

Ephemeroptera, Plecoptera, and Trichoptera fauna of Churchill (Manitoba, Canada): insights into biodiversity patterns from DNA barcoding

Authors: Zhou, Xin, Jacobus, Luke M., DeWalt, R. Edward, Adamowicz, Sarah J., and Hebert, Paul D. N.

Source: Journal of the North American Benthological Society, 29(3): 814-837

Published By: Society for Freshwater Science

URL: https://doi.org/10.1899/09-121.1

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at <u>www.bioone.org/terms-of-use</u>.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Ephemeroptera, Plecoptera, and Trichoptera fauna of Churchill (Manitoba, Canada): insights into biodiversity patterns from DNA barcoding

Xin Zhou^{1,4}, Luke M. Jacobus^{2,5}, R. Edward DeWalt^{3,6}, Sarah J. Adamowicz^{1,7}, AND Paul D. N. Hebert^{1,8}

¹ Biodiversity Institute of Ontario, University of Guelph, 50 Stone Road East, Guelph, Ontario, Canada N1G 2W1

² Department of Biology, Indiana University, Bloomington, Indiana 47405 USA ³ Illinois Natural History Survey, 1816 S Oak St., Champaign, Illinois 61820 USA

Abstract. The insect orders Ephemeroptera, Plecoptera, and Trichoptera (EPT) are particularly important for freshwater ecological and biomonitoring studies, but difficulties in their identification to species level impede research. DNA barcoding provides a solution to this problem by linking newly collected specimens to a reference library of authoritatively identified specimens. Here, we consider the ways in which patterns of intraspecific and interspecific genetic divergences in the barcode region can provide rapid insights into the taxonomic identity, morphological features, and geographical distributions of species. Our study led to a $>5\times$ increase in the EPT fauna, including 68 caddisfly, 37 mayfly, and 7 stonefly species, recorded from Churchill. DNA barcoding also aided detection of rare taxa, allowed identification of otherwise unidentifiable life stages, revealed several potentially new species of caddisflies and mayflies, and suggested the presence of cryptic species. The new insights into this fauna and the strong congruence between morphological and molecular characters affirm the utility of DNA barcoding for rapid characterization of the diversity of EPT faunas. We also explore the phenology and habitat preferences of Churchill's trichopterans and demonstrate that comprehensive sampling is important for documenting biodiversity through DNA barcoding.

Key words: aquatic insects, COI, biodiversity inventory, species boundaries, species checklist, phenology, habitat preference.

Community ecology studies on Arctic aquatic insects have been limited despite their value for understanding the impacts of climate change (Quinlan et al. 2005, Wrona et al. 2006a, b), for monitoring water quality (Bowman et al. 2009, Rosenberg and Resh 1993), and for improving knowledge of biodiversity in the Arctic. Among other practical difficulties, such as funding and availability of expertise, species identification using morphological characters presents a serious impediment to community analysis and is one of the reasons why so few studies have been conducted. This impediment arises because description of most aquatic insect species has been

- ⁵ luke.jacobus@gmail.com
- 6 edewalt@inhs.uiuc.edu
- ⁷ sadamowi@uoguelph.ca
- ⁸ phebert@uoguelph.ca

based on the morphologically variant life stage, typically adult males (with some exceptions, e.g., in some mayflies). As a result, the identification of females and immature stages often is not possible (although mayfly larvae are often more easily identified than adults). Nevertheless, females typically contribute at least ½ of the adult population, and immature forms are typically the predominant stage in the water.

Characterizing an unknown fauna is always challenging, even in areas with only moderate diversity, because qualified taxonomists are often unavailable, especially for less charismatic taxa that comprise the bulk of biodiversity. Moreover, routine identification of large numbers of specimens is not the primary goal for academic taxonomists (Holzenthal et al. 2010) and diverts them from finding and describing new species, writing identification keys, building electronic resources, and reconstructing phylogenies. Investment in the

⁴ Email: xinzhou@uoguelph.ca

training and employment of additional nonacademic taxonomists is much needed, and deoxyribonucleic acid (DNA) barcoding—the use of a short standardized mitochondrial cytochrome c oxidase subunit I (COI) gene region for animal species discrimination (Hebert et al. 2003)—shows much promise as an alternate route for the rapid analysis of biodiversity. This method simply requires access to sequencing technology and a DNA library against which to compare sequences.

Both global and local studies are underway to establish DNA reference libraries and to document biodiversity using DNA barcodes. Global DNA barcode reference libraries-barcode sequences linked to voucher specimens identified by qualified taxonomists and housed in permanent repositoriesare being built rapidly for selected eukaryote groups, including some key aquatic insect orders, e.g., Ephemeroptera (mayflies), Plecoptera (stoneflies), Trichoptera (caddisflies) (EPT), and Odonata (dragonflies and damselflies). A complementary approach, termed barcoding biotas, is being used to survey all eukaryote species at particular sites. Such endeavors contribute to the global libraries of the target taxonomic groups, and provide a better overall understanding of biodiversity in focal ecosystems. One low-Arctic Canadian site (Churchill, Manitoba) has been chosen as a testing ground for documenting all multicellular life via DNA barcoding. The barcode reference library for EPT groups at Churchill (Zhou et al. 2009) represents the 1st order-level installment of this large-scale initiative. Zhou et al. (2009) investigated the overall genetic divergence patterns in COI sequences and demonstrated the utility of DNA barcode clusters for constructing species accumulation curves as a complementary approach to documenting unknown local diversity. Species boundaries and life-history traits of the Churchill EPTs are discussed in detail in the present work.

Churchill is an ideal location for an intensive study of the EPT fauna because of its modest taxonomic diversity, complex biogeographical linkages reflecting its location at the transition zone between the boreal and tundra biomes, and its wide variety of habitats. The drainage system at Churchill is characterized by abundant lentic water bodies (ponds, lakes, and temporary pools), small streams, and one major river, the Churchill River. Tundra ponds and lakes are the predominant freshwater habitats by area. Most of these habitats form isolated pools, but some are linked by fast, shallow creeks. The Churchill River is a large, slow-flowing river with a width of 1 km where it empties into Hudson Bay.

During our previous work on Churchill's EPT fauna, DNA barcodes greatly facilitated sorting of

morphospecies, differentiation of closely related species, discovery of potentially new species, and revelation of cryptic species. The present paper examines genetic divergence patterns in greater detail and explores how DNA barcodes perform in discriminating among EPT species compared to using morphological species boundaries. In addition, an updated checklist for Churchill EPT species is provided. During the course of this work, clear differences among species in phenology and habitat preferences were observed. Because caddisflies, the most species-rich order among EPTs at Churchill, were collected intensively for 5 seasons, the temporal and spatial distributions of the Churchill trichopterans were analyzed. Last, the feasibility of conducting rapid biotic surveys-initially using DNA barcodes alone-is discussed. Several critical factors that might affect the efficiency of efforts to barcode entire biotas are summarized.

Methods

Specimen collection, identification, and sequencing

We first compiled prior records for the EPT fauna at Churchill to generate a draft checklist. This list was subsequently enlarged through collection programs led by researchers at the Biodiversity Institute of Ontario and by collaborators from 2002 to 2007 (except 2003, when none were made). The timing of expeditions in different years was varied to help ensure seasonal coverage. Adult samples were collected using ultraviolet (UV) light traps, sweep and aerial nets, Malaise traps, and pitfall traps. Larval samples were collected with a kicknet and by handpicking. Sampling efforts were structured to maximize microhabitat diversity. Adult specimens were pinned or preserved in 95% ethanol, and all larval samples were kept in 95% ethanol. Trichoptera specimens were deposited in the Biodiversity Institute of Ontario, University of Guelph, at the University of Manitoba, and in the University of Minnesota Insect Collection. Ephemeroptera and Plecoptera specimens were deposited in the Biodiversity Institute of Ontario.

EPT specimens were identified with current morphological keys by 3 authors of this paper (XZ, LMJ, and RED). Representative specimens of certain caddisfly species also were examined by R. Blahnik, J. Morse, and D. Ruiter, and burrowing mayfly identifications were confirmed by W. P. McCafferty. Morphological identifications were independent of molecular analysis. When morphological identification was impossible (e.g., some larvae, females, and damaged specimens), DNA barcodes were used to associate these specimens with identified representatives of the taxon in question.

Sequences analyzed in our study have been published in a companion paper (Zhou et al. 2009). COI barcodes were acquired at the Canadian Centre for DNA Barcoding at the University of Guelph. Standard barcoding protocols for DNA extraction (Ivanova et al. 2006), polymerase chain reaction (PCR) amplification, and sequencing (deWaard et al. 2008, Hajibabaei et al. 2005) were followed. Detailed DNA analysis protocols and a new reverse primer designed for recovering a shorter fragment (325 base pairs [bp] on the 5' end) of the barcode region for EPTs are provided in Zhou et al. (2009).

Barcode sequences and associated trace files and voucher information (taxonomy, image, collection information) are accessible in the Barcode of Life Data System (BOLD) (Ratnasingham and Hebert 2007), within the projects 'Ephemeroptera of Churchill', 'Plecoptera of Churchill', and 'Trichoptera of Churchill 2002/2004/2005/2006/2007'. COI sequences are also available in GenBank under accession numbers GU113533–GU115809. Detailed voucher information for all analyzed EPT specimens also is provided in supplementary table S1 in the companion paper by Zhou et al. (2009).

Genetic data analysis

COI sequences for each order were aligned in MEGA 4.0 (Tamura et al. 2007) with the integrated ClustalX method with default parameters, and alignments were verified using amino acid translation. Unique haplotypes were recognized using analytical tools available at the DNA Barcoding Tools website (www.ibarcode.org; Singer and Hajibabaei 2009) and imported into MEGA for tree construction using the Neighbor-Joining method (Saitou and Nei 1987) with pairwise deletion of missing sites and Kimura-2-Parameter (K2P) distances (Kimura 1980). Bootstrap values were obtained using 1000 replicates.

Genetic distances were calculated using the Nearest Neighbor Summary option in BOLD with K2P distances for all sequences >420 bp to examine intraspecific vs interspecific divergence patterns. Maximum intraspecific divergences and minimum distances to nearest neighbors were grouped into a series of categories: 0– 1%, 1–2%, 2–3%, etc. The frequencies of each of these categories were plotted for each of the 3 orders.

Species discovery in caddisflies: temporal and spatial pattern analyses

All caddisfly samples collected from 2002 to 2007 were used to examine the extent of temporal overlap

in species occurrence. Pooled across years, the emergence season was divided into 3 nearly equal periods (P1–P3, each consisting of 18–20 d). Overlap in species composition between these 3 periods was displayed with a Venn diagram.

All 2007 Trichoptera samples were collected within a 1-mo period and with detailed habitat information and, thus, were used in an analysis of habitat use. Three habitat types were designated for the Churchill area: tundra lentic environments (collected adjacent to ponds or lakes), tundra creeks (collected adjacent to fast-moving creeks), and river (collected next to the Churchill River). The division of Churchill's aquatic habitats is not based on their surface areas or water volumes, but rather on their physical attributes, which are most important in determining the distribution of caddisfly larvae (e.g., Wiggins 1996). Overlap in species composition also was displayed with a Venn diagram.

Caddisflies were most intensively collected in 2006 and 2007 (84% of total caddisfly specimens), and the collecting dates in these 2 y were complementary (5– 26 August 2006 and 30 June–27 July 2007). Thus, the 2006 and 2007 samples were used to build species accumulation curves to explore the trends in species occurrence across the season. Accumulated species number, counting all individuals, was plotted against collection dates (sampling bias uncorrected). To eliminate potential bias caused by heterogeneity in species abundance or collecting/processing effort, the accumulated species number divided by the total number of individuals collected on each collection date also was included (sampling bias corrected) in the same figure.

Results

Tracing species boundaries in Churchill EPTs with DNA barcodes

A total of 2277 COI sequences were collected from 2436 individuals, including 68 caddisfly, 37 mayfly, and 7 stonefly species (morphologically distinguishable taxa confirmed by DNA barcodes, summarized in Table 1; but see later discussion regarding 2 *Nemoura* species). COI was effective for separating all species of EPTs at Churchill. Members of each morphological species formed a monophyletic cluster in the COI Neighbor-Joining tree. Furthermore, all morphospecies with multiple representatives had bootstrap values >99%, except the caddisfly *Limnephilus partitus* Walker (92%) (Figs 1–3). A few morphological species showed relatively high intraspecific divergence (Table 1) with multiple haplotype groups represented in such taxa. Those morphospecies with intraspecific divergence

Species	No. seq	Mean Intra	Max Intra	Nearest neighbor	Distance to NN	Processing ID of NN	Dist	Literature
Trichoptera Apataniidae *							IN FI	10000 D
* Apatania crymophila McLachlan							N-H	KOSS 1938
Apatania stigmatella (Tottoretodt)	2	0.80	0.80	Apatania zonella	10.47	CUCAD446-07	N-H	Lehmkuhl and Kerst 1979
Δ Apatania zonella (Zetterstedt) Brochrosseti Actor	1	N/A	N/A	Apatania stigmatella	10.47	CUCAD538-07	N-H	
Brachycentrus		0.00	0.00	Brachycentrus	13.02	CUCAD742-07	N-H	
Brachycentrus (Datuks) Brachycentrus fuliginosus (Walker)	1	N/A	N/A	Juugunosus Brachycentrus americanus	13.02	CUCAD711-07	NA	
Glossosonna Glossosonna intermedium (Klanalek)	13	0.51	1.54	Glossosoma velonum	11.47	CUCAD736-07	N-H	
Δ Glossosoma velonum		0.74	1.43	Glossosoma	11.47	CUCAD657-07	NA	
Protoptila tenebrosa (Valker) Hudronscrobidae	33	0.18	0.64	nuermeuum Agrypnia macdunnoughi	25.79	CUCAD544-07	NA	
Arctopsyche ladogensis	12	0.00	0.00	Ceratopsyche	21.82	LCHIP116-07	N-H	
(Noterial) Ceratopsyche alhedra (Rosc)	1	N/A	N/A	utternuns Ceratopsyche broata	13.54	CUCAD075-07	NA	
Ceratopsyche alternans (Walker)	101	0.44	1.44	oronta Ceratopsyche alhedra	15.92	CUCAD187-07	NA	Flannagan and Flannagan 1982, Lehmkuhl and Kerst 1979, as <i>Hydropsyche</i>
Ceratopsyche bronta	1	N/A	N/A	Ceratopsyche	11.76	CUCAD073-07	NA	Fecuroutu (Datuss) Flannagan and Flannagan 1000
(Ross) Ceratopsyche vexa (Ross)	1	N/A	N/A	veza Ceratopsyche hronta	11.76	CUCAD075-07	NA	Denning 1943
* Cheumatopsyche analis		I			I		NA	Lehmkuhl and Kerst 1979, as
Cheunatopsyche Cheunatopsyche Underseitid 2005	13	2.38	5.25	Ceratopsyche alhedra	21.33	CUCAD187-07	NA	C. petitu baliks
Hydroptila consimilis	17	0.85	2.70	Hydroptila	20.12	CUCAD779-08	NA	
Hydroptila spatulata	1	N/A	N/A	epututut Hydroptila oonsimilis	20.12	CUCAD793-08	NA	
Ovidethira sn CHI 11				011111101100				

DNA BARCODING SUBARCTIC EPTs

NA H-N Lehmkuhl and Kerst 1979 H-N NA NA NA NA 1964 NA NA H-N								
Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964 T. <i>griseus</i> Banks T. <i>griseus</i> Banks	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964 T. <i>griseus</i> Banks T. <i>griseus</i> Banks	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964 T. <i>Briseus</i> Banks T. <i>Briseus</i> Banks	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964 T. <i>griseus</i> Banks T. <i>griseus</i> Banks	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964 T. <i>griseus</i> Banks T. <i>griseus</i> Banks Lehmkuhl and Kerst 197	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964 T. <i>griseus</i> Banks T. <i>griseus</i> Banks Lehmkuhl and Kerst 197 Lehmkuhl and Kerst 197	Lehmkuhl and Kerst 197 Yamamoto and Wiggins 1964 T. <i>griseus</i> Banks T. <i>griseus</i> Banks Lehmkuhl and Kerst 197 Lehmkuhl and Kerst 197
Yamamoto and Wiggin 1964	Yamamoto and Wiggin 1964	Yamamoto and Wiggin 1964 Lehmkuhl and Kerst 15 T. <i>griseus</i> Banks	Yamamoto and Wiggin 1964 Lehmkuhl and Kerst 15 T. <i>griseus</i> Banks	Yamamoto and Wiggin 1964 Lehmkuhl and Kerst 15 <i>T. griseus</i> Banks	Yamamoto and Wiggin 1964 Lehmkuhl and Kerst 15 T. griseus Banks	Yamamoto and Wiggin 1964 Lehmkuhl and Kerst 15 <i>T. griseus</i> Banks	Yamamoto and Wiggin 1964 Lehmkuhl and Kerst 15 <i>T. griseus</i> Banks Lehmkuhl and Kerst 19	Yamamoto and Wiggin 1964 Lehmkuhl and Kerst 15 <i>T. griseus</i> Banks Lehmkuhl and Kerst 14
964 964	964 964	964 mmoto and Wiggins 964 mkuhl and Kerst 19	964 wiggins mukuhl and Kerst 19.	964 mmoto and Wiggins 964 mmkuhl and Kerst 19 <i>Sriseus</i> Banks	964 mmoto and Wiggins 964 mkuhl and Kerst 19 <i>Sriseus</i> Banks	964 and Wiggins mkuhl and Kerst 19 . <i>Sriseus</i> Banks mkuhl and Kerst 19	964 unkuhl and Wiggins . <i>Sriseus</i> Banks mkuhl and Kerst 19 mkuhl and Kerst 19	964 unkuhl and Wiggins . <i>Sriseus</i> Banks . <i>Banks</i> . mkuhl and Kerst 19
Yamamoto and Wigg 1964	Yamamoto and Wigg 1964	Yamamoto and Wigg 1964 Lehmkuhl and Kerst T. <i>griseus</i> Banks	Yamamoto and Wigg 1964 Lehmkuhl and Kerst <i>T. griseus</i> Banks	Yamamoto and Wigg 1964 Lehmkuhl and Kerst <i>T. griseus</i> Banks Lehmkuhl and Kerst	Yamamoto and Wigg 1964 Lehmkuhl and Kerst <i>T. griseus</i> Banks Lehmkuhl and Kerst			
1964 1964	1964 1964	1964 tamamoto and Wigg ehmkuhl and Kerst T. griseus Banks	1964 1964 and Wigg Lehmkuhl and Kerst <i>T. griseus</i> Banks	1964 camamoto and Wigg Lehmkuhl and Kerst T. griseus Banks	1964 Lehmkuhl and Kerst <i>T. griseus</i> Banks	1964 and Wigg Lehmkuhl and Kerst <i>T. griseus</i> Banks	1964 and Wigg Lehmkuhl and Kerst <i>T. griseus</i> Banks Lehmkuhl and Kerst	1964 and Wigg Lehmkuhl and Kerst <i>T. griseus</i> Banks Lehmkuhl and Kerst
+00-1	+001	170 1 Lehmkuhl and Kerst <i>T. griseus</i> Banks	170 1 Lehmkuhl and Kerst <i>T. griscus</i> Banks	170 1 Lehmkuhl and Kerst <i>T. griseus</i> Banks	Lehmkuhl and Kerst <i>T. griseus</i> Banks	Lehmkuhl and Kerst <i>T. griscus</i> Banks Lehmkuhl and Kerst	Lehmkuhl and Kerst <i>T. griscus</i> Banks Lehmkuhl and Kerst	Lehmkuhl and Kerst <i>T. griscus</i> Banks Lehmkuhl and Kerst
		Lehmkuhl and Kers T. griseus Banks	Lehmkuhl and Kers T. <i>griseus</i> Banks	Lehmkuhl and Kers <i>T. griseus</i> Banks	Lehmkuhl and Kers T. <i>griseus</i> Banks	Lehmkuhl and Kers T. <i>griseus</i> Banks Lehmkuhl and Kers	Lehmkuhl and Kers T. <i>griseus</i> Banks Lehmkuhl and Kers	Lehmkuhl and Kers <i>T. griseus</i> Banks Lehmkuhl and Kers
		Lehmkuhl and Ke T. <i>griseus</i> Bank	Lehmkuhl and Ke T. <i>griseus</i> Bank	Lehmkuhl and Ke <i>T. griseus</i> Bankı	Lehmkuhl and Ke T. <i>griseus</i> Bankı	Lehmkuhl and Ke T. <i>griseus</i> Bank	Lehmkuhl and Ke T. <i>griseus</i> Bank	Lehmkuhl and Ke T. <i>griseus</i> Bank
		Lehmkuhl ar T. griseus I	Lehmkuhl an T. <i>griseus</i> F	Lehmkuhl ar T. griseus I	Lehmkuhl ar T. griseus I	Lehmkuhl an T. griseus I	Lehmkuhl ar T. griseus F	Lehmkuhl ar T. griseus F
N-H	N-H							
1717107	CUCAD161-07	AD161-07 AD729-07	AD161-07 AD729-07	AD161-07 AD729-07 RI534-07	AD161-07 AD729-07 RI534-07 088-05	AD161-07 AD729-07 RI534-07 .088-05 	AD161-07 AD729-07 RI534-07 088-05 —	AD161-07 AD729-07 RI534-07 088-05 RI362-07 AD402-07
		CUCAD729-07	CUCAD7294	CUCAD729-6 DSTRI534-07	CUCAD7294 DSTRI534-07 COC088-05	CUCAD7294 DSTRI534-07 COC088-05	CUCAD729-(CUCAD729-(DSTRI534-07 COC088-05 — DSTRI362-07	CUCAD729-07 DSTRI534-07 COC088-05 — DSTRI362-07 CUCAD402-07
7 45	7.45	7.45 17.00	7.45 17.00	7.45 17.00 12.66	.45 .00 .66 .38			
CHUZ Oscatio of	CHU2 Oecetis cf. ochracea	сиа			7 7 12 10	7.45 17.00 12.66 10.38	7.45 17.00 12.66 10.38 	7.45 17.00 12.66 10.38 9.34 9.34
0.30	0	CHU2 Oecetis cf. ochracea CHU1 Oecetis cf.	CHU2 Oecetis cf. ochracea CHU1 Oecetis cf. inconspicua CHU2	CHU2 Oecetis cf. ochracea CHU1 Oecetis cf. inconspicua CHU2 Asynarchus montanus	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	i 1 s cua us 1 s 1 s s 1 s s 1	r cua s us us s s s s s s s s s s s s s s s
	0.3	0.30 Oecetis cf. ochracea CHU1 0.15 Oecetis cf. inconspi	0000	O_{ℓ}	CHU2 Oecetis cf. ochracea CHU1 Oecetis cf. inconspicua CHU2 Asynarchus Masynarchus Limnephilus Limnephilus 1	CHU2 Oecetis cf. ochracea CHU1 Oecetis cf. inconspicua CHU2 Asynarchus Montanus Limnephilus I hageni -	Asynarchus <i>Asynarchus</i> <i>Linnephilus</i> <i>Asynarchus</i> <i>Innephilus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>Asynarchus</i> <i>As</i>	Asynarchus Asynarchus CHU1 Oecetis cf. inconspicua Asynarchus Limnephilus Asynarchus montanus Asynarchus Asynarchus Mageni Mageni Asynarchus
\subset	0.20 0.3	0.30 <i>Oe</i> 0.15 <i>Oe</i>	0.30 <i>Oe</i> 0.15 <i>Oe</i>	0.30 Oe 0.15 Oe 0.96 As	0.30 Occetis cf. ochracea CHU1 0.15 Occetis cf. inconspicua CHU2 0.96 Asynarchus 1.43 Limnephilus 1.43 Limnephilus 1.43 Limnephilus	0.30 Oecetis cf. CHU1 0.15 Oecetis cf. inconspicua 1.43 Limnephilus 1.43 Limnephilus 1.43 Limnephilus 1.43 Limnephilus 1.43 Limnephilus	0.30 Oecetis cf. CHU1 0.15 Oecetis cf. 11.43 Decetis cf. 1.43 Limnephilus 0.66 Asynarchus 1.64 Asynarchus 1.64 Asynarchus 0.66 Asynarchus	 0.30 Occetis cf. 0.15 Occetis cf. 0.15 Occetis cf. 0.15 Occetis cf. 1.43 inconspicua 1.43 Limnephilus 1.43 Limnephilus 1.43 Limnephilus 1.43 Limnephilus 1.43 montanus 3.92 Asynarchus
		0.20 0.30 <i>Oe</i> 0.10 0.15 <i>Oe</i>	0.20 0.30 <i>Oe</i> 0.10 0.15 <i>Oe</i>	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.20 0.30 Oecetis cf. 0.10 0.15 ochracea 0.10 0.15 Oecetis cf. 1 0.10 0.15 Oecetis cf. 1 0.20 0.96 Asynarchus 1 0.33 1.43 Linnephilus 1	0.20 0.30 Oecetis cf. 0.10 0.15 Oecetis cf. 1 0.10 0.15 Oecetis cf. 1 inconspicua 0.20 0.96 Asynarchus 1 0.33 1.43 Limmephilus 1 	0.20 0.30 Oecetis cf. 0.10 0.15 ochracea 0.10 0.15 Oecetis cf. 1 0.10 0.15 Oecetis cf. 1 0.20 0.96 Asynarchus 1 0.33 1.43 Linnephilus 1 0.33 1.43 Linnephilus 1 0.66 Asynarchus 1	0.20 0.30 Oecetis cf. 0.10 0.15 Ochracea 0.10 0.15 Ocertis cf. 1 0.10 0.15 Oecetis cf. 1 0.20 0.96 Asynarchus 1 0.33 1.43 Linnephilus 1 0.33 1.43 Linnephilus 1 0.66 Asynarchus 1 1 0.65 Asynarchus 1 1 0.68 0.66 Asynarchus 1

Species	0	No. seq	Mean Intra	Max Intra	Nearest neighbor	Distance to NN	Processing ID of NN	Dist	Literature
	Asynarchus rossi	35	0.00	0.00	Anabolia	14.68	DSTRI376-07	NA	
	(Leonard & Leonard)				bimaculata				
	Grammotaulius interrogationis (7ettereted+)	119	0.63	2.83	Limnephilus picturatus	9.92	DSTRI580-07	NA	
∇	(Doming) (Doming)	4	1.01	2.01	Arctopora	11.22	COC034-05	NA	
	Limnephilus canadensis	1	N/A	N/A	putchetta Limnephilus	7.06	MHTRI011-06	NA	
	Dauks Limnephilus externus Urccon	66	0.09	0.62	perpusuus Limnephilus	9.66	DSTRI512-07	N-H	
	Limnephilus femoralis Vistere	94	0.29	1.20	Arctopora	11.58	COC034-05	N-H	Fischer 1968
∇	Limnephilus fischeri Diitor	13	0.13	0.31	puicneuu Limnephilus Laccari	3.46	GUCAD147-08	NA	
∇	Limmephilus hageni Daala	122	0.35	1.43	tiugent Limnephilus Godiani	3.46	DSCNI047-07	NA	
	Dauks Limnephilus infernalis	17	0.00	0.00	Jischert Limnephilus	69.6	DSTRI414-07	NA	
	(Dauks) Limnephilus kennicotti Boole	14	0.13	0.32	externus Limnephilus	9.51	CUCAD289-07	NA	
	Limnephilus moestus Barte	б	0.10	0.15	ser neus Limnephilus canadancie	11.42	CUCAD186-07	NA	
	Limnephilus nigriceps	Ŋ	0.33	0.83	Limnephilus	5.75	DSTRI582-07	N-H	
	(zeuersteut) Limmephilus partitus Walber	12	0.89	1.72	picturutus Limnephilus sausoni	4.09	DSTR1471-07	NA	
	Limnephilus perpusillus Walker	7	0.31	0.31	Limnephilus canadensis	7.06	CUCAD186-07	NA	Flannagan and Flannagan 1982
	Limnephilus picturatus McI achlan	9	0.29	0.62	Limnephilus nioricens	5.75	DSTRI512-07	N-H	1
	Limnephilus rhombicus (Limneeus)	μ	N/A	N/A	Limnephilus externus	10.44	DSTRI075-06	N-H	
∇	Limnephilus sansoni Banks	87	0.37	4.01	Limnephilus nartitus	4.09	DSTRI604-07	NA	
	Limnephilus sericeus	б	0.10	0.15	Limnephilus kennicotti	9.51	DSCN1046-07	N-H	
*	Nemotaulius hostilis							NA	Flannagan and Flannagan
	(Indecity) Onocosmoecus unicolor (Barles)	1	N/A	N/A	Asynarchus	17.73	COC010-05	N-H	1704, 11111 1700
	Philarctus bergrothi McLachlan	9	0.00	0.00	Limnephilus kennicotti	14.05	DSTRI563-07	N-H	Lehmkuhl and Kerst 1979, as P. quaeris (Milne)

TABLE 1. Continued.

Species	No. seq	Mean Intra	Max Intra	Nearest neighbor	Distance to NN	Processing ID of NN	Dist	Literature
Molannidae Molanna flavicornis Banks	45	0.02	0.36	Agrypnia obsoleta	21.74	CUCAD379-07	N-H	Lehmkuhl and Kerst 1979
Philopotamidae Cliimarra socia Hagen	1	N/A	N/A	Hydroptila consimilis	25.76	CUCAD796-08	NA	
Phryganeidae Agrypnia colorata	71	0.07	0.92	Agrypnia	13.36	COC041-05	N-H	
Hagen Agrypnia deflata	15	0.05	0.15	agrypnia Agrypnia	3.45	CUCAD379-07	NA	
(Muture) Agrypnia improba	7	0.00	0.00	obsoleta Agrypnia	8.57	CUCAD524-07	NA	
(rtagen) Agrypnia macdunnoughi	14	0.42	1.54	straminea Agrypnia straminea	10.70	CUCAD145-07	NA	
Δ Agrypnia obsoleta	1	N/A	N/A	Agrypnia deflata	3.45	COC040-05	N-H	
* Agrypnia pagetana				I			N-H	Wiggins 1960
Agrypnia straminea	32	0.01	0.18	Agrypnia	8.57	COC041-05	NA	Lehmkuhl and Kerst 1979
nagen Banksiola crotchi Banks	21	0.44	1.07	aruprova Agrypnia	11.76	CUCAD572-07	NA	
Phryganea cinerea	6	0.16	0.30	mucuunnougni Banksiola crotchi	15.23	CUCAD355-07	NA	
Walker Ptilostomis semifasciata (Sav)	9	0.15	0.46	Banksiola crotchi	12.60	CUCAD100-07	NA	
Polycentropodidae Neureclipsis crepuscularis	ю	0.00	0.00	Polycentropus aureolus	19.92	DSTRI586-07	NA	
(Walker) Polycentropus aureolus (Banks)	4	0.00	0.00	Psychomyia flavida	19.90	CUCAD325-07	NA	
Fsychomyndae Psychomyia flavida Hagen Dharroschiida	4	0.23	0.46	Polycentropus aureolus	19.90	DSTRI586-07	N-H	Lehmkuhl and Kerst 1979
Kiyacopriitidae Rhyacophila angelita	41	0.01	0.17	Agrypnia	22.54	COC041-05	NA	
Datities Average for Trichoptera Ephemeroptera		0.34	0.98	improvu	12.21			
Acentrella turbida	1	N/A	N/A	Acentrella	19.36	EPCHU531-07		
(INICLUUTIOUGII) D Acentrella turbida (McDunnoush) E	7	0.15	0.15	turouu E Plauditus sp. CHU1	18.89	EPCHU506-07		

X. Zhou et al.

[Volume 29

No. Mean Max Intra Max Intra pygmaea 7 1.23 1.93 Pr seq Intra Max Max Pr sp. CHU1 9 0.00 0.00 Ba p. CHU1 11 0.59 1.12 Ac p. CHU1 11 0.59 1.12 Ac p. CHU1 11 0.59 1.12 Ac meicolor 59 0.20 1.86 Ba nough 4 0.15 0.30 Ba dyae 4 0.15 0.30 Ba anough 3 0.31 0.78 Pr anough 5 0.15 0.30 Pa anough 5 0.15 0.30 Pa mough 5 0.15 0.30 Pa anough 5 0.16 Oa Oa anough 1 0.03 0.016 Oa					TABLE 1. Continued.	nued.			
Accrpenta pygrated 7 123 193 Proclocen fragite 2.02 Accrpenta sp. CHU1 9 0.00 0.00 Baetidae sp. CHU1 20.25 Accrpenta sp. CHU1 11 0.59 1.12 Accrpenta sp. CHU1 2.77 Baetida sp. CHU1 11 0.59 1.12 Accrpenta sp. 2.77 2.77 Baetis brunnicolor 59 0.20 1.86 Baetis brundy 2.77 Baetis brundyac 4 0.15 0.30 Baetis brundyac 4.24 Baetis protexa 176 0.01 0.78 Proclocon 20.35 Baetis protexa 176 0.01 0.78 Proclocon 4.24 NcDunwough 176 0.01 0.78 Proclocon 13.03 Proclocon inanum - - - - - - - Proclocon inanum - - - - - - - - - - - - - -	Species	No. seq	Mean Intra	Max Intra	Nearest neighbor	Distance to NN	Processing ID of NN	Dist	Literature
Actronagent Actronagent 2.77 Baetidae sp. CHU1 11 0.59 1.12 Actronamesp. 2.77 Baetidae sp. CHU1 11 0.59 1.12 Actronamesp. 2.77 Baetis brunnough 59 0.20 1.86 Baetis brundone 4.24 Baetis brundone 4 0.15 0.30 Baetis brundone 4.24 Baetis phoebus 176 0.01 0.78 Procheen 4.24 Baetis phoebus 176 0.01 0.78 Procheen 4.24 Baetis phoebus 176 0.01 0.78 Procheen 2.77 Baetis phoebus 176 0.01 0.78 Procheen 2.24 Baetis phoebus 176 0.01 0.78 Procheen 2.036 Provideen titut 2 0.30 Procheen 2.036 1.1789 Procheen titut 2 0.30 Procheen fundar 1.789 1.040 Procheen titut 2 0.30	Acerpenna pygmaea		1.23	1.93	Procloeon fragile	20.25	EPCHU141-07	NA	Harper and Harper 1981
Bactidae sp. CHU111 0.59 1.12 Accrements primation 2.77 Bactidae sp. CHU111 0.59 1.20 1.20 1.20 1.20 1.20 Bactis bundyae4 0.15 0.30 Bactis bundyae 4.24 1.303 Bactis bundyae4 0.15 0.30 Bactis bundyae 4.24 1.303 Bactis ploebus 1.76 0.01 0.78 $Bactis bundyae4.241.303Bactis ploebus1.760.010.78Bactis bundyae4.241.893McDumougha1.760.010.78Pacolocon20.551.80Plauditus sp. CHU120.000.00Accntrella20.361.404Plauditus sp. CHU120.000.00Accntrella20.361.1789Plauditus sp. CHU120.120.30Paclocon1.7891.789Plauditus sp. CHU120.000.00Accntrella1.889Proclocon mandax110.080.31Proclocon1.789Proclocon mandax110.080.31Proclocon madax1.789Proclocon mandax0.160.30Proclocon madax1.789Proclocon mandax0.160.30Proclocon madax1.789Proclocon mandax0.160.30Proclocon madax1.789Proclocon mandax0.160.30Proclocon madax$	(наgen) Acerpenna sp. CHU1	6	0.00	0.00	Baetidae sp. CHI 11	2.77	EPCHU340-07		
Bactis brunneicolor 59 0.20 1.86 Bactis bundyne 4.24 McDumnough 4 0.15 0.30 Bactis bundyne 4.24 Bactis bundyne 4 0.15 0.30 Bactis bundyne 4.24 Bactis phoebus 176 0.01 0.78 Procloson 20.36 Planditius sp. CHU1 2 0.00 0.00 Acentrella 18.89 Planditius sp. CHU2 7 0.15 0.30 Procloson 17.89 Procloson fragite 13 0.30 Procloson fragite 17.89 Procloson fragite 11 0.08 0.31 Procloson fragite 17.89 McDunnough 11 0.08 0.31 Procloson fragite 17.89 McDunnough 11 0.08 0.31 Procloson fragite 17.89	Baetidae sp. CHU1	11	0.59	1.12	Acerpenna sp.	2.77	EPCHU374-07		
Baetis bundyae 4 0.15 0.30 Baetis bundyae 4.24 Lehmkuhl 3 0.31 0.46 Baetis bundyae 4.24 Baetis phoetus 176 0.01 0.78 Broclecon 20.55 Baetis phoetus 176 0.01 0.78 Broclecon 20.55 Baetis phoetus 176 0.01 0.78 Proclecon 20.55 Roburnough 25 0.99 2.80 Acentrelia 20.36 Plauditus sp. CHU1 2 0.00 0.00 Acentrelia 20.36 Proclocon inantum - - - - - - Proclocon fragile 5 0.12 0.30 Proclocon 17.89 Proclocon mendax 11 0.08 0.31 Proclocon fragile 17.89 Baetiscidae 8 0.15 0.30 Proclocon fragile 17.89 McDumough 5 0.12 0.30 Proclocon fragile 17.89 Proclocon mendax 11 0.08 0.31 Proclocon fragile 17.89	Baetis brunneicolor McDunnough	59	0.20	1.86	Baetis hudsonicus	13.03	EPCHU086-07	NA	Harper and Harper 1981, McClure 1943, Morihara
Baetis hudsonicus Ide 3 0.31 0.46 Baetis bunduas 4.24 Baetis plochus 176 0.01 0.78 Proclocon 20.35 Baetis plochus 176 0.01 0.78 Proclocon 20.36 Plauditus cf. dubius 25 0.99 2.80 Acentrelia 20.36 Plauditus sp. CHU1 2 0.00 0.00 Acentrelia 20.36 Plauditus sp. CHU1 2 0.15 0.30 Parcheon 20.36 Plauditus sp. CHU1 2 0.00 0.00 Acentrelia 20.36 Proclocon inanum - - - - - - Proclocon inanum - - - - - - - McDunnough 5 0.12 0.30 Proclocon fragile 17.89 Proclocon inanum - - - - - - McDunnough 5 0.12 0.30 Proclocon fragile 17.89 Walsh Proclocon inanum - - - - - McDunnough 5 0.12 0.30 Proclocon fragile 17.89 Walsh Proclocon inducat 0.16	Baetis bundyae Tehmhihl	4	0.15	0.30	Baetis	4.24	EPCHU086-07	NA	and McCatterty 1979 Harper and Harper 1981
Bartis plochus McDumougha1760.010.78Procleeon mendax20.55McDumougha1760.010.78Procleeon20.55McDumougha250.992.80Acentrella20.36Plauditus sp. CHU120.000.00Acentrella20.36Plauditus sp. CHU120.000.00Acentrella20.36Plauditus sp. CHU270.150.30Plauditus sp.20.23Procloeon inanumProcloeon inanumNocloeon inanumRestiscidae <td>Baetis hudsonicus Ide</td> <td>б</td> <td>0.31</td> <td>0.46</td> <td>Baetis bundyae</td> <td>4.24</td> <td>EPCHU319-07</td> <td>NA</td> <td>Ide 1937, McClure 1943, Morihara and McCafferty</td>	Baetis hudsonicus Ide	б	0.31	0.46	Baetis bundyae	4.24	EPCHU319-07	NA	Ide 1937, McClure 1943, Morihara and McCafferty
Plauditus cf. dubius250.992.80Acentrella20.36 $(Walsh)$ $(Walsh)$ $(Walsh)$ 2 0.00 0.00 $Acentrella$ 18.89 $(Walsh)$ $Plauditus sp. CHU1$ 2 0.00 0.00 $Acentrella$ 18.89 $Plauditus sp. CHU2$ 7 0.15 0.30 $Plauditus sp. CHU1$ 2 20.23 $Procloeon inanum (McDunnough)50.120.30Procloeon inandax17.89Procloeon mendax110.080.31Procloeon fragile17.89Procloeon mendax110.080.31Procloeon fragile17.89McDunnough110.080.31Procloeon fragile17.89Procloeon mendax110.080.31Procloeon fragile17.89McDunnoughProcloeon mendax110.080.31Procloeon fragile17.89McDunnoughProcloeon mendax110.080.31Procloeon fragile17.89McDunnoughProcloeon mendax110.080.31Procloeon fragile17.89McDunnoughProcloeon mendax110.080.31Procloeon fragile12.79Procloeon mendax110.000.000.000.000.02Procloeon mendax11N/AN/ADannella simplex12.79$	Baetis phoebus McDunnough ^a	176	0.01	0.78	Procloeon mendax	20.55	EPCHU037-07	NA	Morihara and McCafferty 1979, as <i>B. flavistriga</i>
$V_{audities}$ V_{a	Plauditus cf. dubius	25	0.99	2.80	Acentrella	20.36	EPCHU531-07		Harper and Harper 1981
Plauditus sp. CHU270.150.30Plauditus sp.20.23Procloeon inanumMcDunnough)50.120.30Procloeon17.89Procloeon fragile50.120.30Procloeon17.89Procloeon mendax110.080.31Procloeon fragile17.89Rocloeon mendax110.080.31Procloeon fragile17.89Baetiscidae0.030.16Metretopus20.23Baetiscidae90.000.00Damella12.79Dantella lita (Burks)30.000.00Damella simplex12.79Dannella simplex40.100.15Dannella simplex12.79Dannella simplex1N/AN/ADannella simplex12.79Enrylophella bicolor1N/AN/ADannella simplex20.24Enrylophella bicolor1N/AN/ADannella simplex12.79Enrylophella bicolor1N/AN/ADannella simplex20.24Enrylophella bicolor1N/AN/ADannella bindex20.24Enrylophella bicolor1N/AN/ADannella bindex20.24Enrylophella bicolor1N/AN/ADannella bindex20.24Ephemeridae20.872.17Pannella bindex20.24Ephemeridae20.872.17Pannella bindex20.24Ephemeridae2	Plauditus sp. CHU1	7	0.00	0.00	Acentrella	18.89	EPCHU531-07		
Proclocon inanumProclocon fragile50.120.30Proclocon17.89Proclocon mendax110.080.31Proclocon17.89(McDunnough)70.030.16Metretopus17.89Proclocon mendax110.080.31Proclocon fragile17.89Retrisca laurentina90.030.16Metretopus20.23Baetisca laurentina90.030.16Metretopus20.23Baetisca laurentina90.000.00Dannella simplex12.79Dannella lita (Burks)30.100.15Dannella simplex12.79Dannella simplex1N/AN/ADannella lita20.24Ephemerella needhami160.060.30Ephemera20.24Ephemerella needhami1N/AN/ADannella lita20.24McDunnough1N/AN/ADannella lita20.24Ephemerella needhami160.060.30Ephemera20.24McDunnough1N/AN/ADannella lita20.24Ephemerella needhami1N/ADannella lita20.24McDunnough1N/AN/ADannella lita20.24McDunnough1N/AN/ADannella lita20.24Ephemerella needhami1N/ADannella lita20.24Ephemerella20.872.17Paraleptophlebia16.76 <td>Plauditus sp. CHU2</td> <td></td> <td>0.15</td> <td>0.30</td> <td>plauditus sp. CHU1</td> <td>20.23</td> <td>EPCHU506-07</td> <td></td> <td></td>	Plauditus sp. CHU2		0.15	0.30	plauditus sp. CHU1	20.23	EPCHU506-07		
Procloeon fragile (McDunnough)50.120.30Procloeon17.89Procloeon mendax (Walsh)110.080.31Procloeon fragile17.89Procloeon mendax (Walsh)110.080.31Procloeon fragile17.89Baetiscidae Baetiscidae90.030.16Metretopus20.23Baetiscidae Baetiscidae90.000.00Dannella12.79Baetiscidae Baetiscidae30.000.00Dannella lita12.79Bannella lita (Burks)30.000.15Dannella simplex12.79Dannella lita (Burks)30.000.15Dannella lita12.79Dannella lita (Burks)30.000.15Dannella lita12.79Dannella lita (Burks)160.060.30Ephemera20.24Dannella lita (Burks)1N/AN/ADannella lita20.24Dannella lita (Burks)1N/ADannella lita20.24Dannella lita (Burks)1N/ADannella lita20.24Dannella simplex1N/ADannella lita20.24Ephemerella reedhami160.060.30Ephemera20.24Ephemeridae20.872.17Panleptophlebia16.76DannelaeDannelae1N/ADannelae20.24Ephemeridae20.872.17Panleptophlebia16.76DannelaeDannelaeDannelae116.76 <td>Pr</td> <td>I</td> <td> </td> <td> </td> <td></td> <td> </td> <td> </td> <td>NA</td> <td>Harper and Harper 1981</td>	Pr	I						NA	Harper and Harper 1981
Proclocon fragile110.080.31Proclocon fragile17.89Proclocon mendax110.080.31Proclocon fragile17.89Retrictidae(Walsh)90.030.16Metretopus20.23Baetiscidae90.030.16Metretopus20.23Baetiscidae90.000.00Donealis20.23Ephemerellidae30.000.00Dannella simplex12.79Dannella lita (Burks)30.000.15Dannella lita12.79Dannella simplex40.100.15Dannella lita12.79Dannella simplex160.060.30Ephemera20.24Ephemerella needhami160.060.30Ephemera20.24Ephemeridae1N/AN/ADannella lita23.42Ephemeridae20.872.17Paraleptophlebia16.76Wolloon0.000.872.17Paraleptophlebia16.76	P_{T}	IJ	0.12	0.30	Procloeon	17.89	EPCHU031-07	NA	
Detendance90.030.16Metretopus20.23Baetisca laurentina90.030.16Metretopus20.23Baphemerellidae0.000.000.00Dannella simplex12.79Dannella lita (Burks)30.000.15Dannella lita12.79Dannella lita (Burks)30.000.15Dannella lita12.79Dannella simplex160.060.30Ephemera20.24Dannella simplex1N/AN/ADannella lita23.42Ephemeridae1N/AN/ADannella lita23.42Ephemeridae20.872.17Paraleptophlebia16.76Wolloon0.000.872.17Paraleptophlebia16.76	Pr	11	0.08	0.31	menuux Procloeon fragile	17.89	EPCHU275-07	NA	
Dannella lita (Burks)30.000.00Dannella simplex12.79Dannella simplex40.100.15Dannella simplex12.79Dannella simplex140.100.15Dannella lita12.79(McDunnough)160.060.30Ephemera20.24Ephemerella needhami160.060.30Ephemera20.24McDunnough1N/AN/ADannella lita23.42Clemens)60.872.17Paraleptophlebia16.76Ephemeridae0.872.17Paraleptophlebia16.76Walton0.000.872.17Paraleptophlebia16.76	Bactiscuae Baetisca laurentina Finhemerollidae	6	0.03	0.16	Metretopus borealis	20.23	EPCHU557-07	NA	
Ephemeral and Control160.060.30Ephemera20.24Ephemeral and the acdhami160.060.30Ephemera20.24McDunnough1N/AN/ADannella lita23.42Eurylophella bicolor1N/AN/ADannella lita23.42(Clemens)60.872.17Paraleptophlebia16.76Ephemeridae0.872.17Paraleptophlebia16.76Walloo0.060.872.17Paraleptophlebia16.76		θ4	0.00 0.10	$0.00 \\ 0.15$	Dannella simplex Dannella lita	12.79 12.79	EPCHU001-07 EPCHU545-07	NA NA	Allen and Edmunds 1961
tella bicolor 1 N/A N/A Dannella lita 23.42 nens) 22 0.87 2.17 Paraleptophlebia 16.76		16	0.06	0.30	Ephemera	20.24	EPCHU244-07	NA	
a simulans 22 0.87 2.17 Paraleptophlebia 16.76	Eurylophella bicolor (Clemens)	1	N/A	N/A	summuns Dannella lita	23.42	EPCHU545-07	NA	
	Ephemeruse Ephemera simulans Walker Heptageniidae	22	0.87	2.17	Paraleptophlebia praepedita	16.76	EPCHU229-07	NA	

DNA BARCODING SUBARCTIC EPTs

Species	50	No. seq	Mean Intra	Max Intra	Nearest neighbor	Distance to NN	Processing ID of NN	Dist	Literature
\bigtriangledown	Ecdyonurus criddlei	4	00.00	0.00	Heptageniidae	16.71	EPCHU218-07	NA	
	(McDunnough) Heptagenia pulla (Clemens)	27	0.11	0.76	AZ sp. CHUI Heptageniidae X7 en CHII	16.53	EPCHU218-07	NA	
	Heptageniidae sp.	Т	N/A	N/A	Heptagenia pulla	16.53	EPCHU540-07		
	Heptageniidae sp.	Ц	N/A	N/A	Rhithrogena manifesta	10.65	EPCHU283-07		
	Leucrocuta jewetti	10	0.58	1.26	Ecdyonurus ewiddlei	16.76	EPCHU541-07	NA	
	Maccaffertium	8	0.47	1.11	Leucrocuta Leucrocuta	20.01	EPCHU553-07	NA	
*	Maccaffertium vicarium			I				NA	Bednarik and McCafferty
*	Rhithrogena jejuna Ertor			I				NA	Ide 1954, Jacobus and
\bigtriangledown	Eaton Eaton	Ŋ	0.00	0.00	Heptageniid XZ sp. CHU2	10.65	EPCHU495-07	NA	INICCALLELLY ZUUL
Lep	Leptohyphidae Tricorythodes mosegus Alba-Tercedor & Flannagan	1	N/A	N/A	Ephemera simulans	24.48	EPCHU241-07	NA	
rep	Leptopruebuuae Leptophlebia cupida	З	0.10	0.15	Paraleptophlebia	17.83	EPCHU229-07	NA	
$\nabla \#$	Paraleptophlebia aquilina Harper &	б	0.31	0.46	praepeutu Leptophlebia cupida	21.08	EPCHU021-07	NA	
	Paraleptophlebia debilis	4	0.00	0.00	Paraleptophlebia	5.58	EPCHU179-07	NA	
	Paraleptophlebia	55	0.19	0.97	sp. CIIUI Metretopus boxadia	16.15	EPCHU557-07	NA	Harper and Harper 1981
	Paraleptophlebia sp. CHU1	4	0.00	0.00	portauts Paraleptophlebia debilis	5.58	MHCOL145-07		
	Metreupoutuae Metretopus borealis Circlenaria	9	0.10	0.15	Siphlonurus phyllis	14.41	EPCHU055-07	N-H	Edmunds 1957
rdie	Depution and the chelifer Benoteson	Ц	N/A	N/A	Paraleptophlebia nraenedita	19.02	EPCHU229-07	N-H	
*	Parameletus midas			l	р исрсини -			NA	Harper and Harper 1981
	Siphlonurus alternatus	49	0.08	0.77	Siphlonurus	7.18	EPCHU085-07	N-H	
	Siphlonurus phyllis McDunnough	IJ	0.71	1.38	pnyuus Siphlonurus alternatus	7.18	EPCHU105-07	NA	

822

TABLE 1. Continued.

				TABLE 1. Continued.	nued.			
Species	No. seq	Mean Intra	Max Intra	Nearest neighbor	Distance to NN	Processing ID of NN	Dist	Literature
Average for Ephemeroptera		0.25	0.65		14.97			
Capniidae							IN II	N 201 1- 50
· Capnia nearctica banks · · · · · · · · · · · · · · · · · · ·							N-H	Kicker 1964 Ricker 1938
Newport							-	
Chloroperlidae Hastaperla brevis	13	0.07	0.30	Haploperla orpha	10.89	PLCHU217-08	NA	Frison 1942
(bauks) Haploperla orpha (Frison)	ю	0.20	0.30	Haploperla brevis	10.89	PLCHU206-08	NA	
Nemouridae Amphinemura linda	180	0.34	2.54	Nemoura arctica	17.68	PLCHU218-08	N-H	Ricker 1952
(Kicker) Nemoura arctica Esben-	4	0.00	0.00	Nemoura sp.	6.05	PLCHU222-08	N-H	Ricker 1952
Peterson Nemoura sp. CHU1	ю	0.22	0.35	CHUI Nemoura arctica	6.05	PLCHU218-08		
<i>Nemoura</i> sp. CHU2 * Shipsa rotunda	- I	N/A -	N/A -	Nemoura arctica —	8.79 —	PLCHU218-08 —	N-H	Ricker 1944
(Claassen)								
* Acroneuria carolinensis	Ι				I		NA	Stark and Gaufin 1976
* Acroneuria lycorias	I	I		I	I		NA	Ricker 1964
(Newman) * Claassenia sabulosa (Banks)	I	I		I	I	I	N-H	Ricker 1964
Perlodidae								
* Diura bicaudata					I		N-H	Ricker 1964
* Isogenoides frontalis			I	I		I	NA	Frison 1942
* Isoperla decolorata				I			N-H	Ricker 1944
(vvalker) Isoperla sp.	6	1.30	3.13	Haploperla brevis	20.57	PLCHU206-08		
* Isoperla marlynia Needham and		I				I	NA	Frison 1942
* Isoperla transmarina	I		I		I		NA	Frison 1942
Pteronarcyidae * Pteronarcyidae	I	I	I	I	I	I	NA	McClure 1943, as T. shelfordi
(Say) Average for Plecoptera		0.35	1.10		11.56			Frison
				5 F				

823

^a Baetis phoebus McDunnough is being reinstated from synonymy with B. flavistriga McDunnough by LMJ, XZ, and PDNH, unpublished data

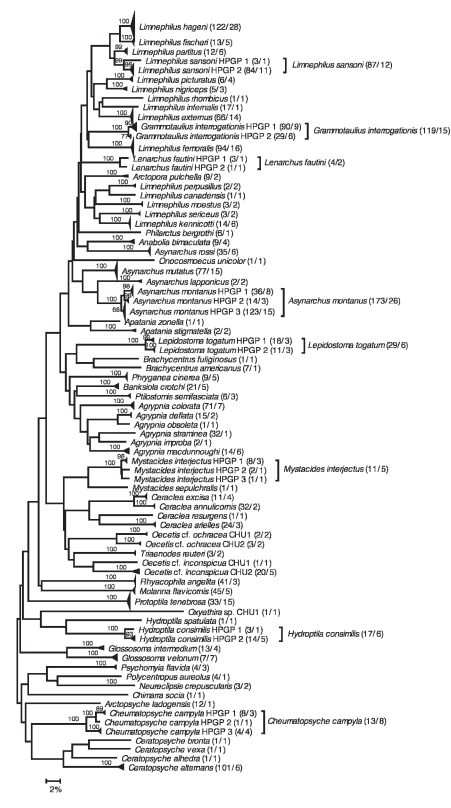


FIG. 1. Neighbor-Joining tree of unique cytochrome c oxidase subunit I (COI) haplotypes of Trichoptera from Churchill. A total of 1500 COI sequences representing 68 species of Trichoptera are represented. Haplotypes of the same morphospecies are collapsed into triangles whose height represents the number of distinct haplotypes and width represents the extent of intraspecific divergence. Species showing relatively large intraspecific divergences (>2%) are highlighted in brackets to the right, and haplotype groups (HPGP) within these species are shown on the tree. Numbers in parentheses represent the total number of sequences and the number of haplotypes for each species or haplotype group. Bootstrap values are provided for all morphospecies and for each distinct haplotype group with multiple representatives. Haplogroups represented by multiple individuals with the same haplotype possess a bootstrap value of 100%, e.g., *Limnephilus infernalis*.

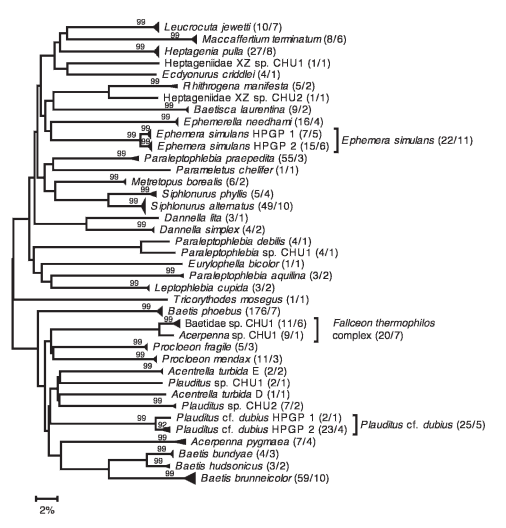


FIG. 2. Neighbor-Joining tree of unique cytochrome c oxidase subunit I (COI) haplotypes of Ephemeroptera from Churchill. A total of 564 COI sequences representing 36 species of Ephemeroptera represented. Figure annotations are the same as those in Fig. 1.

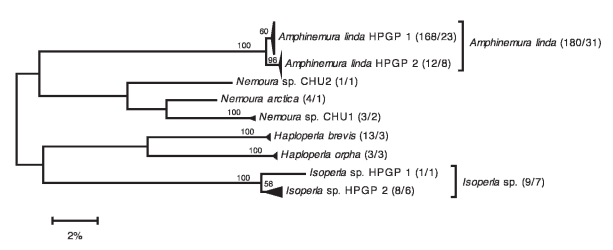


FIG. 3. Neighbor-Joining tree of unique cytochrome c oxidase subunit I (COI) haplotypes of Plecoptera from Churchill. A total of 213 COI sequences representing 7 species of Plecoptera are represented. Figure annotations are the same as those in Fig. 1.

values >2% are shown in brackets in Figs 1-3 (haplogroups within these species are noted as HPGPs). This threshold was used because past studies have shown that intraspecific divergences rarely exceed this value in sympatry (Hebert et al. 2003, 2004, Ball et al. 2005, Hogg et al. 2009). Most cases of large intraspecific divergence reflect cryptic diversity (Williams et al. 2006, Smith et al. 2007, Decaëns and Rougerie 2008, Ståhls and Savolainen 2008, Vaglia et al. 2008, Basquin and Rougerie 2009, Hausmann et al. 2009, Hebert et al. 2009). Of course, exceptions have been observed (Wiemers and Fiedler 2007, Alexander et al. 2009). Therefore, cases of >2% intraspecific divergence are highlighted in our study to emphasize taxa that deserve further attention for potential taxonomic revision (see discussion of morphological characters below), but species names based on morphology are used for all lineages, even those with deep genetic divergence.

For the most part, interspecific distances of Churchill's EPTs did not overlap with intraspecific divergences. In fact, average distance to nearest neighboring species (NN distance) was greater than maximum intraspecific divergences by a factor of $\geq 10 \times$ in each order (Table 1). NN distances were greater than average intraspecific divergences by $\geq 30 \times$. Genetic patterns were similar across the 3 orders. Mean intraspecific divergences for species within each order ranged from 0.25 to 0.35%, whereas the averages of the maximum intraspecific divergences es ranged from 0.65 to 1.10% and NN distances ranged from 11.6 to 14.97%.

The few cases of overlap between intraspecific and interspecific distance values reflected large intraspecific divergences in 3 caddisfly and 2 mayfly species combined with shallow distances between 4 caddisfly and 1 mayfly species pairs (Fig. 4, noted in panels A and B). At least some of these cases probably reflected the presence of cryptic species or a species complex with a recent diversification history (e.g., *Limnephilus sansoni* Banks, *Cheumatopsyche campyla* Ross; see Discussion). In addition, morphological identification of some samples was uncertain (e.g., *Isoperla* sp.; see Discussion), but the relevant samples were temporarily treated as the same morphospecies.

Extended checklists and distribution patterns for Churchill's EPT fauna

Few published studies exist for EPT species at Churchill, and the available records are scattered. McClure (1943) recorded 2 mayfly, 1 stonefly, and several unidentified caddisfly species from this region. For stoneflies, most Churchill records came from the taxonomic treatments, such as Frison (1942), Ricker (1952), and Stark and Gaufin (1976). A study of Churchill caddisflies, motivated by an attempt to solve identification difficulties encountered in an ecological study (Lehmkuhl and Kerst 1979), provided an annotated checklist of 10 species collected from 1971 to 1979. Species records of Churchill EPTs from previous studies are compiled in Table 1 with notes on the original references. Most historical records were extracted from a faunistic review of the EPTs of Manitoba by Flannagan and Flannagan (1982).

Most of the EPT species collected in our study were new records to Churchill, and 18 were new to Manitoba (Table 1), including 1 new record for Canada. Lehmkuhl and Kerst (1979) reported that 7 of the 10 caddisfly species collected at Churchill in their study were endemic to North America. In our study, of the 66 caddisfly species that have been assigned to a nominal species or species complex, 44 are Nearctic, whereas 22 are Holarctic. Three of the identified mayflies (Metretopus borealis Eaton, Parameletus chelifer Bengtsson, Siphlonurus alternatus (Say)) also have Holarctic distributions, but the rest are Nearctic (Table 1). Our collections and literature records suggest that 19 stonefly species occur at Churchill, including those defined by temporary names (Nemoura sp. CHU1, Nemoura sp. CHU2) and the unidentified Isoperla (Table 1). Of the 16 named stonefly species, 3 have Holarctic, circumpolar distributions: Capnia nearctica Banks, Nemoura arctica Esben-Petersen, and Diura bicaudata (Linneaus). Several other species have northern transcontinental distributions: Capnia vernalis Newport, Amphinemura linda (Ricker), Shipsa rotunda (Claassen), Isoperla decolorata (Walker), and Claassenia sabulosa (Banks), and the rest have eastern Nearctic distributions with Churchill as the western limit of their range.

Most of the caddisfly species collected at Churchill have a transcontinental distribution in North America. Eleven species are new for Manitoba (Table 1), among which *Glossosoma velonum* Ross, *Lenarchus fautini* (Denning), *Limnephilus sansoni*, and *Agrypnia obsoleta* (Hagen) represent the easternmost records in North America. *Asynarchus rossi* (Leonard and Leonard) is regarded as a rare species in North America, with a locally abundant disjunct population in Minnesota (Houghton and Holzenthal 2003). This species was collected in Churchill only during 9–26 August 2006, but it was common (n = 46). Five caddisfly species previously recorded from Churchill were not detected during our study (Table 1, marked with *).

Both mayfly larvae and adults were included in our DNA analysis, enabling the association of unknown

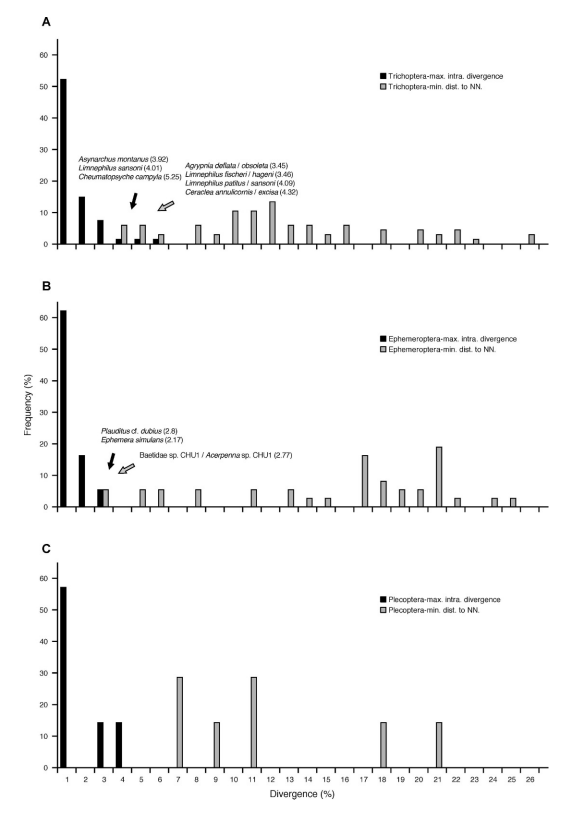


FIG. 4. Intraspecific vs interspecific cytochrome c oxidase subunit I (COI) distances in Trichoptera (A), Ephemeroptera (B), and Plecoptera (C) at Churchill. Frequencies of the maximum intraspecific divergence and minimum distance to nearest neighboring species are plotted for the 3 orders. Identities of the species and nearest neighboring species pairs appearing in the overlapping portion are provided with their divergence values. Arrows indicate cases of overlap between intra- and interspecific distance values. Black and grey arrows in panels A and B indicate species showing large intraspecific divergences and species pairs possessing shallow interspecific distances, respectively. Max. intra. divergence = maximum intraspecific divergence, min. dist. to NN = minimum distance to nearest neighbor. life stages and confirmation of current associations. About 1/3 of the mayfly species in our study were represented by both adults and larvae: Baetis brunneicolor McDunnough, Baetis bundyae Lehmkuhl, Baetis phoebus McDunnough, Baetisca laurentina McDunnough, Dannella lita (Burks), Ecdyonurus criddlei (McDunnough), Ephemera simulans Walker, Heptagenia pulla (Clemens), Leptophlebia cupida (Say), Procloeon fragile (McDunnough), and Siphlonurus alternatus (Say). The larva of *P. fragile* has not been described, but it was associated with adults via barcodes in our study. However, additional material is needed before their formal description because our specimens were damaged. One mayfly species, B. phoebus, is being reinstated from synonymy with B. flavistriga McDunnough based on the present DNA barcode data and concordant morphological evidence from a review of eastern Nearctic species of the genus Baetis Leach (LMJ, XZ, and PDNH, unpublished manuscript). Eight mayfly species are reported from Manitoba for the first time, including 1 new record for Canada (Table 1). Among these, the distribution range of Paraleptophlebia aquilina Harper & Harper, a boreal species previously known only from Oregon (Harper et al. 1995, Harper and Harper 1986, Parsons et al. 1991), is significantly extended eastward. This species is very similar to the eastern US species Paraleptophlebia assimilis (Banks), which might be considered more likely to occur in Manitoba, but the 2 species are differentiated easily by size and color (Harper and Harper 1986). The occurrence of E. criddlei at Churchill represents a significant northeastern range extension from the western US and Alberta, Canada (McDunnough 1927). However, some identifications of Nixe and Ecdyonurus species are problematic in central Canada (Webb and McCafferty 2008). We failed to recollect 4 mayfly species previously recorded from Churchill (Table 1, marked with *), but 2 of these species might be represented in our DNA library by haplotypes that can be identified only to the family Heptageniidae at the moment.

We collected only 5 stonefly species that could be identified to species with morphological traits, and 2 additional genetically distinct Nemourinae (probably *Nemoura*) were each represented by only a larva and females, respectively. The low number of stonefly species reflected our use of collecting techniques that were inefficient for sampling stonefly larvae and that our sampling began too late to collect early-emerging species. For example, 2 early-spring-emerging stoneflies, *Capnia nearctica* and *Capnia vernalis*, occur at Churchill but emerge while snow is still on the ground. Larval exuviae of several large species (Perlidae and Perlodidae of Table 1) were observed during our collections, so we know that these species also occur although we failed to collect them.

Temporal and spatial distributions of Churchill caddisflies

The overall shapes and trends of the bias-corrected (number of accumulated species/specimen) and biasuncorrected (number of accumulated species) species accumulation curves (Fig. 5) were very similar, a result suggesting that the temporal distributions of the detected species are independent of abundances. The shallow slope for the initial sampling dates reflects the combined effect of undersampling (caused by limited collection activity in the early season) and the lower species richness at this time. The subsequent steep slope suggests that this later period is the most critical period for sampling species diversity in Trichoptera.

Emergence of adult Churchill caddisfly species was bimodal with many species emerging around the summer solstice, followed by a lull in activity, then a resumption of species emerging in late summer (Fig. 5). Only 14 species (21%) were collected throughout the entire season (Fig. 6A), and nearly ½ (31/68) were collected within only 1 of the 3 periods. Among caddisfly species detected in 2 periods, adjacent time periods had the greatest overlap in species composition as expected, whereas only 1 species was shared between the earliest and latest time periods.

Of the 68 species of adult caddisflies, 42 were collected from lentic habitats, 29 from the Churchill River, and 8 from small streams (Fig. 6B). Only 4 species (Agrypnia colorata Hagen, Asynarchus montanus (Banks), Ceraclea excisa (Morton), and Mystacides interjectus (Banks)) were detected in all 3 habitats, and these cases might reflect adult dispersal because the larvae of these genera are mostly lentic. Overlap in species was greatest between the Churchill River and lentic habitats, and no species were shared only between creeks and the Churchill River. Only 2 medium- to large-sized caddisflies, Asynarchus mutatus (Hagen) and Phryganea cinerea Walker, were shared between creeks and lentic habitats. They were each represented by a single specimen collected in the creeks, probably because of adult flight. Once larval habitat use is investigated, we expect a substantial decrease in overlap between habitats.

Discussion

We investigated the species diversity of Ephemeroptera, Plecoptera, and Trichoptera (EPTs) at a site in the Canadian subarctic. Patterns of intraspecific and interspecific divergence in DNA barcodes were

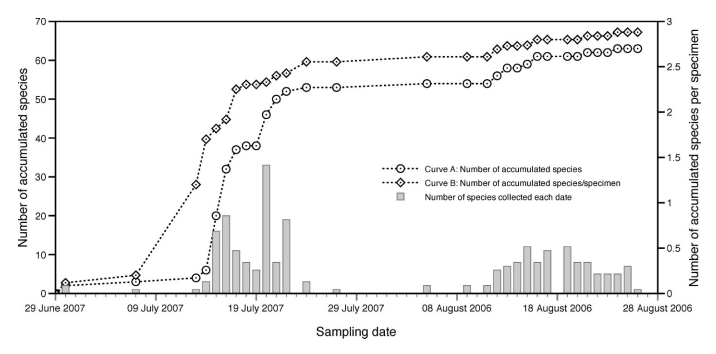


FIG. 5. Temporal distribution of Trichoptera species richness in samples collected in 2006 and 2007. The accumulated number of morphospecies (curve A, represented by circles) is plotted against sampling dates during 5 to 26 August 2006 and 30 June–27 July 2007 to explore the trends in species occurrence in caddisflies at Churchill. The accumulated number of morphospecies divided by the number of samples collected on each day also is included as a correction for potential sampling intensity or abundance bias (curve B).

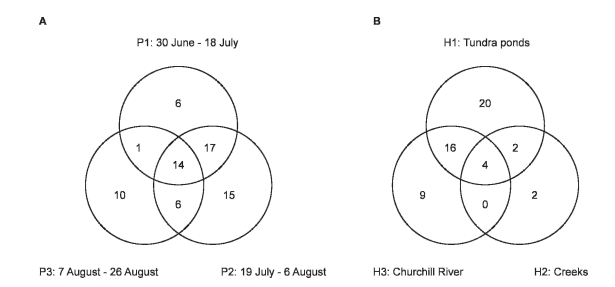


FIG. 6. Temporal overlap and habitat usage in Churchill Trichoptera. A.—Temporal overlap in Trichoptera (2002–2007). Collection dates are divided into 3 periods (P1–P3), each consisting of 18–20 d. The number of species collected during each period is noted in the corresponding area. For example, 14 species were collected throughout P1 to P3, whereas 10 species were collected only during P3. B.—Habitat preference in Trichoptera (2007). Aquatic habitats at Churchill are categorized into 3 types: tundra ponds, tundra creeks, and the Churchill River. The number of species collected in each habitat is noted in the corresponding area. For example, 4 species were collected from all habitats, whereas 9 species were collected only from the Churchill River.

compared with the current taxonomic status and morphological characteristics of species encountered. This combined approach was successful for characterizing this fauna, and resulted in an expanded understanding of the extent of species diversity and distribution, especially among caddisflies.

The barcode- and morphology-based approaches to species recognition were largely concordant. Members of most valid species showed low intraspecific divergences (<2%), much higher interspecific differences, and high bootstrap values for terminal nodes in the Neighbor-Joining tree. The usual discontinuity between maximum intraspecific divergences and distances to the nearest neighboring species indicates the presence of a barcode gap, and supports the use of barcode-based identification systems for future ecological and monitoring studies at Churchill. Moreover, in cases where large intraspecific divergences were observed, morphospecies always formed monophyletic groups so these cases do not represent a problem for a barcode-based identification system built for this site. The DNA barcode results also were instrumental for complementing morphological data, so that rare species were revealed and species pairs with subtle morphological differences were separated.

The balance of this discussion is divided into 3 sections. The 1st considers the new insights gained about the diversity of EPT species at Churchill through DNA barcodes. The 2nd describes how DNA barcoding can aid biodiversity surveys across time, space, and researchers. Last, we discuss how lessons from the present study can improve the efficiency of efforts to build comprehensive barcode libraries for all animal species at Churchill and other sites.

DNA barcoding provides finer taxonomic resolution in biotic surveys

DNA barcoding is crucial in detecting rare taxa and species pairs with subtle diagnosis.—Three closely related Hydropsychidae (Trichoptera) species, Ceratopsyche alhedra (Ross), Ceratopsyche bronta (Ross), and Ceratopsyche vexa (Ross), were each represented in our collections by a single female whose presence was detected during the barcode analysis of numerous specimens of Ceratopsyche alternans because of their distinct COI sequences. Morphological examination by an independent specialist (R. Blahnik, University of Minnesota) had revealed only the presence of C. bronta and C. vexa. The third female of Ceratopsyche remained unidentifiable via morphology, but its COI sequence was very similar to specimens of C. alhedra from sites in Ontario, Minnesota, and Wisconsin (XZ, unpublished data) and was deeply divergent from other congeneric taxa (10.2% sequence divergence from its nearest neighboring species, *Ceratopsyche sparna* (Ross); XZ, unpublished data). Because of the similarity between female hydropsychids and the subtle diagnostic characters (genitalia structures that require dissection and clearing), these locally rare female hydropsychids would have been overlooked in routine morphological sorting. Such oversight probably is a common error in surveys where 1 dominant species co-occurs with several closely related species in much lower abundance. However, this biodiversity heterogeneity can be detected through large-scale DNA barcoding efforts.

DNA barcodes also revealed an unexpected range extension of a caddisfly species by differentiating members of a morphologically similar species pair. A female Agrypnia (BOLD Sample ID: 07PROBE-01729) originally identified as Agrypnia deflata (Milne), was re-examined when it showed 3.45% sequence divergence from its nearest neighboring A. deflata specimen from Churchill. This re-inspection revealed its close affinity to A. obsoleta, a species previously thought restricted to northern and central Europe, Asia, and northwestern North America. Although very similar, the females of these sister species can be differentiated by the shape of segment X. The female in question possesses lighter forewings, more highly developed lateral lobes of tergum X, and a deep terminal notch on the median lobe of tergum X, characters used for recognition of A. obsoleta (Wiggins 1998). In addition, its COI barcode formed a monophyletic lineage with Mongolian A. obsoleta samples, with a within-group divergence of 1.3% and a number of synapomorphic nucleotide variations separating this group from A. deflata (XZ, unpublished data). Because this species pair is very similar in morphology, Wiggins (1998) questioned the validity of certain North American records of A. obsoleta, especially those southeast of the northern Yukon, and concluded that this species was restricted to Beringia. In contrast, the new record for this species at Churchill indicates a much larger expansion from Eurasia. A thorough investigation of the dispersal history of these taxa is beyond the scope of this paper, but more extensive geographic coverage of COI sequences, perhaps together with other gene markers, should provide detailed insights. Moreover, the larvae of North American A. deflata have not been positively associated. The barcode reference library for Trichoptera will assist with the association of larval samples collected from a geographical range of populations and might provide additional larval characters to distinguish A. deflata and A. obsoleta.

Two members of the Oecetis inconspicua (Walker) complex also were differentiated via morphology and DNA barcodes. Considerable variation in the shape of the aedeagus and the inferior appendages on the Xth abdominal segment were observed among the Churchill specimens, but most individuals followed descriptions and illustrations of the species (Ross 1944). This species complex requires revision (Floyd 1995, Ross 1944), so most Churchill O. inconspicua samples were assigned to a provisional identification, O. cf. inconspicua CHU2. However, the COI sequence of 1 specimen (BOLD Sample ID: 06-PROBE-0862) was 12.54% different from its nearest neighboring O. cf. inconspicua CHU2 specimen. Morphological examination revealed that the spine inside its aedeagus was antisymetric to that of typical O. inconspicua, a character that had not been observed previously in this group (J. Morse, Clemson University, personal communication). Because this specimen was otherwise indistinguishable from O. cf. inconspicua CHU2, it probably is an unknown species that would have been overlooked in routine morphological sorting. This single specimen of O. cf. inconspicua CHU1 awaits the collection of additional specimens before it is formally described.

The mayflies Baetis hudsonicus Ide and B. bundyae that we examined were very similar morphologically, with primary differences being the relative length of caudal filaments. Baetis bundyae is widespread in northern and western North America, but B. hudsonicus has a distribution restricted to the far north (Giberson et al. 2007, McCafferty and Randolph 1998). DNA barcode analyses at Churchill provide evidence for maintaining the validity of these sibling species and refute past suggestions of their synonymy (Morihara and McCafferty 1979). Furthermore, B. bundyae was considered as a subspecies of B. macani Lehmkuhl until McCafferty (1994) reinstated it to full species status based on anecdotal evidence from Europe, where the 2 species might co-occur. Further genetic analyses of all eastern Nearctic Baetis species, including B. hudsonicus and B. bundyae, and published European Baetis vernus Curtis-group species, including B. macani (from Ståhls and Savolainen 2008 and Williams et al. 2006), are being conducted by authors of this paper (LMJ, XZ, and PDNH, unpublished data). The preliminary results indicate that our Churchill *B. bundyae* are nested within 1 of the cryptic European *B. macani* haplotype groups that is confined to lotic habitats and possesses narrow gills and invisible tracheae. This haplotype group was proposed by Ståhls and Savolainen (2008) as true B. bundyae distributed in Finland. Thus, our findings at Churchill confirm their results.

Two barcode haplogroups of Acentrella turbida (McDunnough) (noted as A. turbida D and E in our paper) were collected at Churchill, but 5 lineages of this species have been detected thus far from North America (XZ and LMJ, unpublished data). Unlike the adults, the Acentrella larvae associated with these species (LMJ and XZ, unpublished data) differ in body coloration, setation, and leg morphology. However, names cannot be assigned to these species until detailed investigation of related species in Acentrella and other genera is complete (Lugo-Ortiz and McCafferty 1998, McCafferty et al. 2005). Unfortunately, our attempts to recover DNA barcodes from dry, pinned specimens contemporaneous with types in this genus (>50 y old) were not successful, a result suggesting that the recovery of full-length barcodes from mayfly types is not feasible at this time.

Similar observations were made in other taxonomic groups, such as *Oecetis* cf. *ochracea* (Curtis) CHU1 and CHU2, *Plauditus* sp. CHU1 and CHU2, and *Fallceon thermophilos* (McDunnough) complex (see following discussion). In each case, cryptic morphotypes were first detected and differentiated by DNA barcodes and then confirmed by additional morphological scrutiny.

DNA barcoding sheds light on species delimitation in a mayfly complex with highly variable morphology.— Hindwings have been very important to speciesand genus-level identifications of mayfly adults from the family Baetidae in middle North America (Traver 1935, Waltz and Burian 2008). However, recent findings have complicated long-held notions of generic boundaries (McCafferty et al. 2008) because some genera previously identified by hindwing characteristics demonstrate considerable variation that overlaps with other genera.

Some specimens that we examined revealed further complications. A group of small minnow mayfly specimens was very difficult to identify because of high variability in hindwing morphology, in contrast to overall morphological similarity including size and coloration. These specimens are collectively referenced in this paper as the "*Fallceon thermophilos* complex" (Fig. 2) because the presence of a long 3rd vein in the hindwing is characteristic of that species (McCafferty et al. 2008). DNA barcoding separated these difficult specimens into 2 haplotype groups, in which sequence variability was concordant with observed morphological divergence, a result that allowed identification of morphological characteristics that separated members of the 2 clusters.

Specimens in Cluster I lacked sequence diversity (9 individuals with an identical barcode sequence) and

demonstrated low variation in hindwing morphology. Members of this cluster possess a very long, marginal intercalary vein on the hindwing that resembles the longitudinal vein parallel to the 2nd longitudinal vein, but this vein does not extend to the base of the wing. The overall shape of the wing is reminiscent of the genus *Acerpenna*, although it does not have the same undulate upper margin. Thus, this cluster is recognized in this study as *Acerpenna* sp. CHU1.

In contrast, Cluster II is much more diverse in its hindwing morphology, to the extent that individuals might be regarded as members of different genera. These specimens also have what appears to be either a long intercalary vein or a 3rd longitudinal vein. In some specimens, this vein runs parallel to the 2nd longitudinal vein and nearly reaches the basal part of the wing. These specimens were the basis for the initial F. thermophilos identification of the 2 clusters that was made without knowledge of barcode results. Among these specimens, the hindwing vein in question matches narrative descriptions of F. thermophilos but is longer than the vein figured by Traver (1935). In other Cluster II specimens, the vein is shorter and angles slightly towards the 2nd vein. These specimens are generally similar to Cluster I, but the shape of the upper margin of the wing is more convexly undulate. In yet other Cluster II specimens, the 3rd vein is angled toward the 2nd vein and connects to it. When this happens, the 2nd vein is bent slightly toward the upper margin of the wing, and creates a distinctive fork that is consistent with the present concept of the genus Diphetor in North America, which contains a single polytypic species, Diphetor hageni (Eaton) (Meyer and McCafferty 2001). However, the hindwings of our specimens appeared to be slightly more narrow and elongate than in Diphetor. In addition, the barcode sequences of the Churchill samples are very different from D. hageni specimens from Indiana, Florida, and Pennsylvania (with a distance of 19.5% to each group's nearest neighboring member), which themselves form a monophyletic group with deep within-group divergence (with a mean intraspecific divergence of 5.1%; XZ, unpublished data). Because of these taxonomic uncertainties, members of Cluster II have simply been treated as Baetidae sp. CHU1 (Fig. 2).

These observations demonstrate the utility of DNA barcode data as a precursory means of specimen sorting, especially for cases where traditional diagnostic characteristics are highly variable or poorly characterized. Our findings shed light on difficulties associated with proper identification of adults in the family Baetidae and, therefore, bring into question some historical taxonomic work, especially those involving single, few, or only a small series of specimens. We emphasize the need for more extensive study of series of adult specimens of both sexes, ideally associated with larvae.

DNA barcoding suggests potential cryptic species.—In a few cases, distinct COI haplotype clusters were present among individuals assigned to a single species under current species hypotheses. In these cases, no consistent diagnostic morphological characters were found to separate members within each group, but studies have not extended to an examination of larval morphology or habitat selection.

Two caddisfly species, Limnephilus sansoni and Cheumatopsyche campyla, each showed deep intraspecific divergences (maximum 4.01% and 5.25%, respectively), with this variation falling into 2 or 3 distinct COI clusters, respectively. We did not find consistent morphological traits that would differentiate adult members of these distinct COI haplogroups. This result is not surprising because taxonomic ambiguities are known in species of Limnephilus and Cheumatopsyche. Similar intraspecific divergence patterns at barcode loci also were observed in C. campyla collected from a much broader geographic region in eastern North America, where the groupings of COI haplotypes were not correlated to geography (XZ, unpublished data). The high mitochondrial divergences within specimens from single localities, combined with close affinity of the major COI haplotypes across broad geographic areas, suggests that C. campyla might include cryptic species, several of which often occur in sympatry. However, when several closely related Cheumatopsyche species were included in the analysis, the Churchill C. campyla haplotypes seemed to intermix with at least 3 eastern Nearctic Cheumatopsyche species-C. speciosa (Banks), C. ela Denning, and C. pasella Ross (XZ, J. L. Robinson, C. J. Geraci, C. R. Parker, O. S. Flint, D. Etnier, D. Ruiter, REDW, LMJ, and PDNH, unpublished data). Such results suggest that this species complex might have undergone recent speciation and is subject to incomplete lineage sorting and gene introgression, which also has been reported in other caddisfly species (Pauls et al. 2009, Waringer et al. 2007). The taxonomic uncertainty in the C. campyla complex was indeed reflected in our Churchill material, where an independent specialist named the same set of Cheumatopsyche specimens as C. nr. ela. Nevertheless, when considering Churchill material alone, DNA barcoding can precisely assign query samples (e.g., larvae) to the same haplotype groups, despite their taxonomic ambiguity.

No *Limnephilus sansoni* from other regions are currently available to us, but the deep divergence

between the 2 haplogroups (HPGP1 and HPGP2; Fig. 1) at Churchill matches interspecific distances between valid *Limnephilus* species at this locality, e.g., *L. hageni* and *Limnephilus fischeri* Ruiter (see Table 1). Furthermore, both haplogroups of *L. sansoni* are genetically close to some Norwegian specimens of *Limnephilus femoratus* (Zetterstedt) (XZ, unpublished data), a species also known from the Kuril Islands in the Western Pacific (Minakawa et al. 2004). The similarity in COI sequences between Churchill *L. sansoni* and Norwegian *L. femoratus*, and the disjunct records of the latter species, suggest the need for further study to clarify their status. Even though high intraspecific divergence (e.g., bracketed terminals in

Figs 1–3) does not necessarily suggest cryptic speciation, such observations draw attention to the potential need for taxonomic revision.

One *Isoperla* stonefly species could not be identified to species level because the most important diagnostic character, the aedeagus, was not extruded in the field. Thus, members of this cluster of samples were simply treated as *Isoperla* sp. in this paper. One female specimen (07PROBE-02689) had a mean divergence of 2.85% from other members of the cluster and contributed to the large maximum intraspecific divergence within the group (3.13%). This specimen cannot be differentiated from the others by color. Given that diagnosing female *Isoperla* is difficult, the collection of fresh, male specimens with extruded aedeagi is needed to determine the identity of this and other *Isoperla*.

DNA barcoding enables consistency among biodiversity surveys

DNA barcoding has the potential to act as a powerful quality assurance tool for biodiversity surveys by ensuring consistency in taxonomic assignments through time and across space. Taxon concepts change over time (e.g., see Table 1, regarding EPT synonyms in previous studies) and the application of names can vary among specialists (e.g., Cheumatopsyche campyla samples examined in this study) and over time for the same specialist as species concepts change. It is time-consuming and sometimes impractical to incorporate all relevant information about synonymies when interpreting a series of biotic surveys-taxonomic expertise, keys used, and life stages collected vary tremendously across studies. Moreover, barcoding increases the value of biotic surveys conducted by nonspecialists. For example, none of the caddisflies collected at Churchill in an early faunistic study (McClure 1943) was identified even to family level because of the limited taxonomic expertise of the researcher. Such records contribute little to our understanding of the Trichoptera assemblage. By contrast, future studies of Churchill's EPTs will benefit from the present DNA barcode library even if taxonomic assignments are revised over time.

DNA barcode results also can aid the comparison of biodiversity surveys at different locations. In addition to differences in taxonomic concepts across biogeographic regions and a lack of access to comparative or type material, geographical variability in morphology can make it difficult to achieve consistent species-level assignments. DNA barcodes, in combination with morphological and ecological traits, are invaluable for overcoming this problem.

Last, even when species identities remain unknown, barcoding provides a useful interim system for documenting biodiversity. The linking of DNA barcodes to formal species names in biosurveillance is certainly desirable, but is not always possible in the short term if species are undescribed or poorly known. However, DNA barcodes enable comparisons of provisional taxonomic entities collected in various surveys even when specific identification is impossible. For example, several EPT samples (females, larvae, or subimagos) encountered in our study could be identified only to genus or family level because their COI sequences did not match any available reference barcodes. However, the lack of a specific identification did not prevent their detection and registration. The fact that they were readily differentiated by barcodes from related taxa further suggests that these unknowns will gain a species identification as the barcode reference library expands. Linnaean names can be provided to the identifiable specimens acquired in future works and associated with samples collected in the current study via the DNA barcode library. Of course, progress towards achieving a comprehensive DNA barcode reference library of the North American EPT groups also might eventually link currently unknown COI haplotypes to described species.

Critical factors for conducting comprehensive biodiversity surveys: time and space

Several factors might lead to more comprehensive regional biotic surveys. The 2 most critical factors are temporal and habitat coverage. In addition to improvements to our understanding of species diversity through barcoding efforts, the dramatic extension of the Churchill caddisfly and mayfly checklists (and discovery of new species and records for Canada) are obvious benefits of our extensive collecting efforts over time and space for that region.

Our samples covered most of the emergence season for the fauna under investigation. Programmed life cycles characterized by long winter diapause and condensed summer development are one of the typical adaptations of aquatic insects for life in cold climates (reviewed in Danks 2007). Therefore, most adult insects emerge within a short time window in the subarctic. This pattern was reported in the study of biotic communities of the Churchill area (McClure 1943) and was also observed for the caddisflies in our study. The time frame with the highest arthropod diversity in the former study (~July 12) coincided with the peak emergence period in our caddisfly records (July 16-23). However, even within such a short active period, different Churchill caddisfly species showed distinct adult flight patterns. Only 19 of 68 caddisfly species flew for >4 wk. Most species (44) were encountered only over <2 wk, some late in the season (e.g., Anabolia bimaculata (Walker), Asynarchus rossi, Limnephilus externus Hagen, Limnephilus infernalis (Banks), Limnephilus nigriceps (Zetterstedt), Limnephilus sansoni, Rhyacophila angelita Banks, and others). Broad coverage of the emergence season is critical for understanding the biodiversity composition in any biosurveillance study.

Broad habitat coverage is also important for comprehensive species surveys. Strong habitat associations were observed even in Churchill caddisfly adults, whose mobility allowed for travel across habitat types. For example, $\sim \frac{1}{2}$ of the caddisfly species were collected only from 1 of the 3 habitat categories. A clear message from our study is that, as with traditional surveys, sampling in a variety of habitat types should be done in efforts to barcode biota.

Many cold-climate aquatic insects have life histories that extend for ≥ 2 y (Danks 2007), and in such cases, adults might be present in alternate years. Such cases are known for certain Lepidoptera species at Churchill, and our records revealed many caddisflies that were collected in only 1 y. However, a longer sampling program is necessary to test if this pattern was a chance observation or an important characteristic of caddisfly population dynamics in the region.

This contribution has substantially extended the species checklists for Trichoptera and Ephemeroptera at Churchill. However, despite 5 y of collecting, we failed to encounter several species recorded in previous studies, especially for Plecoptera. Moreover species accumulation curves (Zhou et al. 2009) indicate that species richness has not reached an asymptote in any of the 3 groups. This situation is typical of most biodiversity surveys, but rare species, or those with multiyear life cycles, can be incorporated into the reference barcode library over time.

Conclusion

DNA barcoding of the Ephemeroptera, Plecoptera, and Trichoptera at Churchill has proven effective in identifying species, detecting rare taxa, suggesting potential cryptic species, and in documenting α diversity. Our existing knowledge of these 3 groups of relatively well-studied freshwater insects helps interpretation of the barcode data and leads to suggestions for improving biosurveillance techniques and efforts to barcode entire biotas. The value of barcoding EPTs will increase when the DNA library is used to associate all life stages of a species, to improve the comparability of parallel biosurvey results, and to increase the efficiency and coverage of the biological monitoring of freshwater systems.

Acknowledgements

The work was supported by an International Polar Year (IPY) grant from the Natural Sciences and Engineering Research Council of Canada (NSERC) and by grants from Genome Canada through the Ontario Genomics Institute to PDNH. We thank Rob Roughley, Jonathan Witt, Torbjørn Ekrem, and Elisabeth Stur for contributing specimens. The Churchill Northern Studies Centre provided important logistics support. Roger Blahnik, John Morse, and Dave Ruiter helped with the identification and confirmation of some caddisfly species, and W. Patrick McCafferty confirmed preliminary identifications for some mayflies. We also thank staff at the Canadian Centre for DNA Barcoding for their assistance with varied molecular and analytical protocols and Alex Borisenko for suggestions on the accumulation curve analysis.

Literature Cited

- ALEXANDER, L. C., M. DELION, D. J. HAWTHORNE, AND W. O. LAMP. 2009. Mitochondrial lineages and DNA barcoding of closely related species in the mayfly genus *Ephemerella* (Ephemeroptera: Ephemerellidae). Journal of the North American Benthological Society 28:584–595.
- ALLEN, R. K., AND G. F. EDMUNDS. 1961. A revision of the genus *Ephemerella* (Ephemeroptera: Ephemerellidae) III. The subgenus *Attenuatella*. Journal of the Kansas Entomological Society 34:161–173.
- BALL, S. L., P. D. N. HEBERT, S. K. BURIAN, AND J. M. WEBB. 2005. Biological identifications of mayflies (Ephemeroptera) using DNA barcodes. Journal of the North American Benthological Society 24:508–524.
- BASQUIN, P., AND R. ROUGERIE. 2009. Contribution to the knowledge of the genus *Maltagorea* Bouyer, 1993: description of a new species revealed by a combination of morphological characters and DNA barcoding

(Lepidoptera, Saturniinae). Bulletin de la Societe Entomologique de France 114:257–264.

- BEDNARIK, A. F., AND W. P. MCCAFFERTY. 1979. Biosystematic revision of the genus *Stenonema* (Ephemeroptera: Heptageniidae). Canadian Bulletin of Fishery and Aquatic Sciences 201:1–73.
- BOWMAN, M., P. SPENCER, M. DUBE, AND D. WEST. 2009. Regional reference variation provides ecologically meaningful protection criteria for northern world heritage site. Integrated Environmental Assessment and Management 6:12–27.
- DANKS, H. V. 2007. How aquatic insects live in cold climates. Canadian Entomologist 139:443–471.
- DECAËNS, T., AND R. ROUGERIE. 2008. Descriptions of two new species of Hemileucinae (Lepidoptera: Saturniidae) from the region of Muzo in Colombia: evidence from morphology and DNA barcodes. Zootaxa 1944:34–52.
- DENNING, D. G. 1943. The Hydropsychidae of Minnesota (Trichoptera). Entomologica Americana (New Series) 23:101–171.
- DE WAARD, J. R., N. V. IVANOVA, M. HAJIBABAEI, AND P. D. N. HEBERT. 2008. Assembling DNA barcodes: analytical protocols. Pages 275–283 in C. C. Martin (editor). Environmental Genomics, Methods in Molecular Biology. Volume 410. Humana Press, Totowa, New Jersey.
- EDMUNDS, G. F. 1957. *Metretopus borealis* (Eaton) in Canada (Ephemeroptera: Ametropodidae). Canadian Journal of Zoology 35:161–162.
- FISCHER, F. C. J. 1968. Limnephilidae Pars 2. Trichopterorum Catalogus 9. Nederlandsche Entomologische Vereeniging, Amsterdam, The Netherlands.
- FLANNAGAN, P. M., AND J. F. FLANNAGAN. 1982. Present distribution and the post-glacial origin of the Ephemeroptera, Plecoptera and Trichoptera of Manitoba. Manitoba Department of Natural Resources Fisheries Technical Report 82-1:1–79.
- FLINT, O. S. 1960. Taxonomy and biology of Nearctic limnephilid larvae (Trichoptera), with special reference to species in eastern United States. Entomologica Americana 40:1–120.
- FLOYD, M. A. 1995. Larvae of the caddisfly genus *Oecetis* (Trichoptera: Leptoceridae) in North America. Bulletin of the Ohio Biological Survey, New Series 10:1–85.
- FRISON, T. H. 1942. Studies of North American Plecoptera with special reference to the fauna of Illinois. Bulletin of the Illinois Natural History Survey 22:1–355.
- GIBERSON, D. J., S. K. BURIAN, AND M. SHOULDICE. 2007. Life history of the northern mayfly *Baetis bundyae* in Rankin Inlet, Nunavut, Canada, with updates to the list of mayflies of Nunavut. Canadian Entomologist 139: 628–642.
- HAJIBABAEI, M., J. R. DEWAARD, N. V. IVANOVA, S. RATNASING-HAM, R. T. DOOH, S. L. KIRK, P. M. MACKIE, AND P. D. N. HEBERT. 2005. Critical factors for assembling a high volume of DNA barcodes. Philosophical Transactions of the Royal Society of London Series B: Biological Sciences 360:1959–1967.
- HARPER, F., N. H. ANDERSON, AND P. P. HARPER. 1995. Emergence of lotic mayflies (Ephemeroptera) in the

Cascade Range of Oregon. Pages 207–222 *in* L. D. Corkum and J. J. H. Ciborowski (editors). Current directions in research on Ephemeroptera. Canadian Scholars' Press, Inc., Toronto, Ontario.

- HARPER, F., AND P. P. HARPER. 1981. Northern Canadian mayflies (Insecta: Ephemeroptera), records and descriptions. Canadian Journal of Zoology 59:1784–1789.
- HARPER, F., AND P. P. HARPER. 1986. An annotated key to the adult males of the northwestern Nearctic species of *Paraleptophlebia* Lestage (Ephemeroptera: Leptophlebiidae) with descriptions of a new species. Canadian Journal of Zoology 64:1460–1468.
- HAUSMANN, A., P. D. N. HEBERT, A. MITCHELL, R. ROUGERIE, M. SOMMERER, T. EDWARDS, AND C. J. YOUNG. 2009. Revision of the Australian *Oenochroma vinaria* Guenee, 1858 speciescomplex (Lepidoptera: Geometridae, Oenochrominae): DNA barcoding reveals cryptic diversity and assesses status of type specimen without dissection. Zootaxa 2239:1–21.
- HEBERT, P. D. N., J. R. DEWAARD, AND J.-F. LANDRY. 2009. DNA barcodes for 1/1000 of the animal kingdom. Biology Letters. doi:10.1098/rsbl.2009.0848.
- HEBERT, P. D. N., E. H. PENTON, J. M. BURNS, D. H. JANZEN, AND W. HALLWACHS. 2004. Ten species in one: DNA barcoding reveals cryptic species in the neotropical skipper butterfly *Astraptes fulgerator*. Proceedings of the National Academy of Sciences of the United States of America 101:14812–14817.
- HEBERT, P. D. N., S. RATNASINGHAM, AND J. R. DE WAARD. 2003. Barcoding animal life: cytochrome *c* oxidase subunit 1 divergences among closely related species. Proceedings of the Royal Society of London Series B: Biological Sciences 270:S96–S99.
- HOGG, I. D., B. J. SMITH, J. C. BANKS, J. R. DE WAARD, AND P. D. N. HEBERT. 2009. Testing use of mitochondrial COI sequences for the identification and phylogenetic analysis of New Zealand caddisflies (Trichoptera). New Zealand Journal of Marine and Freshwater Research 43:1137–1146.
- HOLZENTHAL, R. W., D. R. ROBERTSON, S. U. PAULS, AND P. K. MENDEZ. 2010. Taxonomy and systematics: contributions to benthology and J-NABS. Journal of the North American Benthological Society 29:147–169.
- HOUGHTON, D. C., AND R. W. HOLZENTHAL. 2003. Updated conservation status of protected Minnesota caddisflies. Great Lakes Entomologist 36:35–40.
- IDE, F. P. 1937. Descriptions of eastern North American species of baetine mayflies with particular reference to the nymphal stages. Canadian Entomologist 69:219–243.
- IDE, F. P. 1954. The nymph of *Rhithrogena impersonata* (Ephemerida) and a new closely related species from the same locality in southern Ontario. Canadian Entomologist 86:348–356.
- IVANOVA, N. V., J. R. DEWAARD, AND P. D. N. HEBERT. 2006. An inexpensive, automation-friendly protocol for recovering high-quality DNA. Molecular Ecology Notes 6: 998–1002.
- JACOBUS, L. M., AND W. P. MCCAFFERTY. 2001. Additions to the Canadian Ephemeroptera. Journal of the New York Entomological Society 109:367–371.

- KIMURA, M. 1980. A simple method for estimating evolutionary rate of base substitutions through comparative studies of nucleotide sequences. Journal of Molecular Evolution 16:111–120.
- LEHMKUHL, D. M., AND C. D. KERST. 1979. Zoogeographical affinities and identification of Central Arctic caddisflies (Trichoptera). Musk-Ox 25:12–28.
- LUGO-ORTIZ, C. R., AND W. P. MCCAFFERTY. 1998. A new North American genus of Baetidae (Ephemeroptera) and key to *Baetis* complex genera. Entomological News 109: 345–353.
- McCAFFERTY, W. P. 1994. Additions and corrections to the Ephemeroptera of Alaska. Proceedings of the Entomological Society of Washington 96:177.
- McCAFFERTY, W. P., M. D. MEYER, R. P. RANDOLPH, AND J. M. WEBB. 2008. Evaluation of mayfly species originally described as *Baetis* Leach (Ephemeroptera: Baetidae) from California. Proceedings of the Entomological Society of Washington 110:577–591.
- McCAFFERTY, W. P., AND R. P. RANDOLPH. 1998. Canada mayflies: a faunistic compendium. Proceedings of the Entomological Society of Ontario 129:47–97.
- McCAFFERTY, W. P., R. D. WALTZ, J. M. WEBB, AND L. M. JACOBUS. 2005. Revision of *Heterocloeon* McDunnough (Ephemeroptera: Baetidae). Journal of Insect Science 35: 1–11.
- McClure, H. E. 1943. Aspection in the biotic communities of the Churchill area, Manitoba. Ecological Monographs 13:1–35.
- McDunnough, J. 1927. A new *Heptagenia* from the Yellowstone region (Ephemeroptera). Canadian Entomologist 59:261.
- MEYER, M. D., AND W. P. MCCAFFERTY. 2001. Hagen's small minnow mayfly (Ephemeroptera: Baetidae) in North America. Entomological News 112:255–263.
- MINAKAWA, N., T. I. AREFINA, T. ITO, T. NOZAKI, N. KUHARA, H. NISHIMOTO, M. UENISHI, V. A. TESLENKO, D. J. BENNETT, R. I. GARA, K. L. KUROWSKI, P. B. H. OBERG, T. I. RITCHIE, AND L. J. WEIS. 2004. Caddisflies (Trichoptera) of the Kuril Archipelago. Bulletin of the Hokkaido University Museum 1:49–80.
- MORIHARA, D. K., AND W. P. McCAFFERTY. 1979. The *Baetis* larvae of North America (Ephemeroptera: Baetidae). Transactions of the American Entomological Society 105:139–221.
- PARSONS, G. L., G. CASSIS, A. R. MOLDENKE, J. D. LATTIN, N. H. ANDERSON, J. C. MILLER, P. HAMMOND, AND T. D. SCHOWALTER. 1991. Invertebrates of the H. J. Andrews Experimental Forest, western Cascade Mountains, Oregon: an annotated list of the insects and other arthropods. PNW-GTR-290. Pacific Northwest Research Station, US Department of Agriculture Forest Service, Portland, Oregon.
- PAULS, S. U., K. THEISSINGER, L. UJVAROSI, M. BÁLINT, AND P. HAASE. 2009. Patterns of population structure in two closely related, partially sympatric caddisflies in Eastern Europe: historic introgression, limited dispersal, and cryptic diversity. Journal of the North American Benthological Society 28:517–536.

- QUINLAN, R., M. S. V. DOUGLAS, AND J. P. SMOL. 2005. Food web changes in arctic ecosystems related to climate warming. Global Change Biology 11:1381–1386.
- RATNASINGHAM, S., AND P. D. N. HEBERT. 2007. BOLD: the Barcode of Life Data System (www.barcodinglife.org). Molecular Ecology Notes 7:355–364.
- RICKER, W. E. 1938. Notes on specimens of American Plecoptera in European collections. Transactions of the Royal Canadian Institute 22:129–156.
- RICKER, W. E. 1944. Some Plecoptera from the far North. Canadian Entomologist 76:174–185.
- RICKER, W. E. 1952. Systematic studies in Plecoptera. Indiana University Publications, Science Series No. 18, 1–200.
- RICKER, W. E. 1964. Distribution of Canadian stoneflies. Gewässer und Abwässer 34/35:50–71.
- ROSENBERG, D. M., AND V. H. RESH. 1993. Freshwater biomonitoring and benthic macroinvertebrates. Chapman and Hall, New York.
- Ross, H. H. 1938. Description of Nearctic caddis flies (Trichoptera) with special reference to the Illinois species. Bulletin of the Illinois Natural History Survey 21:101–183.
- Ross, H. H. 1944. The caddisflies or Trichoptera of Illinois. Bulletin of the Illinois Natural History Survey 23:1–326.
- SAITOU, N., AND M. NEI. 1987. The Neighbor-Joining method: a new method for reconstructing phylogenetic trees. Molecular Biology and Evolution 4:406–425.
- SINGER, G., AND M. HAJIBABAEI. 2009. iBarcode.org: web-based molecular biodiversity analysis. BMC Bioinformatics 10(Suppl 6):S14.
- SMITH, M. A., D. M. WOOD, D. H. JANZEN, W. HALLWACHS, AND P. D. N. HEBERT. 2007. DNA barcodes affirm that 16 species of apparently generalist tropical parasitoid flies (Diptera, Tachinidae) are not all generalists. Proceedings of the National Academy of Sciences of the United States of America 104:4967–4972.
- STÅHLS, G., AND E. SAVOLAINEN. 2008. MtDNA COI barcodes reveal cryptic diversity in the *Baetis vernus* group (Ephemeroptera, Baetidae). Molecular Phylogenetics and Evolution 46:82–87.
- STARK, B. P., AND A. R. GAUFIN. 1976. The Nearctic species of *Acroneuria* (Plecoptera: Perlidae). Journal of the Kansas Entomological Society 49:221–253.
- TAMURA, K., J. DUDLEY, M. NEI, AND S. KUMAR. 2007. MEGA4: molecular evolutionary genetics analysis (MEGA) software, version 4.0. Molecular Biology and Evolution 24: 1596–1599.
- TRAVER, J. R. 1935. Part II: Systematics. Pages 239–739 *in* J. G. Needham, J. R. Traver, and Y. C. Hsu. The biology of mayflies, with a systematic account of North American species. Comstock Publishing Company, Ithaca, New York.
- VAGLIA, T., J. HAXAIRE, I. J. KITCHING, I. MEUSNIER, AND R. ROUGERIE. 2008. Morphology and DNA barcoding reveal three cryptic species within the *Xylophanes neoptolemus* and *loelia* species-groups (Lepidoptera: Sphingidae). Zootaxa 1923:18–36.
- WALTZ, R. D., AND S. K. BURIAN. 2008. Ephemeroptera. Pages 181–236 *in* R. W. Merritt, K. W. Cummins, and M.

B. Berg (editors). An introduction to the aquatic insects of North America. 4th edition. Kendall Hunt, Dubuque, Iowa.

- WARINGER, J., W. GRAF, S. PAULS, AND V. LUBINI. 2007. The Larva of *Drusus nigrescens* Meyer-Dur, 1875 (Trichoptera: Limnephilidae: Drusinae) with notes on its ecology, genetic differentiation and systematic position. Annales de Limnologie – International Journal of Limnology 43:161–166.
- WEBB, J. M., AND W. P. MCCAFFERTY. 2008. Heptageniidae of the world. Part II. Key to the genera. Canadian Journal of Arthropod Identification 7:1–55.
- WIEMERS, M., AND K. FIEDLER. 2007. Does the DNA barcoding gap exist? A case study in blue butterflies (Lepidoptera: Lycaenidae). Frontiers in Zoology 4:8.
- WIGGINS, G. B. 1960. A preliminary study of the North American larvae of the caddisfly family Phryganeidae (Trichoptera). Canadian Journal of Zoology 38:1153– 1170.
- WIGGINS, G. B. 1996. Larvae of the North American Caddisfly Genera (Trichoptera). University of Toronto Press, Toronto, Ontario.
- WIGGINS, G. B. 1998. The caddisfly family Phryganeidae (Trichoptera). University of Toronto Press, Toronto, Ontario.

- WILLIAMS, H. C., S. J. ORMEROD, AND M. W. BRUFORD. 2006. Molecular systematics and phylogeography of the cryptic species complex *Baetis rhodani* (Ephemeroptera, Baetidae). Molecular Phylogenetics and Evolution 40: 370–382.
- WRONA, F. J., T. D. PROWSE, J. D. REIST, J. E. HOBBIE, L. M. J. LEVESQUE, AND W. F. VINCENT. 2006a. Climate change effects on aquatic biota, ecosystem structure and function. Ambio 35:359–369.
- WRONA, F. J., T. D. PROWSE, J. D. REIST, J. E. HOBBIE, L. M. J. LEVESQUE, AND W. F. VINCENT. 2006b. Climate impacts on Arctic freshwater ecosystems and fisheries: background, rationale and approach of the Arctic Climate Impact Assessment (ACIA). Ambio 35:326–329.
- YAMAMOTO, T., AND G. B. WIGGINS. 1964. A comparative study of the North American species in the caddisfly genus *Mystacides* (Trichoptera: Leptoceridae). Canadian Journal of Zoology 42:1105–1126.
- ZHOU, X., S. J. ADAMOWICZ, L. M. JACOBUS, R. E. DEWALT, AND P. D. N. HEBERT. 2009. Towards a comprehensive barcode library for arctic life—Ephemeroptera, Plecoptera, and Trichoptera of Churchill, Manitoba, Canada. Frontiers in Zoology 6:30.

Received: 2 September 2009 Accepted: 29 March 2010