Shards, sequences, and shorelines: two new species of Bembidion from North America (Coleoptera, Carabidae)

David R. Maddison

Department of Integrative Biology, Oregon State University, Corvallis, OR 97331, USA

Corresponding author: David R. Maddison (david.maddison@oregonstate.edu)

Abstract

Two new species of Bembidion are described from river shores in North America. One, Bembidion mimbres sp. nov., from the Gila River watershed in the lands of the Mimbres culture in New Mexico and Arizona, is closely related to the widespread Bembidion levigatum. DNA sequences from several linkage groups and morphology provide evidence of the distinctiveness of B. mimbres. The second, Bembidion corgenoma sp. nov., has been the subject of recent genomic and transcriptomic studies. It belongs in the Bembidion transversale subgroup, and occurs from California north to British Columbia, east to Montana and Nevada. The B. transversale subgroup as a whole is reviewed, and morphological characters that distinguish B. corgenoma from the similar and sympatric B. transversale and B. erosum are described and illustrated. DNA sequences of these three species show no consistent differences in 28S, COI, CAD, and Topoisomerase, and a coalescent species delimitation analysis reveals no notable structure within the complex. This is the first known trio of species within Bembidion for which those genes provide no clear signal of species boundaries. A neotype is designated for the one name in the group that lacks a primary type, Bembidium haplogonum Chaudoir. Chromosomes of the new species and their relatives are as is typical for Bembidion, with eleven pairs of autosomes and an XY/XX sex chromosome system.

Keywords

Bembidiini, ground beetle, molecular systematics, species delimitation, taxonomy, Trechinae

Copyright David R. Maddison. This is an open access article distributed under the terms of the Creative Commons Attribution License (CC BY 4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.
Introduction

*Bembidion* is a very large genus of small beetles with more than 1,200 species worldwide (Lorenz 2005). Most species of these small predators live along the edges of bodies of water and can be abundant in their habitats.

In the course of an ongoing project revising the bembidiine carabids of America north of Mexico, a number of undescribed species have been discovered. Most of these will be described in due course within complete revisions of subgenera or species groups. However, two of these new species are or will be soon discussed in the scientific literature, and warrant description more quickly, in order to provide them with names. These two are also especially significant, as they have cultural connections to humans, implicit or explicit, of very different sorts.

One of them is a member of the subgenus *Hydrium*, a group of relatively large *Bembidion* that is widespread in the Northern Hemisphere. The new species (Fig. 1) is only known from the Gila River watershed of southeastern Arizona and southwestern New Mexico, where it lives along the banks of rivers and creeks, on the ground a few meters away from the shoreline (Fig. 2), most commonly under willows (*Salix*). The distribution of this new species is within that of the Mimbres culture, which flourished in that area one thousand years ago. This culture is perhaps best known for black-on-white Mimbres pottery, the designs of which depicted people, cultural icons, and organisms (Hegmon et al. 2018). The people of the Mimbres culture were deeply aware of the arthropods in their environment, as indicated by the astonishing array of images on their pottery of insects, including among others geometrid larvae (https://core.tdar.org/image/383483/1452-style-iii-bowl-from-cameron-creek; https://doi.org/10.6067/XCV8Z60P2N), Orthoptera (https://core.tdar.org/image/383111/2685-style-iii-bowl-from-swarts; https://doi.org/10.6067/XCV80Z7364), dragonflies, and ant lions (Hegmon, et al. 2018). In honor of these peoples who were so connected to the small organisms in their midst, this elegant beetle species is given the name *Bembidion mimbres*.

The second species described here is connected to humans via modern biological research: it is becoming the first model species of *Bembidion* for genomic and transcriptomic studies. It is member of the *B. transversale* species group of the *Ocydromus* complex of *Bembidion*, containing some of the largest *Bembidion* in North America (Lindroth 1963; Maddison and Swanson 2010). The *B. transversale* group consists of two subgroups, the *B. transversale* subgroup and the *B. mexicanum* subgroup (Maddison 2012; Maddison and Swanson 2010). Maddison and Swanson (2010) considered the *B. transversale* subgroup to contain three species (*B. transversale* Dejean, *B. perspicuum* (LeConte), and *B. sarpedon* Casey), although they noted that “*B. transversale*” showed enough morphological variation to suggest that it may contain multiple species. Because of the especially complex pattern of variation, it only recently become clear that the genomic and transcriptomic model species was undescribed. The new species (Fig. 3) is common in Oregon and California, with some populations in neighboring regions, living along cobble and gravel shores of rivers and creeks (Fig. 4). It is the best sequenced *Bembidion* genomically and transcriptomically (Gustafson et al. 2019; Gustafson et al. 2020; Pflug...
Two new *Bembidion* species

Figure 1. Paratype male of *Bembidion mimbres* (voucher number V100327) from the type locality.

Figure 2. Habitat of *Bembidion mimbres* at USA: New Mexico: Grant Co., Gila River, Billings Vista, 1320 m, 32.8163°N, 108.6032°W (type locality). Arrow indicates approximate location of most specimens. Found in the same habitat were *Bembidion aratum* (LeConte), *B. impotens* Casey, *B. scintillans* Bates, *B. horni* Hayward, *B. rupicola* (Kirby), *B. clemens* Casey, *B. rapidum* (LeConte), and *Omophron oblitteratum* Horn.
et al. 2020), and has been used as one of the models for developing a UCE probe set for adephagan beetles (Gustafson, et al. 2019; as B. haplogonum Chaudoir). It is a centerpiece of ongoing and future studies of genome size in carabids (e.g., Pflug, et al. 2020). In this paper, it is described as Bembidion corgenoma.

An important step enabling future research about these beetles is providing the species with stable names. Although we now know the two new species in many ways unimaginable to those who lived a thousand years ago, including detailed aspects of their

Figure 3. Paratype male of Bembidion corgenoma (voucher number V101452) from the type locality. Scale bar: 1 mm.
Two new Bembidion species

Figure 4. Habitats of Bembidion corgenoma A USA: Oregon: Benton Co., Corvallis, Willamette River, 60 m, 44.5491°N, 123.2449°W (type locality). B. corgenoma and B. recticolle are both common in this habitat. B. corgenoma is more common near the water in areas with vegetation growing nearby among the gravel and cobbles B USA: California: Del Norte Co., Wilson Creek, 3 m, 41.6051°N, 124.1005°W, habitat of B. corgenoma and B. erosum, as well as Bembidion haruspex Casey, B. vandykei Blaisdell, B. curtulatum Casey, B. platynoides Hayward, and B incrematum LeConte.
DNA and genomes, we know very little about these two species in nature. With the decreasing emphasis on natural history in modern biology, it is possible that a person of the Mimbres culture knew aspects of the daily life of *B. mimbres* much better than we ever will. However, perhaps this paper, in giving names to the two species and presenting identification tools allowing them and their near relatives to be distinguished, will inspire research about these beetles, including into their way of life along river shores.

**Materials and methods**

Members of *Bembidion* were examined from or will be deposited in the collections listed below. Each collection’s listing begins with the code used in the text.

- **CAS** California Academy of Sciences, San Francisco, USA;
- **CNC** Canadian National Collection, Ottawa, Canada;
- **CSCA** California State Collection of Arthropods, Sacramento, USA;
- **EMEC** Essig Museum Entomology Collection, University of California, Berkeley, USA;
- **MCZ** Museum of Comparative Zoology, Harvard University, Cambridge, USA;
- **NHMUK** The Natural History Museum, London, UK;
- **MNHN** Muséum National d’Histoire Naturelle, Paris, France;
- **MSBA** Museum of Southwestern Biology, University of New Mexico, Albuquerque, USA;
- **OSAC** Oregon State Arthropod Collection, Oregon State University, Corvallis, USA;
- **UAIC** University of Arizona Insect Collection, Tucson, USA;
- **UASM** University of Alberta Strickland Museum, Edmonton, Canada;
- **UBCZ** Spencer Entomological Museum, University of British Columbia, Vancouver, Canada;
- **USNM** National Museum of Natural History, Smithsonian Institution, Washington, DC, USA;
- **ZMUM** Zoological Museum, Moscow State University, Moscow, Russia.

**Collecting methods**

Specimens were collected by hand or using an aspirator; specimens were found during the day in their habitat, or with the aid of a headlamp at night, when the beetles are more actively moving on the surface. Specimens for morphological studies were killed and preserved in maple (*Acer*) sawdust to which ethyl acetate was added. Specimens for DNA sequencing were collected into 95% or 100% ethanol. For chromosomal studies, live specimens were placed in simple Carnoy’s solution (three parts 100% ethanol : one part glacial acetic acid), and the abdomens were opened up shortly after death to allow better penetration of the fixative; the specimens were stored in Carnoy’s in a -20 °C freezer, with vials contained within multiple layers of plastic to prevent the release of acid fumes.
Two new Bembidion species

Morphological methods

General methods of specimen preparation for morphological work, and terms used, follow Maddison (1993; 2008). Genitalia were prepared, after dissection from the body, by treatment in 10% KOH at 65 °C for 10 minutes followed by a series of multi-hour baths of distilled water, 5% glacial acetic acid, distilled water, and then ethanol. Male genitalia were then mounted in Euparal between two small coverslips attached to archival-quality heavyweight watercolor paper, and, once dried, pinned beneath the specimen. Male genitalia were examined for 30–60 specimens each of *B. transversale*, *B. erosum*, and *B. corgenoma*, and four or five specimens each of *B. levigatum* and *B. mimbres*.

Photographs of entire beetles and antennae were taken with a Leica M165C dissecting scope and a Sony NEX-7 camera, and of male genitalia with either a Leica Z6 Apo lens and DMC4500 camera or a Leica DM5500B compound microscope and DMC425C camera. A stack of images from different focal positions was then merged using the PMax procedure in Zerene Systems’s Zerene Stacker; the final images thus potentially have some artifacts caused by the merging algorithm. Measurements were made using Leica Application Suite v4.9 from images acquired using these imaging systems.

Cytogenetic methods

Twenty-two males were examined for chromosome number and sex-chromosome system. Methods used were as outlined by Maddison (1985; 2008). Males examined were: 1 *Bembidion mimbres* from USA: New Mexico: Grant Co., Billings Vista, Gila River, 1310 m, 32.8137°N, 108.6031°W; 2 *B. transversale* from USA: Colorado: Fremont Co., Arkansas River at Texas Creek 1880 m, 38.4100°N, 105.5854°W; 3 *B. erosum* from USA: California: Del Norte Co., Wilson Creek , 3 m, 41.6051°N, 124.1005°W; 3 *B. perspicuum* from USA: Arizona: Cochise Co., San Pedro R at Charleston, 31.6239°N, 110.1722°W; 3 *B. sarpedon* from USA: Colorado: Las Animas Co., Purgatoire R., 2.7 km W Cokedale 1910 m, 37.1295°N, 104.6398°W; 3 *B. mexicanum* from USA: Arizona: Pima Co., Santa Rita Mtns, Box Canyon, 1455 m 31.7981°N, 110.7767°W; 3 *B. lugubre* from USA: Arizona: Pima Co., Santa Rita Mtns, Box Canyon, 1455 m, 31.7981°N, 110.7767°W. In addition, the vouchers of the *B. transversale* group studied in Maddison (1985) were re-examined and re-identified.

Molecular methods

DNA extraction and sequencing

Genes studied, and abbreviations used in this paper, are: 28S: 28S ribosomal DNA (D1-D3 domains); 18S: 18S ribosomal DNA; COI: cytochrome c oxidase I; wg: wingless; CAD: carbamoyl phosphate synthetase domain of the rudimentary gene; ArgK: arginine kinase; Topo: topoisomerase I.
DNA was extracted using a Qiagen DNeasy Blood and Tissue Kit. Fragments for the seven genes were amplified using the polymerase chain reaction (PCR) on an Eppendorf Mastercycler ProS Thermal Cycler, using TaKaRa Ex Taq and the basic protocols recommended by the manufacturers. Primers and details of the cycling reactions used are given in Maddison (2012) and Maddison and Cooper (2014). The amplified products were then cleaned, quantified, and sequenced at the University of Arizona's Genomic and Technology Core Facility using a 3730 XL Applied Biosystems automatic sequencer. Assembly of multiple chromatograms for each gene fragment and initial base calls were made with Phred (Green and Ewing 2002) and Phrap (Green 1999) as orchestrated by Mesquite’s Chromaseq package (Maddison and Maddison 2018a, c) with subsequent modifications by Chromaseq and manual inspection. Multiple peaks at a single position in multiple reads were coded using IUPAC ambiguity codes.

**Taxon sampling for DNA studies**

For the phylogenetic study of *Bembidion* (*Hydrium*), 19 specimens of the subgenus *Hydrium*, as well as five species serving as outgroups (Table 1) were used. Of the 152 sequences examined, 97 were newly acquired, with 55 being from previous publications (Maddison 2012; Maddison and Cooper 2014; Maddison et al. 2019; Maddison and Maruyama 2019; Maddison and Ober 2011). For the phylogenetic study of the *B. transversale* group, I sampled 56 specimens of the *B. transversale* subgroup, as well as three species of the *B. mexicanum* subgroup (Table 2). Of the 237 sequences examined, 109 were newly sequenced, with 128 from previous publications (Kanda et al. 2015; Maddison 2012; Maddison and Swanson 2010; Wild and Maddison 2008). Sequences of the two holotypes listed in Tables 1 and 2 are “genseq-1”, of paratypes “genseq-2”, and the remainder are all “genseq-4” (Chakrabarty et al. 2013). In addition, sequences of the newly designated neotype of *Bembidium haplogonum* Chaudoir were acquired and deposited in GenBank (accession numbers MW151478, MW151506, and MW151563), and are “genseq-1”. Localities of the sequenced specimens of *B. levigatum*, *B. mimbres*, *B. transversale*, *B. erosum*, and *B. corgenoma* are given in Table 3.

**Sequence alignment**

Alignment was not difficult for any of the protein-coding genes. There were no insertion or deletions (indels) evident in the sampled CAD, ArgK, Topo, *wg*, or COI sequences. An alignment of 28S was conducted in MAFFT version 7.130b (Katoh and Standley 2013), using the L-INS-i search option and otherwise default parameter values.

**Molecular phylogenetic analysis**

Maximum likelihood analysis was conducted for each gene individually using IQ-TREE version 1.6.7.1 (Nguyen et al. 2015), as orchestrated by Mesquite’s Zephyr package (Maddison and Maddison 2018b, c). The ModelFinder feature within IQ-TREE
Table 1. Sampling of members of *Bembidion* (*Hydrium*) and related subgenera for DNA-based study. Four-digit numbers under “#” are D.R. Maddison DNA voucher numbers; the specimen marked with an * is the holotype of *B. mimbres*. All other specimens listed of *B. mimbres* are paratypes. An abbreviation for state or province of capture is given under “Loc”; further information on *B. levigatum* and *B. mimbres* specimens is given in Table 3. The last eight columns contain GenBank accession numbers.

| #     | Loc | 28S     | COIBC | COIPJ | COI-like | wg       | CAD       | ArgK       | Topo       |
|-------|-----|---------|-------|-------|----------|----------|-----------|------------|------------|
|       |     |         |       |       |          |          |           |            |            |
| **Outgroups** |     |         |       |       |          |          |           |            |            |
| B. incisum Andrewes | 1131 | JN170365 | JN171052 | MW151374 |          | JN171434 | JN170833 | JN170588 | JN171232 |
| B. dyschirinum LeConte | 0896 | JN170332 | JN171029 | MW151370 |          | GU556029 | JN170799 | JN170553 | JN171210 |
| B. lampros (Herbst) | 1727 | JN170370 | MF616889 | MW151371 |          | JN171439 | JN170838 | JN170593 | MF616755 |
| B. propens (Stephens) | 1279 | JN170408 | JN171088 | MW151372 |          | JN171482 | JN170833 | JN170640 | JN171270 |
| B. luridicorne Solsky | 1122 | JN170377 | JN171064 | MW151373 |          | JN171447 | JN170847 | JN170602 | JN171245 |
| **Bembidion (Hydrium)** |     |         |       |       |          |          |           |            |            |
| B. interventor Lindroth | 1693 | VA | MW151391 | JN171059 | NW151374 | | | | |
| B. nitidum (Kirby) | 1255 | IN | MW151392 | MW151377 | | | | | |
| B. oliquium LeConte | 1299 | IA | MW151395 | KJ624355 | MW151376 | | | | |
| B. levigatum Say | 1694 | IA | MW151393 | MW151378 | MW151378 | | | | |
| B. mimbres sp. nov. | 0763 | NE | GU560883 | MW151379 | | | | | |
| B. mimbres | 2343 | TX | MW151395 | MW151381 | | | | | |
| B. mimbres | 1220 | NM | MW151396 | MW151382 | | | | | |
| B. mimbres | 1267 | NM | MW151405 | MW151382 | | | | | |
| B. mimbres | 1944 | NM | MW151403 | MW151389 | | | | | |
| B. mimbres | 2117 | NM | MW151397 | MW151383 | MW151383 | | | | |
| B. mimbres | 2118 | NM | MW151398 | MW151384 | MW151384 | | | | |
| B. mimbres | 2119 | NM | MW151399 | MW151385 | MW151385 | | | | |
| B. mimbres | 2131 | NM | MW151400 | MW151386 | MW151386 | | | | |
| B. mimbres | 2134 | NM | MW151401 | MW151387 | MW151387 | | | | |
| B. mimbres | 2135 | NM | MW151402 | MW151388 | MW151388 | | | | |
Table 2. Sampling of members of *Bembidion transversale* group for DNA-based study. Four-digit numbers under "#" are D.R. Maddison DNA voucher numbers. Under "T", the holotype of *B. corgenoma* is indicated by "H", and paratypes by "P". An abbreviation for state or province of capture is given under "Loc"; further information on specimens of *B. transversale*, *B. erosum*, and *B. corgenoma* is given in Table 3. The last four columns contain GenBank accession numbers.

| #  | T  | Loc | 28S        | COI        | CAD        | Topo |
|----|----|-----|------------|------------|------------|------|
| 1712 | T | AZ  | JN170375   | JN171062   | JN170845   | JN171243 |
| 2192 | T | NM  | GU454739   | GU454769   | JN170854   | JN171250 |
| 2483 | T | CO  | JN170403   | JN171082   | JN170875   | JN171263 |
| 2484 | T | CO  | JN170432   | JN171106   | JN170908   | JN171285 |
| 2514 | T | CO  | KU233761   | KU233815   | KU233926   | KU234052 |
| 3009 | T | CO  | KU233764   | KU233818   | KU233929   | KU234055 |
| 3761 | T | UT  | KU233771   | KU233823   | KU233933   | KU234059 |
| 3776 | T | UT  | KU233775   | KU233827   | KU233937   | KU234063 |
| 1120 | T | AZ  | GU454740   | GU454770   | JN170877   | JN171265 |
| 2318 | T | CA  | GU454743   | GU454782   | KU233923   | KU234049 |
| 3774 | T | CA  | KU233773   | KU233825   | KU233935   | KU234061 |
| 3775 | T | CA  | KU233774   | KU233826   | KU233936   | KU234062 |
| 2485 | T | CO  | GU454748   | GU454778   | KU233924   | KU234050 |
| 2182 | T | NM  | GU454749   | KU233810   | KU233922   | KU234047 |
| 2596 | T | CA  | MW151550   | MW151493   | MW151522   | MW151465 |
| 2607 | T | CA  | MW151551   | MW151494   | MW151523   | MW151466 |
| 3561 | T | CA  | MW151552   | MW151495   | MW151524   | MW151467 |
| 3562 | T | CA  | MW151553   | MW151496   | MW151525   | MW151468 |
| 3584 | T | CA  | MW151554   | MW151497   | MW151526   | MW151469 |
| 4050 | T | CA  | MW151549   | MW151492   | MW151521   | MW151464 |
| 4212 | T | CA  | MW151555   | MW151498   | MW151527   | MW151470 |
| 4033 | T | OR  | MW151556   | MW151499   | MW151528   | MW151471 |
| 2162 | T | BC  | KU233749   | KU233803   | KU233915   | KU234040 |
| 2194 | T | BC  | KU233757   | KU233811   | MW151529   | MW151440 |
| 2160 | T | NS  | GU454762   | KU233802   | KU233914   | KU234048 |
| 2486 | T | CO  | GU454755   | GU454785   | KU233925   | KU234039 |
| 2157 | T | WY  | EU677688   | GU454797   | EU677541   | KU234051 |
| 4690 | T | MT  | MW151561   | MW151504   | MW151534   | MW151476 |
| 4927 | T | MT  | MW151562   | MW151505   | MW151535   | MW151477 |
| 5064 | T | WA  | MW151560   | MW151503   | MW151533   | MW151475 |
| 5613 | T | OR  | MW151557   | MW151500   | MW151530   | MW151472 |
| 4219 | T | OR  | MW151558   | MW151501   | MW151531   | MW151473 |
| 5612 | T | OR  | MW151559   | MW151502   | MW151532   | MW151474 |
| 4052 | T | CA  | KU233782   | KU233831   | KU233941   | KU234067 |
| 2181 | T | CA  | KU233755   | KU233809   | KU233921   | KU234046 |
| 4054 | T | CA  | KU233783   | KU233832   | KU233942   | KU234068 |
| 2180 | T | CA  | KU233754   | KU233808   | KU233920   | KU234045 |
| 3772 | T | P   | KU233772   | KU233824   | KU233934   | KU234060 |
| 4961 | T | P   | MW151536   | MW151479   | MW151508   | MW151451 |
| 4962 | T | P   | MW151537   | MW151480   | MW151509   | MW151452 |
| 4218 | T | CA  | MW151544   | MW151487   | MW151516   | MW151459 |
| 2597 | T | CA  | MW151538   | MW151481   | MW151510   | MW151453 |
| 2608 | T | CA  | MW151539   | MW151482   | MW151511   | MW151454 |
| 3559 | T | CA  | KU233769   | KU233821   | KU233931   | KU234057 |
| 3560 | T | CA  | KU233770   | KU233822   | KU233932   | KU234058 |
| 3583 | T | CA  | MW151540   | MW151483   | MW151512   | MW151455 |
| 4959 | T | NV  | MW151541   | MW151484   | MW151513   | MW151456 |
| 2346 | T | NV  | GU454763   | GU454793   | MW151507   | MW151450 |
| 5670 | T | OR  | MW151545   | MW151488   | MW151517   | MW151460 |
Two new *Bembidion* species

Table 3. Localities of capture of *Bembidion* specimens of *B. levigatum*, *B. mimbres*, and the *B. transversale* subgroup whose DNA was sequenced. Four-digit numbers at the start of each row are D.R. Maddison DNA voucher numbers.
| Location | Coordinates |
|----------|-------------|
| USA: Washington: Whatcom Co., Nooksack River 1.4 mi S of Deming, 70 m, 48.808°N, 122.2019°W |
| USA: California: Marin Co., Nicasio Reservoir, 70 m, 38.088°N, 122.7383°W |
| USA: California: Del Norte Co. rt 101 @ Wilson Creek, 41.60530°N, 124.10060°W |
| USA: California: Del Norte Co. rt 101 @ Wilson Creek, 41.60530°N, 124.10060°W |
| USA: California: Del Norte Co., Wilson Creek, 3 m, 41.6051°N, 124.1005°W |
| USA: California: Tehama Co., Red Bluff, Sacramento River, 73 m, 40.1759°N, 122.229°W |
| USA: Oregon: Coos Co., Crooked Creek S of Bandon, 7 m, 43.0814°N, 124.4251°W |
| USA: Oregon: Curry Co., Floras Creek at route 124 SE Langlois, 21 m, 42.9132°N, 124.4251°W |
| USA: California: Monterey Co., Big Sur River, Andrew Molera State Park, 7 m, 36.285°N, 121.8544°W |
| USA: California: San Luis Obispo Co., Pismo State Beach, 4 m, 35.0999°N, 120.6267°W, 29.iv.2014 |
| USA: California: San Luis Obispo Co., San Simeon State Park, San Simeon Creek, 4 m, 35.5955°N, 121.1233°W |
| USA: Nevada: Carson City Co., Carson River at Silver Saddle Ranch, SE Carson City, 1405 m 39.1315°N, 122.229°W |
| USA: California: Tehama Co., Red Bluff, Sacramento River, 73 m, 40.1759°N, 122.229°W |
| USA: Oregon: Linn Co., Willamette River, Truax Island, 60 m 44.5853°N, 123.1913°W |
| USA: Oregon: Corvallis, Willamette River, 60 m 44.5491°N, 123.2451°W |

(Kalyaanamoorthy et al. 2017) was used to find the optimal character evolution models. The MFP model option was used for 28S, and the TESTMERGE option for the protein-coding genes. The TESTMERGE option sought the optimal partition of sites, beginning with the codon positions in different parts. Twenty searches were conducted for the maximum-likelihood tree for each matrix.

For the *B. transversale* group, a multi-species coalescent approach was conducted with the 28S, COI, CAD, and Topo data to provide an algorithmic analysis of species boundaries. STACEY version 1.2.5 (Jones 2017) was used as implemented in BEAST version 2.6.2 (Bouckaert et al. 2014), with the epsilon value set to 1*10–4, CollapseWeight parameters to 0.5 and 10, and with a Beta prior. I evaluated sampling sufficiency using ESS values in Tracer version 1.7.1 (Rambaut et al. 2018); after four independent runs of 1E9 generations each, all ESS values exceeded 200, except for mutationRate.s:Topo, whose ESS value was 191. As I saved trees every 100,000 generations, with the first 10% of the trees discarded as the burn-in period, this yielded a sample of 72,000 trees.

**Data resources**

Sequences have been deposited in GenBank with accession numbers MW151366–MW151563. Aligned data for each specimen as well as files containing inferred trees for each gene are available in Supplementary material S1 and S2, and have been deposited in the Dryad Digital Repository, [https://doi.org/10.5061/dryad.18931zcw1](https://doi.org/10.5061/dryad.18931zcw1).
Two new Bembidion species

Results

Molecular results

In the analysis of DNA data for the subgenus *Hydrium*, *B. levigatum* and *B. mimbres* sp. nov. differed in all genes except COI (Fig. 5), providing evidence that they are two separate species. In all gene trees except 28S, *B. nitidum* was the sister group of *B. levigatum* + *B. mimbres* (Suppl. materials 1).

The majority of specimens of *B. levigatum* and *B. mimbres* were indistinguishable in COI, but there were four sequences of *B. mimbres* that formed a separate clade (for...
specimens 1220, 1267, as well as the second sequences of 1944 and 2117). These four sequences have 29 sites at which they differ from all other sampled *B. levigatum* + *B. mimbres*, at 20 of which these four sequences have the same base as in at least one other sampled *Hydrium* species. These four sequences might be nuclear copies or numts (Thalmann et al. 2004), or they could represent the effects of *Wolbachia* infections (Smith et al. 2012). Although it is possible these are the true mitochondrial copies of COI, and that the other sequences are numts, the evidence points to the four unusual sequences being something other than true mitochondrial copies: the chromatograms for these four unusual sequences have several double peaks, indicating polymorphism within the PCR products for non-synonymous differences. These four sequences have been deposited in GenBank as “COI-like” sequences.

In each of the four genes studied in the *B. transversale* group, the maximum likelihood tree showed a monophyletic *B. transversale* subgroup (Suppl. materials 2), with *B. perspicuum* and *B. transversale* s. l. (= *B. transversale* + *B. erosum* + *B. corgenoma*) forming a clade, the sister of which is *B. sarpedon*. None of the three species within *B. transversale* s. l. form a clade in any of the four genes studied (Fig. 6). The multispecies coalescent STACEY tree also showed no distinction between these species based upon the combined analysis of 28S+COI+CAD+Topo (Fig. 7).

Cytogenetic results

All males examined are inferred to have 22 autosomes (i.e., 11 pairs of autosomes) and an XY/XX sex chromosome system (Table 4).

Re-examination of voucher specimens identified as *B. transversale* in Maddison (1985) showed that they belong to three species. The specimens from Alberta and near Fernie, BC, are *B. transversale*; the specimen from near Cache Creek, BC, is *B. corgenoma*; the specimen from Salmon Valley, BC, is *B. erosum*. The specimens called “*B. sp nr transversale-Nr 1*” are *B. pernotum*, and the specimens called “*B. sp nr transversale-Nr 2*” are *B. lugubre*. The specimens reported as belonging to “*B. sp.nr. transversale*” in Pflug et al. (2020) are all *B. corgenoma*. These new identifications are incorporated into the summary shown in Table 4.

Morphological results

Morphological results for *Bembidion* (*Hydrium*) are presented in the taxonomic section below.

Members of the *B. transversale* subgroup are very similar morphologically. DNA sequence data of 28S, COI, CAD, and Topo do not reveal any consistent phylogenetic structure within *B. transversale* s. l. (Figs 6, 7), suggesting that it is perhaps a single species. In all other bembidiines investigated to date (e.g., Maddison 2008; Maddison and Cooper 2014; Maddison and Sproul 2020; Sproul and Maddison 2017), every form judged by morphological evidence as a distinct species is revealed as a clade in the tree of at least one of these four genes. I detected no variation in chromosomes within
Two new Bembidion species

Table 4. Chromosome numbers and sex chromosomes of Bembidion (Hydrium) and Bembidion transversale group males. The Sample column indicates the total number of specimens examined in this paper and in previous papers. “1” in Reference indicates Maddison (1985); “2” indicates Pflug et al. (2020).

| Species         | 2n male | Sample | Locality | Reference |
|-----------------|---------|--------|----------|-----------|
| B. levigatum    | 22+XY   | 2♂     | TX       | 1         |
| B. mimbres      | 22+XY   | 1♂     | NM       | this paper|
| B. transversale | 22+XY   | 5♂, 2♀ | AB, BC, CO | 1, this paper|
| B. erosum       | 22+XY   | 4♂     | BC, OR   | 1, this paper|
| B. corgenoma    | 22+XY   | 15♂    | OR, BC   | 1, 2      |
| B. peripiculum  | 22+XY   | 4♂     | CO, AZ   | 1, this paper|
| B. urpedon      | 22+XY   | 3♂     | CO       | this paper|
| B. perspicuum   | 22+XY   | 6♂     | CO       | 1, this paper|
| B. mexicanum    | 22+XY   | 5♂     | CO, AZ   | 1, this paper|
| B. lugubre      | 22+XY   | 11♂    | AZ, CA, Mexico | 1, this paper|

the group (Table 4); although Bembidion species typically have similar chromosomes (Maddison 1985), some subgenera have species that differ by chromosome number (Maddison 2008). However, in spite of the lack of genetic evidence supporting multiple species, the morphological results reveal that this complex consists of at least three distinct but very similar species.

Examination of primary types (documented in the Taxonomic Treatment section, below) indicates that two of the species have names (Bembidion transversale and B. erosum), and the third is described here as B. corgenoma; these names will be used in advance of the Taxonomic Treatment to simplify the text.

The morphological evidence indicating that there are three species includes color (Figs 11A–C, 12), mentum shape (Fig. 13), and male genitalic structure (Figs 14–17), with the patterns summarized below and graphically in Fig. 18.

The dark and pale western species (B. erosum and B. corgenoma) are broadly sympatric from southern California through British Columbia (Figs 19, 20), and are found microsympatrically (on the same gravel bank) at Wilson Creek, Del Norte County, California (41.6051°N, 124.1005°W), as well as along Floras Creek, Curry County, Oregon (42.9132°N, 124.4251°W), and the Siletz River E of Kernville, Oregon (44.8720°N, 123.9223°W). They differ consistently in size of a sclerotized lobe of the internal sac (Fig. 15B vs. C); the thickness of the tip of the flagellar sheath, with B. corgenoma having a somewhat triangular sclerotized region of the tip (arrow in Fig. 16E), as opposed to a thin dark line in B. erosum (Fig. 16C, D); and color, with B. erosum being generally darker (Fig. 11B) than B. corgenoma (Fig. 11C), especially the appendages (Fig. 12B vs C). In addition, most males of B. erosum have the ventral surface of the apex of the aedeagus more strongly curved downward (Fig. 14C, D). Although there are no universal distinctions between the two species in the genes sequenced, the six B. erosum and five B. corgenoma sequenced from the gravel bank shown in Fig. 4B consistently differ in one base in Topoisomerase, suggesting, combined with differences in genitalia and color, that there is no or extremely limited gene flow at that locality.

The ranges of the two generally paler species (B. transversale and B. corgenoma) overlap in Nevada, Washington, Idaho, and Montana (Fig. 19), and there are three localities
Table 1. Maximum likelihood gene trees of the Bembidion transversale subgroup.

| Species                     | COI             | 28S             |
|-----------------------------|-----------------|-----------------|
| B. transversale s. lat.     |                 |                 |
| B. corgenoma                |                 |                 |
| B. erosum                   |                 |                 |

Figure 6. Maximum likelihood gene trees of the Bembidion transversale subgroup. B. mexicanum subgroup and B. sarpedon were present in the analysis, and reconstructed outside this clade, but were graphically removed to simplify this figure. Holotype of B. corgenoma indicated by a star. Scale bar as reconstructed by IQ-TREE.
Two new *Bembidion* species

Figure 7. Majority rule consensus tree of trees found from a STACEY analysis. Numbers on branches are estimates of the Bayesian posterior probability of a clade, expressed as a percentage.
at which they co-occur (16 mi W Lolo Pass, Ravalli Co., Montana; Walla Walla, Washington; Spokane, Washington; all in OSAC). There are subtle but consistent differences in the male genitalia, as well as striking differences in the mentum, and I am convinced any gene exchange in the region of overlap is minimal. In the overlap region there are a very few specimens of B. transversale with paler antennae (similar to typical B. corgenoma), and there is one population at Hayden Lake, Idaho (CAS), which contains B. transversale typical in all regards except for one specimen that has the anterior margin of the mentum somewhat intermediate between the two species (Fig. 13B). With these minor exceptions, the differences in mentum, genitalia, and color are consistent throughout the overlap range among the many males whose genitalia were dissected.

**Bembidion erosum** and **B. transversale** are the two most similar species within the trio, differing most notably in the anterior lateral region of the mentum: in B. erosum this region is large and triangular, similar to that standard in **Bembidion** (i.e., like those shown in Fig. 13A), in contrast to the modified mentum of B. transversale (Fig. 13B–D), in which the anterior lateral region is much reduced. The male genitalia are very similar, both having a larger lobe on the basal sclerite (Fig. 15A, B), and a thin, non-triangular apex to the flagellar sheath (Fig. 16A–D). However, the flagellar sheath is more dorso-ventrally compressed in B. transversale (Fig. 16A, B), and the flagellar complex is thinner (arrow in Fig. 17A, B). With one exception, the known ranges of B. erosum and B. transversale do not overlap, with B. erosum in the United States being restricted to the Cascades and west, and B. transversale only known from east of the Cascades (Figs 19, 20). The one exception is in eastern Washington: there is a series of seven specimens of B. erosum labeled “Colbert, Wash / V 30 1937 / Dan Bonnell”. The two males from this series have typical B. erosum genitalia, and the mentum has the anterior lateral region large and triangular. Colbert is less than 20 km NNE of Spokane, where both B. transversale and B. corgenoma have been collected. As the only record of B. erosum east of the Cascades, some doubt is cast upon its authenticity, but there is no reason otherwise to question the locality data.

In addition, there is a form in the Sierra Nevada of California that requires further examination. My limited study of it indicates that it has all of the morphological features of B. transversale, except that it has a normal mentum shape, with large and triangular anterior lateral regions.

**Taxonomic Treatment**

**Subgenus Hydrium**

The subgenus **Hydrium**, as defined by Maddison (2012), contains seven species in the Palaearctic region (Marggi et al. 2017) as well as five species in the Nearctic region (Lindroth 1963): **Bembidion nitidum**, **B. interventor**, **B. obliquulum**, **B. levigatum**, and the new species described here.

The species key in Lindroth (1963) can be modified as follows to take into account the new species.
Two new Bembidion species

19 More than two setae on the clypeus, and at least one long seta on the front angle of the prothorax. Elytra without the typical pair of distinguishable dorsal punctures but most intervals with a row of small punctures, each carrying a long seta ................................................................. 19A

– Clypeus with only two setae. Elytra with two dorsal punctures on third interval, otherwise without setigerous punctures ........................................ 20

19A Elytra with a row of long setae on all intervals; prothorax wide, sides very rounded (Fig. 8A) .............................................................................................................. B. levigatum

– Elytra with a row of long setae on most intervals, but lacking on intervals 2 and 4. Prothorax narrower, with straighter sides (Fig. 8B) ........ B. mimbres

Bembidion mimbres sp. nov.
http://zoobank.org/5F95B4F6-9FFD-4841-8E69-794A258381A2

Holotype. Male, in OSAC, labeled: “USA: New Mexico: Grant Co., Gila River, Billings Vista, 1320 m, 32.8163°N, 108.6032°W, 11.viii.2005. DRM 05.043. D.R. & J.H.A. Maddison”, “David R. Maddison DNA2131 DNA Voucher” [pale green paper], “HOLOTYPE Bembidion mimbres David R. Maddison” [partly handwritten, on red paper], “Oregon State Arthropod Collection OSAC_0002000007 [matrix code]” [printed on both sides of white paper]. Genitalia in glycerin vial pinned beneath specimen; extracted DNA stored separately. GenBank accession numbers for DNA sequences of the holotype are MW151386, MW151400, MW151414, MW151425, MW151432, and MW151443.

Paratypes (116). USA: New Mexico: Grant Co., Gila River, Billings Vista, 1320 m, 32.8163°N, 108.6032°W [Type locality] (44: OSAC, USNM, MCZ, NHMUK, MNHM, MSBA); USA: New Mexico: Grant Co., Billings Vista, Gila River, 1310 m, 32.8137°N, 108.6031°W (28: OSAC, CAS, UAIC, EMEC); USA: New Mexico: Grant Co., Gila River near Cliff, 1350 m, 32.9124°N, 108.5897°W (12: OSAC); USA: New Mexico: Grant Co., Gila River near Gila, 1370 m, 32.9692°N, 108.5868°W (3: OSAC); USA: New Mexico: Grant Co., Gila River near Gila, 1370 m, 32.969°N, 108.587°W (4: OSAC); USA: New Mexico: Grant Co., Gila River, Gila National Forest, 1315 m, 32.8167°N, 108.6035°W (14: OSAC); USA: New Mexico: Gila R., jct US 260, nr. Gila (1: UASM); USA: Arizona: Navajo Co., Carrizo Ck nr. Carrizo (10: UASM).

Type locality. USA: New Mexico: Grant Co., Gila River, Billings Vista, 32.8163°N, 108.6032°W.

Derivation of specific epithet. Bembidion mimbres is named in honor of the people of the Mimbres culture, who lived alongside this species, including at the type locality, and who illustrated the insects in their world on their pottery (Hegmon, et al. 2018). The name is to be treated as a noun in apposition.

Diagnosis and description. Adults of this species are relatively large Bembidion (5.3–6.3 mm in length), with a striking appearance because of the smooth and shiny dorsal surface with its metallic reflections (Figs 1, 8B). Body piceous, with an aeneous,
green, or blue metallic reflection. Legs with tarsi and tibiae testaceous, femora infuscated. First three antennomeres testaceous, with the tip of the third infuscated in some specimens; fourth basally testaceous. Palps testaceous except for the penultimate maxillary article, which is infuscated. Mentum with anterior lateral regions large and triangular as typical for a *Bembidion*; mentum tooth incised at tip, and thus bifid (similar to *B. levigatum*). Prothorax with relatively straight sides, especially posteriorly (Fig. 8B); with distinct posterolateral carina close to the lateral margin; posterior region of pronotum smooth, impunctate. Lateral bead of elytra extending inside shoulder well toward the midline, sharply angulate. Striae consisting of rows of distinct punctures, without an associated groove; on stria 1 complete; striae 2–6 absent in the posterior third; stria 7 virtually absent, represented by at most minute punctulae. Dorsal surface lacking microsculpture, and thus very shiny. More than two setae on the clypeus, and extra setae on the frons, at least anteriorly; at least one long seta near the front angle of the prothorax; prosternum with at most two setae. Elytra without the typical pair of distinguishable setose punctures associated with interval 3, but with a row of long setae on all intervals except 2 and 4. Aedeagus (Fig. 9C,D) with ventral margin thinner, and with internal sac sclerites very similar to those of *B. levigatum*, but with slight differences, especially basally.
Figure 9. Male genitalia of subgenus *Hydrium* A. *B. levigatum* (voucher DNA1693, Virginia: Danville City Co., Danville) B. *B. levigatum* (voucher DNA2343, Texas: Bastrop Co., Colorado River near Utley) C. *B. mimbres* (voucher DNA2134, USA: New Mexico: Grant Co., Gila River, Billings Vista) D. *B. mimbres* (voucher DNA2135, USA: New Mexico: Grant Co., Gila River, Billings Vista). Scale bar: 0.1 mm.
Most easily distinguished from B. levigatum by the narrower prothorax with straighter sides (Fig. 8B), the lack of setae on elytral intervals 2 and 4, and having at most one or two setae on the prosternum.

**Additional characteristics.** Diploid chromosome number 24, with 11 pairs of autosomes and an XY/XX sex chromosome system.

**Geographic distribution.** Known from the Gila River watershed in Arizona and New Mexico (Fig. 10).

**Habitat.** At the type locality, found at night on damp sandy soil about 2–4 meters from the river shore, in the shade of a large willow (Fig. 2). At a site a few meters away, 28 specimens were found at night on damp clay/sand soil among small *Salix* and *Populus* saplings 2–8 meters from the water’s edge; in spite of extensive searching in the same area, only one specimen was found during the day. At other sites along the Gila River, found in habitats similar to those in which B. levigatum is found, on damp sand and silt on the steep upper bank of the river, mostly at night.

*Bembidion levigatum* Say

*Bembidium levigatum* Say, 1823: 84. Lectotype female in MNHN, designated by Linderoth and Freitag (1969). Type locality Missouri.

*Bembidion laevigatum delawarens* Casey, 1924: 24. Holotype male in USNM (type number 36814), examined. Type locality Pennsylvania.
Diagnosis and geographic distribution. Adults of this species are large and distinctively wide, with a broad, rounded prothorax (Fig. 8A), and with a very shiny dorsal surface, with a green or bluish metallic reflection. The clypeus, frons, anterior corner of the prothorax, and all elytral intervals have long setae in addition to the standard set in Bembidion; the prosternum has four or more setae. Aedeagus as in Fig. 9A, B. A very widespread species, found throughout most of the eastern United States and a small region of southern Canada, from Maine to Florida, north and west to Alberta and Montana, south to Utah, New Mexico, Texas, and Mexico (Bousquet 2012). I have also seen specimens from the Grand Canyon in northern Arizona (two specimens in MSBA labeled “USA AZ Coconino Co Grand Canyon Nat. Park, N36.77 W111.655 RMBL 29-30 August 2002 coll. Cobb, Brantley, Lightfoot”).

Bembidion transversale species group

The Bembidion transversale group contains large Bembidion found primarily on river shores of cobble, gravel, and sand from Canada to Guatemala. Members of the group are characterized by large size (5.8–8 mm); posterolateral carina of pronotum lacking or indistinct and somewhat oblique; lateral bead of elytra not prolonged onto shoulder; crista clavicularis absent; elytral striae distinct and mostly complete; elytral microsculpture transverse; two discal setae of elytra in or close to third stria. It belongs to what has been called the Nearctic Ocydromus Clade (Maddison 2012), although that group is not closely related to subgenus Ocydromus. The only subgeneric name available for the Nearctic Ocydromus Clade is Leuchydrium Casey, although the type species (Bembidion tigrinum LeConte) is quite distant from the B. transversale group (Maddison 2012).

There are now eight recognized species in the B. transversale group in the United States and Canada:

Bembidion transversale subgroup

Bembidion transversale Dejean, 1831  
Bembidion erosum (Motschulsky, 1850)  
Bembidion corgenoma Maddison, sp. nov.  
Bembidion perspicuum (LeConte, 1848)  
Bembidion sarpedon Casey, 1918  

Bembidion mexicanum subgroup  

Bembidion mexicanum Dejean, 1831  
Bembidion lugubre LeConte, 1857  
Bembidion pernotum Casey, 1918

There is a total of 20 species-group names that have been applied to members of the B. transversale group (for details beyond those provided below, see Maddison and Swanson (2010)). I have examined detailed photographs of the primary type of one (the holotype of Bembidium transversale Dejean, in the MNHN), and the
primary types themselves of 18. The twentieth lacked a type series, and a neotype is designated below.

The species key in Lindroth (1963) can be modified as follows to take into account species in the *B. transversale* subgroup. Specimens from this group are not easy to identify.

145  Prothorax (Lindroth 1963: figs 168a–b) without or with very faint, oblique latero-basal carina ................................................................. 145A

–  Prothorax with latero-basal carina well developed, less oblique...........146

145A Mentum with anterior lateral region reduced, not triangular, each consisting of a mesal denticle and a more lateral rounded bump (Fig. 13B–D). Antenna with at least the second and third antennomeres apically infuscated. Tip of aedeagus not abruptly curved downward (Fig. 14A, B); basal sclerotized lobe large (Fig. 15A); apex of flagellar sheath with long, thin dark line (Fig. 16A, B)............................................................................ *B. transversale*

–  Mentum with anterior lateral region as typical for a *Bembidion*: triangular, large, and with anterior margin significantly anteriad of the central tooth (Fig. 13A). Other characteristics either as mentioned above or not ......145B

145B Paler, with antennae gradually becoming slightly darker toward the apex; pronotum in most specimens dark rufous. Prothorax with lateral margins more rounded, very shiny, with weaker microsculpture and less punctuation. Relatively flat elytral intervals with small punctures in striae. Aedeagus with ventral margin having a slight downward bulge. Internal sac sclerite complex of male genitalia narrower in side view, with relatively long and thin flagellar complex. Known from NM, CO, WY, UT, AZ.........................*B. sarpedon*

–  Darker, with at least antennomeres 4–11 infuscated; pronotum rufous or piceous. Prothorax with lateral margins less rounded, less shiny, and in most specimens with more punctures. Aedeagus without ventral bulge. Internal sac sclerite complex less narrow, with a dorso-ventrally wider flagellar complex........145C

145C Prothorax with later margins less sinuate, with more notable punctures in the basal region (Maddison and Swanson 2010: fig. 4B); aedeagus with ostial flag more dorsal, and with a more abrupt curve at its anterior end (Maddison and Swanson 2010: fig. 6B).....................................................*B. perspicuum*

–  Prothorax with lateral margins more sinuate, flatter, with a smoother basal region (Maddison and Swanson 2010: fig. 4A); aedeagus with ostial flag extending further ventrally, and with gentler curvature (Maddison and Swanson 2010: fig. 6A)..................................................................................145D

145D Legs and basal three antennomeres pale, testaceous or rufo-testaceous. Elytral striae deeper. Tip of aedeagus not abruptly curved downward (Fig. 14E, F); basal sclerotized lobe small (Fig. 15C); apex of flagellar sheath with dark area triangular (Fig. 16E, F)......................................................................*B. corgenoma*

–  Legs in most specimens darker (in southern specimens infuscated); second and third antennomeres infuscated, at least apically. Elytral striae shallower.
Tip of aedeagus abruptly curved downward (Fig. 14C, D); basal sclerotized lobe large (Fig. 15B); apex of flagellar sheath with long, thin dark line (Fig. 16C, D).............................................................................................................**B. erosum**

**Bembidion transversale** Dejean, 1831

*Bembidium transversale* Dejean, 1831:110. Holotype female, in MNHN, examined by Kipling Will, who provided photographs that confirmed the identification. Type locality restricted to Nipigon, Ontario, by Lindroth (1963).

**Diagnosis and geographic distribution.** Adults of this species (Fig. 11A) are characterized by the reduced anterior lateral regions of the mentum (Fig. 13B–D), and the antenna with at least the second and third antennomeres apically infuscated (Fig. 12A). The prothorax is more cordate than in other species. The aedeagus has its ventral surface relatively straight, with the tip not abruptly curved downward (Fig. 14A, B); the basal sclerotized lobe is large (Fig. 15A), and the apex of the flagellar sheath has a long, thin dark line (Fig. 16A, B). This is the easternmost species, found from Newfoundland and Nova Scotia west through Ontario to southeastern British Columbia, central Oregon, northeastern Nevada, northern Utah, and Colorado (western portion of distribution shown in Fig. 19).

**Bembidion erosum** (Motschulsky, 1850)

*Peryphus erosus* Motschulsky, 1850:10. Lectotype female, in ZMUM, examined, designated by Bousquet and Larochelle (1993), labeled “type” [handwritten], “California” [handwritten on red paper], “Peryphus erosus Mots California” [handwritten on green paper], [a rectangle of blank red paper], “LECTOTYPE Peryphus erosus Motschulsky Des. by Y. Bousquet ’91” [partly handwritten on red paper]. Type locality California.

*Bembidion marinicum* Casey, 1918:57. Holotype female in USNM (type number 36919), examined. Type locality Marin County, California.

**Diagnosis and geographic distribution.** Most adults of this species are the darkest members of this group (Fig. 11B), with the second and third antennomeres infuscated (Fig. 12B), at least apically, and with dark femora, although the more northern populations (e.g., from mainland British Columbia) have paler legs and paler ground color of the body. Prothorax moderately cordate, with a smooth basal region with few punctures. Tip of aedeagus abruptly curved downward (Fig. 14C, D); basal sclerotized lobe large (Fig. 15B); apex of flagellar sheath with long, thin dark line (Fig. 16C, D). This species is coastal, occurring from Haida Gwaii in British Columbia south along the coast to Big Sur and neighboring areas of California, with only one record from east of the Cascades (Fig. 20).
Figure 11. Adult males of *Bembidion transversale* subgroup members **A** *B. transversale* (voucher V101454, Canada: Alberta: Burbank, junction of Red Deer and Blindman Rivers, 52.3542°N, 113.7556°W) **B** *B. erosum* (voucher V101453, USA: California: Del Norte Co., Wilson Creek, 3 m, 41.6051°N, 124.1005°W). **C** *B. corgenoma* (voucher V101452, from type locality) **D** *B. perspicuum*, light form (voucher V101461, USA: Arizona: Cochise Co., San Pedro R at Charleston, 31.6239°N, 110.1722°W) **E** *B. perspicuum*, dark form (neotype of *Bembidion haplogonum* Chaudoir, USA: California: Lake Co., North Branch Cache Creek at hwy 20, 305 m 38.9881°N, 122.5400°W) **F** *B. sarpedon* (voucher V101459, USA: Colorado: Las Animas Co., Cokedale, Reilly Canyon, 37.1346°N, 104.6114°W). Scale bar 1.0 mm.

*Bembidion corgenoma* sp. nov.
http://zoobank.org/BF5E001D-F543-4149-8081-0BF7B99A8484

**Holotype.** Male, in OSAC, labeled: “USA: Oregon: Benton Co., Corvallis, Willamette River, 60 m, 44.5491°N, 123.2449°W, 7.x.2019. DRM 19.210. D.R. Maddison”, “David R. Maddison DNA5673 DNA Voucher” [pale green paper], “HOLOTYPE"
Two new Bembidion species

Figure 12. Antennae of *B. transversale* subgroup  
A *B. transversale* (voucher V101457, Canada: Ontario: Thunder Bay Dist., Rossport)  
B *B. erosum* (voucher V101456, USA: California: Del Norte Co., Wilson Creek, 3 m, 41.6051°N, 124.1005°W)  
C *B. corgenoma* (voucher V101455, USA: Oregon: Coos Co., Crooked Creek S of Bandon, 43.0814°N, 124.4335°W). Scale bar 0.1 mm.

Bembidion corgenoma David R. Maddison” [partly handwritten, on red paper], “Oregon State Arthropod Collection OSAC_0002000008 [matrix code]” [printed on both sides of white paper]. Genitalia mounted in Euparal in between coverslips pinned with specimen; extracted DNA stored separately. GenBank accession numbers for DNA sequences of the holotype are MW151449, MW151463, MW151491, MW151520, and MW151548.

Paratypes (193). USA: Oregon: Benton Co., Corvallis, Willamette River, 44.5491°N, 123.2449°W, 60 m [type locality] (78: OSAC, CNC, CAS, UAIC, UASM, MCZ, EMEC, CSCA); USA: Oregon: Benton Co., Corvallis, Willamette River, 60 m, 44.5491°N, 123.2451°W (10: OSAC); USA: Oregon: Benton Co., Corvallis, Willamette River, 44.5475°N, 123.2428°W, 60 m (35: OSAC, USNM,
NHMUK, MNHM, UBCZ); USA: Oregon: Benton Co., Corvallis, Willamette River, 60 m, 44.5478°N, 123.2430°W (2: OSAC); USA: Oregon: Benton Co., Corvallis, 62 m, 44.5491°N, 123.2449°W (6: OSAC); USA: Oregon: Linn Co., Willamette River, Truax Island, 44.5853°N, 123.1913°W, 60 m (12: OSAC); USA: Oregon: Lane Co., Goodman Creek, Willamette NF, 43.8441°N, 122.6736°W, 290 m (2: OSAC); USA: Oregon: Coos Co., Crooked Creek S of Bandon, 43.0814°N, 124.4335°W, 7 m, 24.iii.2014 (26: OSAC); USA: California: Tehama Co., Red Bluff, Sacramento River, 40.1759°N, 122.229°W, 73 m (22: OSAC, CAS, EMEC).

**Type locality.** USA: Oregon: Benton Co., Corvallis, Willamette River, 44.5491°N, 123.2449°W.

**Derivation of specific epithet.** The specific epithet is formed from the Latin word *cor*, meaning heart, and *genoma*, a modification (for easier pronunciation) of the coined word “genome”. *Corgenoma* refers to this species being the heart or current focus of genomic studies in small carabid beetles. *Cor* also alludes to the type locality of Corvallis, Oregon, whose name is derived from Latin, and means “heart of the valley”. It is to be treated as a noun in apposition.

**Diagnosis and description.** Length (5.8–7.4). Relatively light in color compared to *B. transversale* and *B. erosum*, with legs and basal three antennomeres pale, testaceous or rufo-testaceous. Head and prothorax piceous, with metallic reflections, on pronotum green or aeneous, on head bluish or green. Elytra paler, with shoulders and most of the anterior half testaceous with an orange tint, bordered posteriad by a dark band (with intervals 1–3 in this region dark rufous), followed by a pale testaceous band that either extends to the apex or that is bounded posteriad by dark lateral spots which in the darkest individuals merge in the middle. Mentum with anterior lateral region as typical for a *Bembidion*: triangular, large, and with anterior margin significantly anteriad of the central tooth (Fig. 13A); central tooth trapezoidal, rounded. Prothorax cordate, with more sinuate margins than *B. perspicuum*, with a relatively smooth basal region, with minute punctures; pronotum without or with very faint, oblique posterolateral carina as in other members of this group. Elytral striae 1–5 complete; stria 6 distinct and strong through much of its length; stria 7 shallower, less impressed than 6, but distinct. Microsculpture of elytra very transverse, with little tendency to form meshes. Two discal setae on each elytron, close to third stria. Tip of aedeagus not abruptly curved downward (Fig. 14E, F); basal sclerotized lobe small (Fig. 15C); apex of flagellar sheath with dark area triangular (Fig. 16E, F).

**Additional characteristics.** Diploid chromosome number 24, with 11 pairs of autosomes and an XY/XX sex chromosome system (Pflug, et al. 2020). Genome size (1C), as measured by flow cytometry, 2118 Mb in males and 2193 Mb in females (Pflug, et al. 2020). Most specimens with singleton (non-conjugated) sperm (Gómez and Maddison 2020).

**Available genomic and transcriptomic data.** Transcriptomic data for one specimen is available on NCBI’s Sequence Read Archive at accession SRR8801541, and genomic data of four specimens at accessions SRR8518612, SRR8518625, SRR8518626, and SRR8518631 (Pflug, et al. 2020).
Two new *Bembidion* species

Notes. This species was called *Bembidion haplogonum* Chaudoir in Gustafson et al. (2019; 2020), and *B. sp. nr. transversale* in some other publications (Gómez and Madison 2020; Kanda, et al. 2015; Pflug, et al. 2020).

Geographic distribution. This species occurs from central British Columbia south through the Willamette Valley of Oregon, the Central Valley of California, with some records in Nevada, Idaho, eastern Washington, and Montana (Fig. 19), thus overlapping with the range of *B. transversale*.

Geographic variation. The specimens on or close to the beaches of the Pacific Ocean appear on average slightly paler than more inland specimens.
Figure 14. Male genitalia of *B. transversale* subgroup A *B. transversale* (voucher DNA4219, USA: Oregon: Harney County, Banks of Silver Creek, 1379 m, 43.7278°N, 119.6256°W) B *B. transversale* (voucher DNA2161, Canada: Alberta: Lethbridge, Oldman River, 800 m, 49.7043°N, 112.866°W) C *B. erosum* (voucher DNA4033, USA: Oregon: Curry Co., Floras Creek at route 124 SE Langlois, 21 m, 42.9132°N, 124.4251°W) D *B. erosum* (voucher DNA3562, USA: California: Del Norte Co., Wilson Creek, 3 m, 41.6051°N, 124.1005°W) E *B. corgenoma* (voucher DNA2180, USA: California: Sonoma Co., Russian River, 3 mi NE Healdsburg) F *B. corgenoma* (holotype, voucher DNA5673, USA: Oregon: Benton Co., Corvallis, Willamette River, 60 m, 44.5491°N, 123.2449°W). Scale bar: 0.1 mm.

Figure 15. Basal sclerotized lobe of internal sac of male *Bembidion transversale* group members A *B. transversale* (voucher DNA4219, USA: Oregon: Harney County, Banks of Silver Creek, 1379 m, 43.7278°N, 119.6256°W) B *B. erosum* (voucher DNA4033, USA: Oregon: Curry Co., Floras Creek at route 124 SE Langlois, 21 m, 42.9132°N, 124.4251°W) C *B. corgenoma* (voucher DNA2180, USA: California: Sonoma Co., Russian River, 3 mi NE Healdsburg). Scale bar: 0.1 mm.
Two new *Bembidion* species

**Figure 16.** Central sclerite complex of *B. transversale* subgroup **A, B** *B. transversale* (vouchers V101437 and V101436, USA: Michigan: Port aux Pins, Bois Blanc Isl.) **C, D** *B. erosum* (vouchers V101440 and V101439, USA: California: Del Norte Co., Wilson Creek, 3 m, 41.6051°N, 124.1005°W) **E, F** *B. corgenoma* (vouchers V101428 and V101430, USA: Oregon: Benton Co., Corvallis, Willamette River, 44.5475°N, 123.2428°W). Scale bar: 0.1 mm.

**Habitat.** This species occurs on gravel or cobble shores of the rivers and creeks (Fig. 4), more often where the bank is relatively flat and has small amount of clay and silt mixed in with sand and gravel under the rocks. They also can be common under cobbles on the shores of small creeks on the upper portions of beaches of the Pacific Ocean.

*Bembidion perspicuum* (LeConte, 1848)

*Ochthedromus perspicuus* LeConte, 1848: 466. Holotype male, in MCZ (type number 5510), external structure and aedeagus examined. Type locality “Rocky Mountains”.
Ochthedomus mannerheimii LeConte, 1852:190. Lectotype female, designated by Maddison and Swanson (2010), in MCZ (type number 35571). Type locality San Diego, California. Although a male in the same unit tray as the lectotype is, by genitalic characters, a member of Bembidion corgenoma, I am uncertain about the lectotype. The almost complete absence of the seventh stria and the pronotal shape suggests Bembidion perspicuum, but the base of the pronotum is not as punctured as typical for that species. I tentatively place it as a synonym of Bembidion perspicuum. However, as Bembidion mannerheimii LeConte, 1852, is a junior homonym of Bembidion mannerheimii Sahlberg, 1827, this name is unavailable in any event.

Bembidium haplogonum Chaudoir, 1868: 241. Neotype male, in MNHN, here designated, labeled “USA: California: Lake Co., North Branch Cache Creek at hwy 20, 305 m 38.9881°N, 122.54°W, 5.viii.2010. DRM 10.090. K.W. Will & D.R. Maddison”, “David R. Maddison DNA5681 DNA Voucher” [pale green paper], “NEOTYPE Bembidium haplogonum Chaudoir designated D.R. Maddison” [partly handwritten, on white paper, bordered by red lines]. Genitalia mounted in Euparal in between coverslips pinned with specimen; extracted DNA stored separately. GenBank accession numbers for DNA sequences of the neotype are MW151478, MW151506, and MW151563. Details about the choice of neotype are provided below.

Bembidion acomanum Casey, 1918: 59. Lectotype female, designated by Lindroth (1975), in USNM (type number 36916), examined. Type locality Jemez Springs, New Mexico (Lindroth 1975).

Bembidion excursum Casey, 1918: 59. Holotype female, in USNM (type number 36915), examined. Type locality Tucson, Arizona.

Bembidion tuolumne Casey, 1924:30. Lectotype male, designated by Lindroth (1975), in USNM (type number 36917), external structure and aedeagus examined. In Maddison and Swanson (2010), this was treated as tentatively a synonym of B. transversale. Further examination of the lectotype, including of its genitalia, reveal that this is a specimen of B. perspicuum. Type locality Tuolumne, California.

Designation of a neotype for Bembidium haplogonum Chaudoir. Lindroth (1963: 341) could not find the original type series for Bembidium haplogonum Chaudoir in the MNHN. Thierry Deuve and David Kavanaugh have both searched for it, and could not find it in the Chaudoir collection, although other specimens were found that had been collected in California by Pierre Joseph Michel Lorquin. As the original type series is presumed lost, I here designate a neotype.

Chaudoir’s (1868) description of Bembidium haplogonum is detailed enough to make determination of the species he was describing clear. In his description, the large size (8 mm) and absence of a carina near the hind angle of the prothorax could only apply (within California, the type locality) to a member of the B. transversale group. The color pattern of the elytra (“sur les élytres, une petite tache au milieu du bord antérieur de chacune, et une bande transversale un peu arquée aux trois quarts, d’un jaune testacé pâle, peu distinctes, surtout la tache basale”, which translates to “on the elytra, a small spot on the front
Two new *Bembidion* species

Figure 17. Flagella of *B. transversale* subgroup **A, B** *B. transversale* (vouchers V101438 and V101435, USA: Michigan: Port aux Pins, Bois Blanc Isl.) **C, D** *B. erosum* (vouchers V101442 and V101441, USA: California: Del Norte Co., Wilson Creek, 3 m, 41.6051°N, 124.1005°W) **E, F** *B. corgenoma* (vouchers V101434 and V101431, USA: Oregon: Benton Co., Corvallis, Willamette River, 44.5475°N, 123.2428°W). Scale bar: 0.1 mm.

edge of each, and a slightly curved transverse band at three-quarters, of a pale testaceous yellow, indistinct, especially the basal spot”) can only apply to some specimens of *B. erosum*, *B. lugubre*, or the northern, dark form of *B. perspicuum* (Fig. 11E), as the other species in California have the entire basal third to half of the elytra pale. The description of the appendages as having the first 3.5 antennomeres, palps, and legs all pale yellow
eliminates *B. erosum*, as the palps, femur, and antennomeres 2–11 are dark in California specimens of that species. I have seen no specimens of *B. lugubre* with an isolated small spot on the front edge of each elytron; the only specimens that have a pale area in the basal half of the elytra have the entire sides and basal regions of the elytra a pale rufous.

**Figure 18.** Summary of morphological differences between three species in the *B. transversale* subgroup in A mentum B antennal color C curvature of ventral margin of the aedeagus D basal sclerotizes lobe size E apex of flagellar sheath, F flagellum.
Two new *Bembidion* species

with a darker disc; this paler form of *B. lugubre* occurs north of Los Angeles in California. The metallic coloration of the pronotum ("*Le dessus d’un vert brillant un peu cuivreux*") is characteristic of *B. perspicuum*, but not *B. lugubre*; the latter has no metallic sheen in

---

**Figure 19.** Geographic distribution of *B. transversale* and *B. corgenoma* (eastern portion of distribution of *B. transversale* not shown).
Figure 20. Geographic distribution of *B. erosum*. 
any specimens I have seen from California, and in only very few individuals elsewhere. The flatter prothorax with less rounded sides and a wider lateral gutter is also distinctive of *B. perspicuum* relative to all three other species from California (*B. erosum*, *B. lugubre*, and *B. corgenoma*), as is the distinctly punctured pronotal base (“distinctement ponctué tout le long de la base”). The large size (8 mm) is more characteristic of *B. perspicuum*; I have seen no specimens of *B. corgenoma* longer than 7.5 mm, but have seen specimens of *B. perspicuum* that are 7.9 mm in length, and specimens of *B. perspicuum* are, in general, larger than those of *B. corgenoma*. Finally, the virtual absence of the seventh elytral stria is characteristic of *B. perspicuum* relative to *B. corgenoma*. As Lorquin travelled extensively in areas where the dark form of *B. perspicuum* occurs (Grinnell 1904), it is certainly reasonable that a specimen of that form could have been seen by Chaudoir.

My early interpretations of Chaudoir’s descriptions were in error, and led me to believe that Chaudoir’s specimen was a member of what I here call *Bembidion corgenoma*; that mistake led me to call the species studied in Gustafson et al. (2019) *Bembidion haplogonum*. As a correct reading of the original description shows that *Bembidion haplogonum* refers to the dark form of *B. perspicuum*, I have designated a specimen from northern California with a color pattern matching Chaudoir’s description as the neotype (shown in Fig. 11E).

**Diagnosis and geographic distribution.** Adults of this species are large, and have a pronotum that is flatter than in other members of the group, with less rounded sides, and with the basal region more evidently punctate (Maddison and Swanson 2010: fig. 4). At least antennomeres 4–11 infuscated. Specimens from most areas are relatively pale (Fig. 11D), with the front half of the elytra pale, but specimens from northern California and Oregon are much darker (Fig. 11E), with only elytral apices being pale. Aedeagal characters are described in Maddison and Swanson (2010). This species is known from Texas, Kansas, Colorado, New Mexico, Arizona, Utah, Nevada, California, and Oregon.

**Bembidion sarpedon** Casey, 1918

*Bembidion sarpedon* Casey, 1918: 58. Lectotype male, designated by Lindroth (1975), in USNM (type number 36914); external structure and aedeagus examined. Type locality Cañon City, Colorado.

*Bembidion animatum* Casey, 1918: 62. Lectotype female, designated by Lindroth (1975), in USNM (type number 36918), examined. Type locality Jemez Springs, New Mexico (Lindroth 1975).

**Diagnosis and geographic distribution.** Adults of this species (Fig. 11F) are the palest members of this group, with legs entirely testaceous or rufo-testaceous, with antennae gradually becoming slightly darker toward the apex, and pronotum in most specimens dark rufous as opposed to the piceous or black of other species. The dorsal surface is shinier than in other species, especially the pronotum, because of the nearly effaced microsculpture. The prothorax is moderately cordate; the elytral intervals are
flatter than in related species, with small punctures in the striae. The ventral margin of the aedeagus has a slight downward bulge, and the internal sac sclerite complex of male genitalia is narrow in lateral view, with a long and thin flagellar complex. Known from New Mexico and Colorado west to Arizona and Utah, and north to Mammoth Hot Springs, Wyoming (OSAC).

Concluding remarks

The pathways that led to the recognition of the two species described here were very different. When I encountered *Bembidion mimbres* for the first time, as pinned specimens at the University of Alberta’s Strickland Museum in 1981–1982, I immediately recognized them as an undescribed species. They shared the large size, setose elytra, shiny surface, and striking color of the distinctive *Bembidion levigatum*, but did not share *B. levigatum*’s unusual prothorax shape and width.

In contrast, it took at least 12 years of study for me to become confident that *B. corgenoma* was a new species, and that the *B. transversale* subgroup consisted of at least five species (*B. sarpedon, B. perspicuum, B. transversale, B. erosum, and B. corgenoma*). The distinctiveness of *B. sarpedon* and *B. perspicuum* was recognized many years ago. The specimens that remained (*B. transversale s. l.*), however, were so complex in their variation patterns, so lacking in a differentiating signal in DNA sequences, and with such similar genitalia, that at times I thought there was just one species in *Bembidion transversale s. l.*, and at other times more.

I had become so accustomed to the clarity provided by DNA sequences in my other taxonomic projects on bembidiines that I had become somewhat skeptical of the value of traditional taxonomic methods utilizing only patterns of morphological variation. Two events changed my mind, as they caused the patterns to become evident at last. The first event was Kip Will’s collecting of both dark and light specimens from the shores of Wilson Creek in north coastal California. They were so obviously different in color that I expected them to have clearly different genitalia, and different DNA sequences. My cursory inspection revealed only the slightest difference in the overall shape of the aedeagus (I had not yet noticed the differences in the structures of the internal sac), not notable enough to be significant in itself. In addition, all six sequenced dark specimens from that gravel bar differed from all five sequenced light specimens in one base in Topoisomerase, but they did not differ in 28S, COI, and CAD. The correlation between color, aedeagal shape, and that single base in Topoisomerase convinced me that there were likely two species at that site in northern California, although if so they would be much more similar than are most other pairs of closely related, sympatric *Bembidion* species. Examination of Motschulsky’s specimens eventually revealed that the dark species had a name, *Bembidion erosum*, but the pale species at Wilson Creek and elsewhere continued to trouble me: other than the normal mentum, I saw no consistent differences from the more eastern *Bembidion transversale*. The distinctiveness of the pale western form (here called *B. corgenoma*) did not become evident until
Two new *Bembidion* species

The basic morphological work was done: thorough examination of the genitalia of 63 *B. corgenoma* males and 33 *B. transversale* males, focused on the area of geographic overlap, revealed the consistent differences shown in Fig. 18, especially the shape of the basal sclerotized lobe (Fig. 18D) and the tip of the flagellar sheath (Fig. 18E). This confirmed that sequences of four genes will not necessarily reveal the presence or absence of gene flow, and that even in *Bembidion*, a group in which DNA sequences often work very well for species delimitation, species boundaries are sometimes more quickly uncovered by traditional morphological methods.

The lack of observed differentiation in DNA sequences between *Bembidion transversale*, *B. erosum*, and *B. corgenoma* suggests that these are young, recently differentiated species. The contrast is striking between this trio and other bembidiines; in most bembidiines, sequences in at least one of the handful of standard genes provides a clear signal of lack of gene flow between species (e.g., Maddison 2008; Maddison and Cooper 2014; Maddison and Sproul 2020; Sproul and Maddison 2017). Why is the signal of species boundaries so clear in most bembidiine groups, but not *B. transversale* s. l.? Given genomic resources now available for this group, one fruitful and available avenue of future research would be comparison of coalescent patterns of thousands of regions of the genome within both this trio of *Bembidion transversale* group species and other groups of bembidiines with similar levels of morphological divergence.

### Acknowledgements

I am very thankful to the many people who helped out with this project. I am grateful to Will G. Russell, Michelle Hegmon, and Janet C. Berlo for engaging in a discussion about the naming of *Bembidion mimbres*, and helping me become more aware of respectful ways to honor the people of the Mimbres culture and their descendants. I am especially grateful to Stewart Koyiyumptewa, Tribal Historic Preservation Officer of the Hopi Tribe, for giving me permission to give the name *Bembidion mimbres* to the species that lives in the Gila River watershed. I thank Stephen H. Lekson for providing information about the distribution of the Mimbres culture, and its presence around the type locality of *B. mimbres*. I also thank Karen A. Ober for her early help curating specimens of subgenus *Hydrium*.

For their help in looking for the original type series of *Bembidium haplogonum* Chaudoir, I thank Thierry Deuve and David Kavanaugh. For their aid in translating the original description of that name, I am very thankful to Thierry Deuve and Arnaud Faille, as they helped me correct my earlier unfortunate misinterpretation of the original French. For his examination of the primary type of *Bembidium transversale* Dejean, I am grateful to Kipling Will.

For permission and information that helped with my efforts to sample specimens from various State Parks in California, I thank Jay Harris, Vince Cicero, Ronnie Glick, Amy Palkovic, and Stephen Bachman of California State Parks. Thanks as well to Stephen Gaimari for his efforts in obtaining a California State Park permit for me. For
permission to collect *B. levigatum* from the Matheson Wetlands Preserve in Moab, Utah, and advice about the lands, I am grateful to Linda Whitham of the Nature Conservancy. For permission to collect at Wallowa State Park and Bandon Beach, I am thankful to Shawnae Stanton, Ben Fisher, Sara Griffith, and all of the other helpful people at the Oregon Parks and Recreation Department.

For help in collecting specimens, I thank Julia H. Amerongen Maddison, Pamela R. Tripplet, A.E. Arnold, Wayne P. Maddison, W. Moore, Kip W. Will, Janine N. Caira, John S. Sproul, J.C. Oliver, Kojun Kanda, James M. Pflug, A.J. Baker, M.D. Baker, W.E. Hall, J.K. Moulton, D.H. Kavanaugh, F.A.H. Sperling, C.J. Marshall, and M.L. Jameson.

I am thankful to all of the curators of collections from which I borrowed material, including Philip Perkins (MCZ), the late Terry Erwin (USNM), Nikolai Nikitsky (ZMUM), as well as Danny Shpeley and the late George E. Ball (UASM). I also thank Max Barclay and Beulah Garner for shipping the ZMUM specimens to me. I am especially grateful to David Kavanaugh for his finding, in the collection of the California Academy of Sciences, specimens that proved important for my understanding of the distribution of species of the *B. transversale* group.

For their performing many of the PCR reactions on which the DNA sequence data in this paper is based, I thank Danielle L. Mendez, Estany Campbell-Dunfee, Caitlin E. Hudecek, Lili S. Adams, Joseph J. Dubie, Chris M. Cohen, Kalyn M. Hansen, and Tiffany A. Soto.

Thanks as well to Wayne P. Maddison and Julia H. Amerongen Maddison for very helpful comments on the manuscript. I am also thankful to Kipling W. Will, Michael J. Raupach, Paolo Bonavita, and Dmitry Fedorenko for their valuable reviews.

This project was supported by National Science Foundation grant DEB-1258220, and the Harold E. and Leona M. Rice Endowment Fund at Oregon State University.

References

Bouckaert R, Heled J, Kühnert D, Vaughan T, Wu C-H, Xie D, Suchard MA, Rambaut A, Drummond AJ (2014) BEAST 2: A Software Platform for Bayesian Evolutionary Analysis. PLoS Computational Biology 10: e1003537. https://doi.org/10.1371/journal.pcbi.1003537

Bousquet Y (2012) Catalogue of Geadephaga (Coleoptera, Adepahga) of America, north of Mexico. ZooKeys 245: 1–1722. https://doi.org/10.3897/zookeys.245.3416

Bousquet Y, Larochelle A (1993) Catalogue of the Geadephaga (Coleoptera: Trachypachidae, Rhysodidae, Carabidae including Cicindelini) of America north of Mexico. Memoirs of the Entomological Society of Canada 167: 1–397. https://doi.org/10.4039/entm125167fv

Casey TL (1918) A review of the North American Bembidiinae. Memoirs on the Coleoptera 8: 1–223.

Casey TL (1924) Additions to the known Coleoptera of North America. Memoirs on the Coleoptera 11: 1–347. https://doi.org/10.5962/bhl.title.48776
Two new *Bembidion* species

Chakrabarty P, Warren M, Page LM, Baldwin CC (2013) GenSeq: An updated nomenclature and ranking for genetic sequences from type and non-type sources. ZooKeys 346: 29–41. https://doi.org/10.3897/zookeys.346.5753

Chaudoir M, de (1868) Observations synonymiques sur les Carabiques de l’Amérique septentrionale et descriptions d’espèces nouvelles de ce pays. Revue et Magazin de Zoologie, Series 2, 20: 239–245.

Dejean PFMA (1831) Spécies général des Coléoptères de la collection de M. le Comte Dejean. Paris, I–VIII, 384 pp.

Gómez RA, Maddison DR (2020) Novelty and emergent patterns in sperm: Morphological diversity and evolution of spermatozoa and sperm conjugation in ground beetles (Coleoptera: Carabidae). Journal of Morphology 281: 862–892. https://doi.org/10.1002/jmor.21144

Green P (1999) Phrap. Version 0.990329. http://phrap.org

Green P, Ewing B (2002) Phred. Version 0.020425c. http://phrap.org

Grinnell F (1904) An early collector in California. Entomological News 15: 202–204.

Gustafson GT, Alexander A, Sproul JS, Pfug JM, Maddison DR, Short AEZ (2019) Ultraconserved element (UCE) probe set design: Base genome and initial design parameters critical for optimization. Ecology and evolution 9: 6933–6948. https://doi.org/10.1002/ece3.5260

Gustafson GT, Baca SM, Alexander AM, Short AEZ (2020) Phylogenomic analysis of the beetle suborder Adephaga with comparison of tailored and generalized ultraconserved element probe performance. Systematic Entomology 45: 552–570. https://doi.org/10.1111/syen.12413

Hegmon M, McGrath JR, O’Hara III FM, Russell WG (2018) Mimbres pottery designs in their social contexts. In: Roth BJ, Gilman PA, Anyon R (Eds) New Perspectives on Mimbres Archaeology: Three Millenia of Human Occupation in the North America Southwest. The University of Arizona Press, Tucson, 149–168. https://doi.org/10.2307/j.ctv5cg8bb.10

Jones G (2017) Algorithmic improvements to species delimitation and phylogeny estimation under the multispecies coalescent. Journal of Mathematical Biology 74: 447–467. https://doi.org/10.1007/s00285-016-1034-0

Kalyaanamoorthy S, Minh BQ, Wong TKF, von Haeseler A, Jermiin LS (2017) ModelFinder: fast model selection for accurate phylogenetic estimates. Nature Methods 14: 587–589. https://doi.org/10.1038/nmeth.4285

Kanda K, Pfug JM, Sproul JS, Dasenko MA, Maddison DR (2015) Successful recovery of nuclear protein-coding genes from small insects in museums using Illumina sequencing. PLoS ONE 10: e0143929. https://doi.org/10.1371/journal.pone.0143929

Katoh K, Standley DM (2013) MAFFT Multiple Sequence Alignment Software Version 7: Improvements in Performance and Usability. Molecular Biology and Evolution 30: 772–780. https://doi.org/10.1093/molbev/mst010

LeConte JL (1848) A descriptive catalogue of the geodephagous Coleoptera inhabiting the United States east of the Rocky Mountains. Annals of the Lyceum of Natural History of New York 4: 173–233, 334–474. https://doi.org/10.1111/j.1749-6632.1848.tb00277.x

LeConte JL (1852) Descriptions of new species of Coleoptera, from California. Annals of the Lyceum of Natural History of New York 5: 185–216. https://doi.org/10.1111/j.1749-6632.1852.tb00123.x
Lindroth CH (1963) The ground-beetles (Carabidae, excl. Cicindelinae) of Canada and Alaska. Part 3. Opuscula Entomologica Supplementum XXIV: 201–408.

Lindroth CH (1975) Designation of holotypes and lectotypes among ground beetles (Coleoptera, Carabidae) described by Thomas L. Casey. The Coleopterists Bulletin 29: 109–147.

Lindroth CH, Freitag R (1969) North American ground-beetles (Coleoptera, Carabidae, excluding Cicindelinae) described by Thomas Say: designation of lectotypes and neotypes. Psyche 76: 326–361. https://doi.org/10.1155/1969/90590

Lorenz W (2005) Systematic list of extant ground beetles of the world (Insecta Coleoptera “Geadephaga”: Trachypachidae and Carabidae incl. Paussinae, Cicindelinae, Rhysodinae). Second edition. Published by the author [Hörmannstrasse 4, D-82327], Tützing, [i–iii,] 530 pp.

Maddison DR (1985) Chromosomal diversity and evolution in the ground beetle genus Bembidion and related taxa (Coleoptera: Carabidae: Trechitae). Genetica 66: 93–114. https://doi.org/10.1007/BF00139715

Maddison DR (1993) Systematics of the Holarctic beetle subgenus Bracteon and related Bembidion (Coleoptera: Carabidae). Bulletin of the Museum of Comparative Zoology 153: 143–299.

Maddison DR (2008) Systematics of the North American beetle subgenus Pseudoperyphus (Coleoptera: Carabidae: Bembidion) based upon morphological, chromosomal, and molecular data. Annals of Carnegie Museum 77: 147–193. https://doi.org/10.2992/0097-4463-77.1.147

Maddison DR (2012) Phylogeny of Bembidion and related ground beetles (Coleoptera: Carabidae: Trechinae: Bembidiini: Bembidiina). Molecular Phylogenetics and Evolution 63: 533–576. https://doi.org/10.1016/j.ympev.2012.01.015

Maddison DR, Cooper KW (2014) Species delimitation in the ground beetle subgenus Lio-cosmius (Coleoptera: Carabidae: Bembidion), including standard and next-generation sequencing of museum specimens. Zoological Journal of the Linnean Society 172: 741–770. https://doi.org/10.1111/zoj.12188

Maddison DR, Kanda K, Boyd OF, Faille A, Porch N, Erwin TL, Roig-Juñent S (2019) Phylogeny of the beetle supertribe Trechitae (Coleoptera: Carabidae): Unexpected clades, isolated lineages, and morphological convergence. Molecular Phylogenetics and Evolution 132: 151–176. https://doi.org/10.1016/j.ympev.2018.11.006

Maddison DR, Maddison WP (2018a) Chromaseq: a Mesquite package for analyzing sequence chromatograms. Version 1.31. http://chromaseq.mesquiteproject.org

Maddison DR, Maddison WP (2018b) Zephyr: a Mesquite package for interacting with external phylogeny inference programs. Version 3.0. http://zephyr.mesquiteproject.org

Maddison DR, Maruyama M (2019) Phylogenetic relationships and convergent evolution of ocean-shore ground beetles (Coleoptera: Carabidae: Trechitae: Bembidion and relatives). Systematic Entomology 44: 39–60. https://doi.org/10.1111/syen.12307

Maddison DR, Ober KA (2011) Phylogeny of minute carabid beetles and their relatives based upon DNA sequence data (Coleoptera, Carabidae, Trechitae). ZooKeys 147: 229–260. https://doi.org/10.3897/zookeys.147.1871
Maddison DR, Sproul JS (2020) Species delimitation, classical taxonomy and genome skimming: a review of the ground beetle genus *Lionepha* (Coleoptera: Carabidae). Zoological Journal of the Linnean Society 189: 1313–1358. https://doi.org/10.1093/zoolinnean/zlz167

Maddison DR, Swanson AP (2010) A preliminary characterization of *Bembidion perspicuum* LeConte, with a reclassification of related species (Coleoptera, Carabidae) north of México. ZooKeys 43: 15–31. https://doi.org/10.3897/zookeys.43.390

Maddison WP, Maddison DR (2018c) Mesquite: a modular system for evolutionary analysis. Version 3.51. http://mesquiteproject.org

Marggi W, Toledano L, Neri P (2017) Tribe Bembidiini Stephens, 1827, subtribe Bembidiina Stephens, 1827. In: Löbl I, Löbl D (Eds) Catalogue of Palearctic Coleoptera. Brill, Leiden, Boston, 294–342.

Motschulsky TV de (1850) Die Kaefer Russlands. Gautier, Moscau, [iv + xi +] 91 pp.

Nguyen L-T, Schmidt HA, von Haeseler A, Minh BQ (2015) IQ-TREE: A fast and effective stochastic algorithm for estimating maximum-likelihood phylogenies. Molecular Biology and Evolution 32: 268–274. https://doi.org/10.1093/molbev/msu300

Pflug JM, Holmes VR, Burrus C, Johnston JS, Maddison DR (2020) Measuring genome sizes using read-depth, k-mers, and flow cytometry: methodological comparisons in beetles (Coleoptera). G3: Genes, Genomes, Genetics 10: 3047–3060. https://doi.org/10.1534/g3.120.401028

Rambaut A, Drummond AJ, Xie D, Baele G, Suchard MA (2018) Posterior summarization in Bayesian phylogenetics using Tracer 1.7. Systematic Biology 67: 901–904. https://doi.org/10.1093/sysbio/syy032

Smith MA, Bertrand C, Crosby K, Eveleigh ES, Fernandez-Triana J, Fisher BL, Gibbs J, Hajibabaei M, Hallwachs W, Hind K, Hrcek J, Huang D-W, Janda M, Janzen DH, Li Y, Miller SE, Packer L, Quicke D, Ratnasingham S, Rodriguez J, Rougerie R, Shaw MR, Sheffield C, Stahlhut JK, Steinké D, Whitfield J, Wood M, Zhou X (2012) *Wolbachia* and DNA barcoding insects: patterns, potential, and problems. PLoS ONE 7: e36514. https://doi.org/10.1371/journal.pone.0036514

Sproul JS, Maddison DR (2017) Cryptic species in the mountaintops: species delimitation and taxonomy of the *Bembidion breve* species group (Carabidae: Coleoptera) aided by genomic architecture of a century-old type specimen. Zoological Journal of the Linnean Society 183(3): 556–583. https://doi.org/10.1093/zoolinnean/zlx076

Thalmann O, Hebler J, Poirier HN, Pääbo S, Vigilant L (2004) Unreliable mtDNA data due to nuclear insertions: a cautionary tale from analysis of humans and other great apes. Molecular Ecology 13: 321–335. https://doi.org/10.1046/j.1365-294X.2003.02070.x

Wild AL, Maddison DR (2008) Evaluating nuclear protein-coding genes for phylogenetic utility in beetles. Molecular Phylogenetics and Evolution 48: 877–891. https://doi.org/10.1016/j.mphye.2008.05.023
Supplementary material 1

Mesquite NEXUS file containing the DNA sequence data and resulting phylogenetic trees from maximum likelihood analyses for the subgenus *Hydrium*

Authors: David R. Maddison
Data type: NEXUS file (DNA sequences, phylogenetic trees)
Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.
Link: https://doi.org/10.3897/zookeys.1007.60012.suppl1

Supplementary material 2

Mesquite NEXUS file containing the DNA sequence data and resulting phylogenetic trees from maximum likelihood analyses for the *Bembidion transversale* group

Authors: David R. Maddison
Data type: NEXUS file (DNA sequences, phylogenetic trees)
Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.
Link: https://doi.org/10.3897/zookeys.1007.60012.suppl2