Testing species limits of Eurytomidae (Hymenoptera) associated with galls induced by Diplolepis (Hymenoptera: Cynipidae) in Canada using an integrative approach

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Abstract—Studies of insect host-parasitoid relationships are often confounded by the difficulties associated with species delimitation in taxonomically challenging groups. Eurytomidae (Hymenoptera) are common parasitoids associated with galls induced by Cynipidae (Hymenoptera) and are difficult to identify due to their small size, morphological conservatism, and unreliable published host records. This study tests the species limits of eurytomids associated with galls induced by *Diplolepis* Geoffroy (Hymenoptera: Cynipidae) in Canada using an integrative taxonomy approach including adult morphology, the mitochondrial gene *cytochrome c oxidase I*, host records, and geographical range. Incongruences between morphological and molecular data were found within the *Eurytoma discordans* Bugbee complex, as *Eurytoma discordans, Eurytoma acuta* Bugbee, and *Eurytoma calcarea* Bugbee were shown to be **new synonyms**. The results also revealed the presence of cryptic species within *Eurytoma spongiosa* Bugbee. Furthermore, issues that have impeded ecological and biological studies of eurytomids associated with rose galls such as host specificity and sex association were resolved using DNA barcodes, providing new insights into the evolutionary history of this difficult group.

Résumé—Les études sur les interactions hôtes-parasitoïdes chez les insectes sont souvent limitées par les difficultés associées à l'identification des espèces, surtout lorsqu'elles proviennent de groupes taxonomiques complexes. Les Eurytomidae (Hymenoptera) sont des parasitoïdes communs associés aux galles formées par les Cynipidae (Hymenoptera), et leur identification peut s'avérer laborieuse en raison de leur taille minuscule, leur similitude morphologique et un recueil peu fiable des espèces d'hôtes connues. Le but de cette étude est de redéfinir les limites taxonomiques des eurytomides associés aux galles formées par Diplolepis Geoffroy (Hymenoptera: Cynipidae) au Canada en utilisant une méthodologie taxonomique intégrée basée sur la morphologie des adultes, le gène mitochondrial cvtochrome c oxidase I, les hôtes exploités et l'étendue géographique des espèces. Nous avons noté certaines incompatibilités entre les données morphologiques et moléculaires dans le complexe de Eurytoma discordans Bugbee, trois espèces s'avérant analogues. Nos résultats révèlent également la présence d'espèces cryptiques au sein d'Eurytoma spongiosa Bugbee. Cette étude démontre donc que les difficultés rencontrées lors de recherches écologiques et biologiques sur les eurytomides associés aux galles des rosiers, comme par exemple leur spécifité d'hôtes et l'identification des sexes, peuvent être contournées par l'utilisation de marqueurs génétiques. Cette méthodologie permettra ainsi d'approfondir nos connaissances de l'histoire évolutive de ce groupe taxonomique complexe.

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Introduction

Communities of insects associated with plant galls are useful model systems for studying trophic interactions among insects (Csóka et al. 2005), and species level identification of the inhabitants is necessary to analyse food webs (Gómez et al. 2011). Besides the inducers, gall communities are commonly comprised of parasitoids that feed on inducers or other inhabitants, and inquilines that feed on galls tissues (Shorthouse 2010). One of the driving forces behind the speciation of gall parasitoids is the phenological patterns of their hosts, as parasitoids can only reach their hosts by oviposition through gall tissues, which proliferate and enlarge as the galls develop (reviewed in Csóka et al. 2005). Several studies have also shown that the host plant, host organ, and the time required for gall development all affect richness of parasitoid species (e.g., Schönrogge et al. 1995, 1996; Plantard et al. 1996; Plantard et al. 1999); however, the reason why many parasitoids are highly host specific, despite the presence of other potential hosts on the same host plant, are largely unknown (Csóka et al. 2005). All known parasitoids that attack cynipid hosts are wasps within the superfamilies Chalcidoidea and Ichneumonoidea (Shorthouse 2010).

The superfamily Chalcidoidea contains an estimated 500 000 species, making it one of the most biologically and morphologically diverse groups of parasitic wasps (Gibson et al. 1997, 1999; Munro et al. 2011). While some chalcidoids are phytophagous, the majority are entomophagous and their hosts include all life-history stages of 13 orders of insects, two orders of arachnids, and one family of nematodes (Gibson et al. 1997, 1999). Chalcids within the family Eurytomidae have over 1400 nominal species in 84 genera and are found in most zoogeographical regions (Lotfalizadeh et al. 2007b; Gates 2008; Noyes 2012). Eurytomids are largely endophytic as seed feeders, gall inducers or parasitoids of phytophagous insects (Lotfalizadeh et al. 2007b). The accurate identification of eurytomids, in particular the genus Eurytoma Illiger, which includes more than 700 described species, has proven difficult using existing morphological keys due to overlapping diagnostic characters and lack of illustrations

(e.g., Bugbee 1967). As a result, phylogenetic, ecological, and evolutionary studies of eurytomids have been impeded. The degree of morphological conservatism is particularly prominent in members of the Eurytoma rosae Nees species group, which parasitises various gall-inducing cynipids, Tephritidae (Diptera), and Curculionidae (Coleoptera) (Lotfalizadeh et al. 2007b). Members of the E. rosae group are often morphologically similar and impossible to segregate into morphospecies, despite being ecologically and genetically distinct (Acs et al. 2002; Lotfalizadeh et al. 2007a; Gómez et al. 2011).

A total of 14 native species of cynipid wasps of the genus Diplolepis Geoffroy (Hymenoptera: Cynipidae) have been recorded in Canada, all of which induce structurally distinct galls on Rosa Linnaeus (Rosaceae) (Shorthouse 2010). These rose galls are heavily attacked by chalcid parasitoids, of which eurytomids are the most abundant (Shorthouse et al. 2005; Shorthouse 2010). Ten species of eurytomids are known to be associated with galls of Diplolepis in Canada feeding as koinobiont ectoparasitoids of either the inducers or cynipid inquilines of the genus Periclistus Förster (Noyes 2012). While most eurytomids are univoltine and overwinter within galls before exiting the following spring, cases of fall emergence have been recorded where mature eurytomids pupate and exit the gall in the fall of the year of gall initiation (Shorthouse 1973, 2010; Brooks and Shorthouse 1997). Several studies have been conducted on both adult (Lotfalizadeh et al. 2007a, 2007b) and larval (Gómez et al. 2011) eurytomids associated with rose gall communities in the Palearctic region, whereas the Nearctic species have received little taxonomic attention. Bugbee (1951a, 1951b, 1973) described the majority of the Nearctic species; however, many of the original species descriptions are brief, and were often based on a limited number of specimens collected from a single locality. Further, morphological variations from the type specimens were assigned as subspecies, resulting in even more indistinguishable taxa. While identification keys were provided for Eurytoma by Bugbee (1951b, 1967), they were based only on females and distinguishing characters used were often ambiguous and generally lacking illustrations. These impediments confound studies



Fig. 1. Map of Canada indicating the sampling locations of Eurytomidae used in this study.

on host-parasitoid relationships within galls induced by *Diplolepis* (*e.g.*, Shorthouse *et al.* 2005; Leggo and Shorthouse 2006; Shorthouse 2010), and thus a novel approach is needed to delimit these morphologically similar species.

With advances in molecular biology, the use of molecular markers has proven essential for delimiting closely related species among Hymenoptera parasitoids (Heraty 2009; Santos et al. 2011). The mitochondrial genome in particular serves as a good model for the study of molecular evolution and population genetics, with high rates of evolution and genome reorganisation observed in known chalcid wasp genomes (Dowton and Austin 1995; Oliveira et al. 2008). A short fragment of the mitochondrial cytochrome c oxidase I (COI) gene is the core DNA barcoding animal gene and has proven successful in species identification, and distinct molecular clades or haplogroups have been used in the identification of morphologically cryptic taxa (e.g., Hebert et al. 2004; Smith et al. 2006, 2007). However, controversy exists on the exclusive reliance of mitochondrial DNA in species delimitation without the inclusion of morphological or ecological datasets (e.g., Cognato 2006; Meier et al. 2006). Thus, an integrative taxonomy approach is preferred using multiple independent

character data sources and avoiding reliance on key characters alone in testing species hypotheses for problematic groups (Dayrat 2005; Will *et al.* 2005). Cytochrome *c* oxidase I has been shown to be a valuable tool in identifying cryptic taxa, in combination with morphological and ecological data, for testing host-specificity and geographical variability for Hymenoptera (*e.g.*, Smith *et al.* 2008; Sheffield *et al.* 2009; Ács *et al.* 2010; Kaartinen *et al.* 2010; Sun *et al.* 2011; Gebiola *et al.* 2012) including members of Eurytomidae (Lotfalizadeh *et al.* 2007a; Li *et al.* 2010).

The purpose this study was to use an integrative taxonomy approach to delimit eurytomids associated with galls of 14 species of *Diplolepis* from Canada by testing congruency of genetic variation, morphological differences, host specificity, and geographical distribution between different populations. Additionally the validity of species limits proposed by Bugbee (1967) was tested using COI sequences.

Materials and methods

Sample collection

Maturing or mature galls were collected from various sites in Canada from 1998 to 2011 (Fig. 1), either in the spring after snow melt for galls induced the previous year, or in the fall after galls had matured. Galls from the previous year were stored in jars at room temperature allowing the inhabitants to exit the galls. Galls collected in the fall were subjected to -5 °C for three to four months to break diapause. All inhabitants were either aspirated or removed with a paint brush, and then stored in 100% ethanol. The specimens used for this study were limited to those with sufficient ecological and geographical data to unambiguously identify host galls (n = 423). Eurytomids were selected from pinned specimens and bulk samples stored in 100% ethanol and identified to the species level based on dichotomous morphological keys by Bugbee (1951a, 1967) in combination with host records whenever possible. Specimens that could not be confidently identified were separated into morphospecies. Localities of the eurytomids used in the study are shown in Figure 1. This map was generated using Simplemappr (Shorthouse 2012).

DNA extraction and PCR amplification of COI barcoding region

DNA extractions were performed at the Canadian Centre for DNA Barcoding (CCDB) in Guelph, Ontario, Canada using a silica-based 96-well automated extraction according to the protocol described by Ivanova *et al.* (2006, 2007) in combination with the nondestructive voucher retrieval method described in Porco *et al.* (2010). A series of primers were used listed in Table 1. PCR amplification and sequencing were performed according to the standard protocol used by CCDB (Ivanova and Grainger 2007a, 2007b).

Phylogenetic inference

Contigs were assembled using Sequencher version 4.5 and aligned by CLUSTALX in MEGA version 5.05 (Tamura et al. 2011) then manually checked by eye. Genetic distances were calculated in the Barcode of Life Data System (BOLD) using Kimura-2-Parameter (K2P) (Ratnasingham and Hebert 2007). Sequences of eurytomids with >350 base pairs were used in all analyses, with the sequences of Orthopelma mediator Thunberg (Hymenoptera: Ichneumonidae), Ormvrus rosae Ashmead (Hymenoptera: Ormyridae), and Torvmus *bedeguaris* (Linnaeus) (Hymenoptera: Torymidae) as outgroups.

Maximum likelihood analyses were performed using the K2P distance model (Kimura 1980) in MEGA 5.05 and visualised as a phylogenetic tree. Branch support was assessed with 1000 bootstrap pseudoreplicates and was considered as supported when bootstrap value was >70%. Similarly, Bayesian inference using gamma-distributed rate variation across sites and a proportion of invariable sites with HKY + I + G model, as selected by JModeltest version 0.1.1 (Posada 2008) was performed using MrBayes 3.2 (Ronquist et al. 2012). Two parallel runs of four simultaneous Monte Carlo Markov chains (three heated and one cold) were run for four million generations, and trees sampled every 1000 generations. The burn-in value was set at 25% of the total sampled topologies, with the phylogeny estimated from a majority-rule consensus of the remaining trees at the threshold for clade acceptance set at 0.95. The trace files sequences and specimen

Table 1. Prime	rs used	for	PCR	and	sequencing.
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Primer name	Direction	Primer sequence $(5'-3')$	Primer source	
LepF1	Forward	ATTCAACCAATCATAAAGATATTGG	Hebert et al. (2004)	
FWPTF1	Forward	CCTGGTTCTTTRATTGGTAATGATC	Li et al. (2010)	
RonMWASPdeg_t1	Forward	TGTAAAACGACGGCCAGTGGWTCW	Smith et al. (2008)	
		CCWGATATAKCWTTTCC		
UEA3	Forward	TATAGCATTCCCACGAATAAATAA	Lunt et al. (1996)	
TL2-N-3014	Reverse	TCCATTGCACTAATCTGCCATATTA	Simon et al. (1994)	
C_ANTMR1D-	Reverse	GGRGGRTARAYAGTTCATCCWGTWCC	Modified from	
RonIIdeg_R			Simon et al. (1994)	
C_ANTMR1D-	Reverse	CAWCCWGTWCCKRMNCCWKCAT	Smith et al. (2005)	
AMR1deg_R				
LepR1	Reverse	TAAACTTCTGGATGTCCAAAAATCA	Hebert et al. (2004)	

information are deposited in the project Eurytomidae associated with galls of *Diplolepis* in Canada (project code MZEDO) on BOLD (www.boldsystems.org), and all sequences have been deposited in GenBank under accession numbers KC685087–KC685296.

Morphological study

Morphospecies were compared with the voucher specimens used for molecular study a posteriori, and sorted according to haplogroups. These specimens were chemically dried using Hexamethyldisilazane (Heraty and Hawks 1998) before being point or card mounted. Scanning electronic microscopy or stereomicroscope photographs were taken using methods described by Gates and Pérez-Lachaud (2012). The vouchers were also compared with type specimens located in the National Museum of Natural History (USNM) in Washington, DC, United States of America or the Canadian National Collection of Insects (CNCI) in Ottawa, Ontario, Canada. The DNA extracts are stored at the Biodiversity Institute of Ontario (Guelph, Ontario, Canada), while the specimens are deposited at USNM and CNCI.

Results

Cytochrome *c* oxidase I species delimitation

A variety of primers were used due to the difficulty in amplifying the COI region of chalcids as a result of the poly-T runs in priming region. This is also the likely cause of the low success rate, as COI sequences were obtained only from 220 of 423 specimens. Sequence lengths ranged from 223 base pairs to 632 base pairs, and show a strong A + T nucleotide bias (mean = 0.752) in comparison to C + G (mean = 0.248). Phylogenetic analyses identified eight haplogroups of eurytomids, seven of which have successfully matched identified females with male conspecifics that were morphologically unidentifiable. All haplogroups were well supported by maximum likelihood bootstrap and Bayesian posterior probabilities (Fig. 2). Both the subfamily Eurytominae (Tenuipetiolus + Eurytoma) and the genus Eurytoma were recovered as monophyletic, and Eurytoma iniquus Bugbee, Eurytoma longavena Bugbee, and *Tenuipetiolus ruber* Bugbee were recovered as distinct clades (Table 2). Deeply divergent lineages were revealed in *Eurytoma spongiosa* Bugbee, including an additional clade "*E. spongiosa* 2". *Eurytoma discordans* Bugbee, *Eurytoma acuta* Bugbee, and *Eurytoma calcarea* Bugbee were grouped together into one genetically variable clade (Fig. 3). In addition, two rare haplogroups were found among unidentified species, *Eurytoma* species 1 with four specimens, and a single male specimen as *Eurytoma* species 2 (Fig. 2; Table 2). The intra-specific variation ranged from 0.2% to 3.8%, whereas the inter-specific divergence was 5.7–20.2% (Table 3).

Morphological study

In total, eight morphospecies of eurytomids were found associated with galls induced by Diplolepis from the collection sites, including five of the 10 species previously known from Canada (Noyes 2012). Eurytoma obtusilobae Ashmead was only observed in a single collection of galls of Diplolepis radicum (Osten Sacken) found near Kelowna, British Columbia, Canada in 1999. These specimens failed to generate sequences and were not used for this study. With the exception of T. ruber Bugbee, seven other morphospecies belong to the genus *Eurytoma* within the *E. rosae* species group, characterised by the presence of postgenal depressions and the raised adscrobal carina which forms the precoxal tooth in front of the mesocoxal cavities in lateral view (Lotfalizadeh et al. 2007a).

Key morphological characteristics traditionally used to distinguish eurytomids were found to be ambiguous, with morphological variations often correlated with size, and the absence or atrophy of key characters in the smaller specimens. Other variation included colour of the scape, forelegs and midlegs (yellow to black) and the ratio of wing vein length. New characters such as the number and arrangement of multiporous plate sensilla, and sculpturing on the petiole were useful in distinguishing male specimens, many of which were previously unidentifiable if encountered singly.

Host and geographical records

Ten new provincial records were established as a result of this study (Table 2), including a new Canadian record for *E. iniquus*. In addition, 18 new host associations were discovered, **Fig. 2.** Phylogenetic tree for species of Eurytomidae associated with rose galls induced by *Diplolepis* in Canada based on cytochrome *c* oxidase I (COI) data. Maximum likelihood bootstrap support (first value) and Bayesian posterior probabilities (second value) are shown at each node. The scale bar represents the number of nucleotide substitutions per site. *Orthopelma mediator* (Hymenoptera: Ichneumonidae), *Ormyrus rosae* (Hymenoptera: Ormyridae), and *Torymus bedeguaris* (Hymenoptera: Torymidae) are used as outgroups.



greatly expanding the known host records in North America. With the exception of *Eurytoma* species 2, which has only been observed in association with galls of *Diplolepis ignota* (Osten Sacken), all other haplogroups are associated with two to six different hosts (Table 4). In addition, two generations of *E. longavena* and *E. spongiosa* 2 were collected, from both spring (*e.g., Diplolepis polita* (Ashmead)) and fall initiated galls (*e.g., Diplolepis nebulosa* (Bassett)).

Discussion

Testing species limits using COI

Accelerated rates of evolution of the chalcid mitochondrial genome have been correlated with parasitic lifestyles (Xiao *et al.* 2011); however, testing the species limits of recently diverged lineages is difficult because the organisms often had insufficient time for the evolution of

diagnostic characters or complete reproductive isolation (Xiao *et al.* 2011; Gebiola *et al.* 2012). Independent lines of evidence were used in the testing of species limits in these studies, thus avoiding the reliance of one particular dataset.

The COI sequences found in present study resolved closely related species of eurytomids that are difficult or impossible to distinguish morphologically. This first "screening" of morphospecies using COI drew attention to problematic clades that required further investigation (Li *et al.* 2010). An additional benefit of molecular analyses is the ability to associate sexually dimorphic eurytomids in a simple and precise way, where males were previously unknown or indistinguishable due to the lack of detailed species descriptions. The results of this study show the COI sequences are taxonomically informative in identifying eurytomid species and species boundaries defined by deep COI

Morphospecies	species Haplogroups Locality Host gall		Host plant	Sample size	
Eurytoma acuta	Eurytoma discordans	Alberta	D. nodulosa (Beutenmueller)	<i>R. woodsii</i> Lindley	4
		Alberta	D. tumida (Bassett)	R. woodsii	7
Eurytoma calcarea		Alberta	D. bicolor (Harris)	R. woodsii	10
		British Columbia	D. variabilis (Bassett)	R. woodsii	3
Eurytoma discordans		Ontario	D. nodulosa	<i>R. blanda</i> Gray	4
		Ontario	D. radicum (Osten Sacken)	<i>R. acicularis</i> Lindley	6
		Ontario	D. spinosa (Ashmead)	R. blanda	20
Eurvtoma	Eurvtoma	Alberta	D. bicolor	R. woodsii	2
iniauus	iniauus	British Columbia	D. variabilis	R. woodsii	1
iniquitio		Ontario	D. bicolor	R blanda	4
		Ontario	D. nodulosa	R. blanda	3
		Ontario	D. rosaefolii (Cockerell)	R. acicularis	2
Eurytoma longavena	Eurytoma longavena	Alberta	D. nebulosa (Bassett)	R. woodsii	2
		British Columbia	D. variabilis	R. woodsii	1
		Quebec	D. polita (Ashmead)	R. acicularis	6
		Ontario	D. nebulosa	R. blanda	6
		Ontario	D. <i>fusiformans</i> (Ashmead)	R. acicularis	2
		Ontario	D. polita	R. acicularis	4
		Ontario	D. rosaefolii	R. acicularis	15
Eurytoma spongiosa	Eurytoma spongiosa 1	Alberta	D. ignota (Osten Sacken)	R. woodsii	2
		Alberta	D. tumida	R. woodsii	5
		British Columbia	D. variabilis	R. woodsii	4
		Ontario	D. spinosa	R. blanda	15
		Ontario	D. spinosa	R. rugosa Thunberg	7
		Ontario	D. triforma Shorthouse and Ritchie	<i>R. canina</i> L.	24
	Eurytoma spongiosa 2	Alberta	D. ignota	<i>R. arkansana</i> Porter	5
	r 8	Ontario	D. nebulosa	R. blanda	1
		Ontario	D. triforma	R. acicularis	7
		Ontario	D. triforma	R. canina	19
		Ouebec	D. polita	R. acicularis	4
Eurvtoma	<i>Eurvtoma</i> sn 1	Ontario	D. nodulosa	R. blanda	1
species		Ontario	D. rosaefolii	R. acicularis	3
Species	<i>Eurvtoma</i> sp 2	Alberta	D. ignota	R. arkasana	1
Tenuipetiolus	Tenuipetiolus	Ontario	D. triforma	R. acicularis	2
ruber	ruber	Quebec	D. polita	R. acicularis	2

 Table 2. Collection locality and host information for eurytomid morphospecies and haplogroups.

Note: New records are indicated in bold.

Fig. 3. Expanded phylogenetic tree for the *Eurytoma discordans* complex. Codes after species identification indicates location of host galls on plant organ (LE, leaf; RO, root; ST, stem). The codes after the underscore are the collection location in Canada (AB, Alberta; BC, British Columbia; ON, Ontario). Maximum likelihood bootstrap support (first value) and Bayesian posterior probabilities (second value) are shown at each node.



Interspecific divergence	Eurytoma discordans	Eurytoma iniquus	Eurytoma longavena	<i>Eurytoma</i> sp 1	<i>Eurytoma</i> sp 2	Eurytoma spongiosa 1	Eurytoma spongiosa 2	Tenuipetiolus ruber	Intraspecific divergence	SE
Eurytoma discordans		0.018	0.017	0.016	0.021	0.023	0.014	0.025	0.014	0.005
Eurytoma iniquus	0.092		0.024	0.021	0.023	0.024	0.019	0.028	0.012	0.004
Eurytoma longavena	0.079	0.126		0.018	0.022	0.023	0.018	0.027	0.002	0.002
Eurytoma sp 1	0.076	0.117	0.090		0.021	0.022	0.018	0.028	0.028	0.008
Eurytoma sp 2	0.097	0.132	0.103	0.112		0.023	0.021	0.029	N/A	N/A
Eurytoma spongiosa 1	0.128	0.134	0.118	0.125	0.120		0.021	0.029	0.004	0.002
Eurytoma spongiosa 2	0.057	0.088	0.076	0.089	0.097	0.103		0.026	0.005	0.002
Tenuipetiolus ruber	0.161	0.197	0.175	0.194	0.202	0.198	0.170		0.038	0.011

Table 3. Intra-specific and inter-specific divergence for all haplogroups. Standard errors are shown in reverse of the matrix for interspecific divergence

Table 4.	Eurytomid	haplogroups	and	associated	rose galls.
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Species	Plant organ	Eurytoma discordans	Eurytoma iniquus	Eurytoma longavena	Eurytoma spongiosa 1	Eurytoma spongiosa 2	<i>Eurytoma</i> sp 1	<i>Eurytoma</i> sp 2	Tenuipetiolus ruber
D. bicolor	Leaf	X	Х						
D. ignota	Leaf				Х	Х		Х	
D. nebulosa	Leaf			Х		Х			
D. polita	Leaf			Х		Х			Х
D. rosaefolii	Leaf		Х	Х			Χ		
D. variabilis	Leaf	Х	Х	Х	Х				
D. fusiformans	Stem			Х					
D. nodulosa	Stem	Х	Х				Х		
D. spinosa	Stem	Х			Х				
D. triforma	Stem				Х	Х			Х
D. tumida	Stem	Х			Х				
D. radicum	Root	Х							

divergences are incongruent with morphological studies by Bugbee (1951a, 1951b, 1967, 1973). While this study did not include additional genes, similar studies on eurytomids have shown that mitochondrial and nuclear genes corroborated each other (Lotfalizadeh et al. 2007a; Li et al. 2010), therefore the results of the current study based only on mitochondrial genes is likely robust when combined with ecological and host records. The species limits of three of the eight haplogroups were resolved by COI sequences although the other five haplogroups showed conflicting results with existing morphological data (Bugbee 1951b, 1967). The intraspecific divergence is < 2%, which is consistent with other published studies on Hymenoptera (e.g., Sheffield et al. 2009; Li et al. 2010). Deep phylogenetic divergences within the COI data support the existence of cryptic genetic species in Eurytoma, consistent with previous studies of eurytomids of Palearctic gall communities (Acs et al. 2002; Lotfalizadeh et al. 2007a; Gómez et al. 2011). The presence of cryptic species within specimens identified as E. spongiosa was expected, as the species was originally described as a morphologically variable generalist that attacked a wide variety of hosts (Bugbee 1951a). A possible hypothesis for the lack of consistent morphological differences between E. spongiosa 1 and E. spongiosa 2 despite host differences could be related to the presence of Wolbachia Hertig (Rickettsiaceae), a common and widespread group of intracelluar bacteria found in the reproductive organs of arthropods that can cause cytoplasmic incompatibility (reviewed in Werren et al. 2008). The presence of Wolbachia is much higher within Hymenoptera in comparison to other groups of arthropods, which has been hypothesised to be the cause of host speciation (Bordenstein et al. 2001; Sun et al. 2011; Smith et al. 2012). As the presence of other cryptic chalcid parasitoids has been reported in association with cynipid galls (Lotfalizadeh et al. 2007a; Nicholls et al. 2010), further investigation on screens for Wolbachia may help to delimit the two E. spongiosa species.

Eurytoma acuta, E. calcarea, and *E. discordans* were described as morphologically distinct from each other based on the colour of the scape and shape of the stigmal club and marginal vein

(Bugbee 1951b). Based on specimens examined in this study, these characters were shown to be extremes of a continuum rather than stable characters and are thus unreliable. The three species were also previously distinguished by their range and host, which has been expanded and now overlap as a result of additional data presented in this study. The high rate of intraspecific divergence of this clade suggests the presence of a species complex, where retention of ancestral polymorphism and hybridisation may have resulted in the failure of molecular tracing of species boundaries (Li et al. 2010). Hence, E. acuta and E. calcarea should be synonymised, under the more senior name E. discordans, new synonyms.

Host specificity and the evolution of eurytomids on roses

The new distribution and host records suggest that eurytomids exhibit a much wider host range than previously reported (Noyes 2012), as the majority of species are either oligophagous or polyphagous and were found wherever their hosts occur. Thus, using host records and range as key characteristics (Bugbee 1951b, 1967, 1973) in species delimitation is likely prone to error. The presence of fall emergents in E. longavena and E. spongiosa 2 in both springinitiated and late summer-initiated galls suggests that these species are bivoltine, where the first generation emerges in the spring and attacks freshly initiated galls, while the second generation develops and exits from spring galls in the late summer to attack the galls of other species of Diplolepis that are maturing at this time (Shorthouse 1973).

The radiation of *Diplolepis* species onto novel host plants and organs was likely in response to selection for exclusion of natural enemies (enemy-free space) such as eurytomids (Stille 1984; Price *et al.* 1987; Stone *et al.* 2002). Most eurytomids are found across a wide geographical range within galls found on multiple species of wild roses; thus, their natural range likely mirrors their hosts. For instance, in cases where *Diplolepis spinosa* (Ashmead) shifted hosts from *Rosa blanda* Aiton to the domestic rose *Rosa rugosa* Thunberg (Shorthouse 1988), the species of eurytomid parasitoids that are normally associated with *D. spinosa* are also found attacking galls on the new host plant (Table 2). In a study by Nicholls *et al.* (2010), evidence was provided for parasitoids of oak galls that have tracked their hosts through space and time, showing radiation into cryptic species together with host radiations at multiple trophic levels. It is likely that eurytomids associated with rose galls also have stable, long-term co-evolutionary interactions with other species in the cynipid community, responding as a single unit to environmental perturbations (Nicholls *et al.* 2010).

Several species of eurytomids examined in this study showed a close evolutionary relationship with their hosts, often only attacking hosts inducing galls on a specific plant organ (Table 4). Eurytoma longavena was observed almost exclusively in single-chambered galls such as those induced on leaves (e.g., D. polita). The only exception was galls induced by Diplolepis fusiformans (Ashmead), a small, single-chambered stem gall that is closely related to the other basal lineages of leaf-gall inducing species (Plantard et al. 1998). Likewise E. discordans was found in multi-chambered stem galls where it sometimes consumes several hosts by tunnelling from one larval chamber to another (Brooks and Shorthouse 1997). Tenuipetiolus ruber was rarely found in galls of D. polita and Diplolepis triforma Shorthouse and Ritchie, and the intraspecific divergence of specimens identified as T. ruber (3.8%) suggests the presence of cryptic species. In addition to Diplolepis, this species has also been found in association with cynipid galls on blackberry induced by Diastrophus Hartig (Hymenoptera: Cynipidae) (Bugbee 1951a). Additional specimens from other hosts to determine the species limit of *T. ruber*. The two unidentified species of Eurytoma and E. iniquus were collected from galls with high levels of attack by inquilines of the genus Periclistus Förster (Hymenoptera: Cynipidae) (Table 4). Eurytoma nigricoxa Provancher is the only species in Canada that has been recorded in association with Periclistusmodified galls (Bugbee 1967); however, none of the three species matches E. nigricoxa upon comparison with the holotype. It is likely these three Eurytoma species are parasitoids of Periclistus; although more specimens are needed to further investigate these host relationships.

This study has established a DNA barcode reference library for eurytomids, particularly Eurytoma associated with galls of Diplolepis in Canada. This is the first phylogenetic study of Nearctic Eurytoma and suggests that many eurytomid species associated with rose galls (Bugbee 1951a, 1951b, 1967, 1973) require further investigation. Detailed studies of E. spongiosa 1, E. spongiosa 2, and the E. discordans species complex will undoubtedly aid in the identification of species. In addition, the larval forms of the eurytomids included in this study have not been described, therefore matching larvae with their corresponding adults using COI could provide valuable useful information on species delineation. Such studies of other eurytomid larvae have been morphologically informative when the adults were difficult to identify (Claridge and Askew 1960; Henneicke et al. 1992; Gómez et al. 2011).

The presence of synonymous and cryptic species likely occurs in other eurytomid species treated by Bugbee and are in need of taxonomic revision as many morphological characters used to distinguish Nearctic eurytomids are highly variable. Issues that have impeded the identification of eurytomids associated with cynipid rose galls such as host specificity and sex association were resolved using DNA barcoding, providing new insights into the evolutionary history of this taxonomically difficult group.

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