likely to lie close to the Jurassic/Cretaceous boundary.

The investigated amber inclusion derives from a former amber mine located near Danai (Tanai) Town (approximately at 26° 150’ N, 96° 340’ E) in the Hukawng Valley, state of Kachin in northern Myanmar. Radiometric U-Pb zircon dating of the volcanioclastic matrix of the amber produced an age of 98.79 ± 0.62 million years (earliest Cenomanian).

**Specimen repository.** All the specimens studied in the course of this work are deposited permanently in publicly owned collections in national museums. The Myanmar amber inclusion, the holotype of *C. mikolajczyki* sp. nov., is deposited in the National Museum of Myanmar in Yangon.
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Author contributions
K.K. took the lead in writing the manuscript. K.K., W.K. were responsible for taxonomic decisions. K.K. was responsible for material preparation and photography. I.K.K. and A.S.M. performed graphic illustration. K.K., W.K., A.S.M. contributed to the interpretation of the results, correction of manuscript after reviews and general discussion. W.K. was the leader who supervised the work and financially supported the project. R.A.C. revised the text. A.S.M. was the corresponding author and coordinated the correction of the manuscript. All authors provided critical feedback and helped shape the research and manuscript.

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Competing interests
The authors declare no competing interests.

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