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ORIGINAL ARTICLE

Phasing in and out of phytophagy: Phylogeny and evolution of the family Eurytomidae (Hymenoptera: Chalcidoidea) based on Ultraconserved Elements

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Abstract

We present the first global molecular phylogenetic hypothesis for the family Eurytomidae, a group of chalcidoid wasps with diverse biology, with a representative sampling (197 ingroups and 11 outgroups) that covers all described subfamilies and 70% of the known genera. Analyses of 962 Ultraconserved Elements (UCEs) with concatenation (IQ-TREE) and multispecies coalescent approaches (ASTRAL) resulted in highly supported topologies in recovering the monophyly of Eurytomidae and its four subfamilies. The taxonomy of Eurytomidae, and in particular the large subfamily Eurytominae, needs major revisions as most large genera are recovered as para- or polyphyletic, and the erection of multiple new genera is required in the future to accommodate these taxa. Here, we synonymize the genera Cathilaria (C. certa, C. globiventris, C. opuntiae and C. rigidae) and the monotypic Aiolomorphus rhopaloides within Tetramesa syn. nov., Parabruchophagus (P. kazakhstanicus, P. nikolskaji, P. rasnitsyni, P. saxatilis and P. tauricus) and Exeurytoma (E. anatolica, E. caraganae and E. kebanensis) within Bruchophagus syn. nov. We also provide 137 DNA barcode COI fragments extracted from the UCE contigs to aid in future identifications of Eurytomidae using this popular genetic marker. Eurytomidae most likely originated in South America with an estimated crown age of 83.37 Ma. Ancestral state reconstruction indicates that secondary phytophagy has evolved at least seven times within the subfamily Eurytominae, showcasing the evolutionary flexibility of these vastly understudied wasps.

KEYWORDS

dating, entomophytophagy, parasitoids, seed chalcid, UCE

INTRODUCTION

The family Eurytomidae encompasses more than 1600 described species distributed across 77 genera, organized into four subfamilies (Gates et al., 2025; UCD Community, 2025). Among these subfamilies, Eurytominae, the largest one, comprises 66 genera and approximately 1500 species (UCD Community, 2025). In contrast, the remaining three subfamilies, Heimbrinae (two genera/eight species) (Stage & Snelling, 1986), Rileyinae (seven genera/78 species) (Gates, 2008) and Buresiinae (two genera/nine species) (Zerova, 1988), are considerably smaller. Eurytomidae are identified by their quadrate pronotum, head and mesosoma with umbilicate sculpture, mesothoracic spiracle not exposed and generally black, brown or yellow coloration. For a more in-depth examination of morphology, refer to Gates et al. (2025). However, the identification of eurytomid genera is often difficult due to the seemingly uniform appearance of the species and the blending

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of character traits that often form morphoclines. The largest genus *Eurytoma* Illiger includes over half of the described eurytomid species, while more than 40 genera are monotypic (Lotfalizadeh et al., 2007). To make matters worse, molecular data are lacking for most eurytomid genera in public databases, making it difficult for non-experts to identify specimens accurately. This is partly due to the low success rate in amplifying the typical barcoding fragment using universal primers (e.g., Li et al., 2010; Zhang et al., 2014), although recently, primer sets with higher success rates have been designed specifically for eurytomids (Jafari et al., 2023; Zhang et al., 2022).

Eurytomids are present on every continent except Antarctica, with the highest diversity observed in tropical regions, followed by temperate climates. They showcase a wide range of biological characteristics and exhibit rapid evolution in both host utilization and feeding behaviour. The majority of eurytomid larvae are endophytic. serving as seed eaters, gall formers or parasitoids of phytophagous insects (Gates et al., 2025). Most species act as primary or secondary parasitoids, targeting eggs, larvae or pupae of various arthropod groups such as Diptera, Coleoptera, Hymenoptera, Lepidoptera, Orthoptera and Araneae. Host-parasitoid relationships for many endophytic species remain unclear, and some Eurytomidae believed to be parasitoids may function as inguilines (Lotfalizadeh et al., 2007). A subset of eurytomid species holds economic significance, either as potential biological control agents or pests. Due to their diverse biology, Eurytomidae play contrasting roles in agriculture. Many are harmful seed eaters affecting cultivated plants, while others are used for the biological control of invasive plants (Gates et al., 2025).

The monophyly of Eurytomidae and the relationships between the subfamilies remain debated as different morphological and molecular data have provided conflicting answers (Gates, 2008; Lotfalizadeh et al., 2007). The phylogenetic analysis of Gates (2008) used 50 morphological characters with a sampling focus on Rileyinae, recovering a monophyletic family. Lotfalizadeh et al. (2007) used 150 morphological characters and focused on the large subfamily Eurytominae, instead recovering a paraphyletic Eurytomidae, with Heimbrinae as the sister to Chalcididae, another family of chalcid wasps used as outgroup, instead of grouping with the rest of the family. Using multilocus data, Campbell et al. (2000) included five eurytomid samples within their Chalcidoidea phylogeny inferred using the 28S-D2 rDNA and recovered a polyphyletic Eurytomidae comprising two disparate groups, Rileyinae and Eurytominae. Chen et al. (2004) published the first molecular phylogeny of the family using 24 species and four genes: 16S,18S, and 28S (ribosomal DNA) along with COI (mitochondrial DNA). While most genes exhibited very low levels of variability and the ribosomal DNA results strongly conflicted, the authors concluded that the family was not monophyletic, with Rileyinae sister to Dirhininae (Chalcididae) and nested within a clade containing other chalcidoid families such as Eunotidae, Perilampidae and Eucharitidae. Munro et al. (2011) also failed to recover a monophyletic Eurytomidae in their molecular analysis of Chalcidoidea using 18S and 28S, as while Rileyinae was recovered as a monophyletic group, Buresiinae and Heimbrinae did not cluster with the other Eurytomidae. Heraty et al. (2013) recovered a monophyletic Eurytomidae using a combination of

morphology with the same two ribosomal genes, with all subfamilies monophyletic except for Heimbrinae which placed internally with *Neorileya* Ashmead (Rileyinae). Using transcriptome data, Peters et al. (2018) recovered a monophyletic Eurytomidae as sister to Chalcididae using transcriptomic data and a reduced representation matrix, but their sampling included only two specimens/genera of Eurytominae.

With the rapid advances in sequencing technology, it is now possible to generate thousands of loci from non-destructively sampled museum specimens (Cruaud et al., 2019), enabling the increasing popularity of museum-based phylogenomic studies dubbed 'museomics'. Target capture approaches such as Ultraconserved Elements (UCEs, Faircloth et al., 2012) have revolutionized our understanding of arthropod phylogenomics, including studies on Hymenoptera and/or Chalcidoidea that have included Eurytomidae (e.g., Blaimer et al., 2023; Branstetter, Longino, et al., 2017; Cruaud et al., 2021, 2024; Zhang et al., 2020, 2022). However, the monophyly of Eurytomidae remained unresolved in these studies until Cruaud et al. (2024) recovered a monophyletic Eurytomidae (18 taxa representing all four subfamilies) with Chalcididae as sister group. Conversely, the Blaimer et al. (2023) study used many of the same samples as Cruaud et al. (2024) but did not recover Eurytomidae as monophyletic despite a wider sampling of 35 taxa; Heimbra opaca (Ashmead) behaved as a rogue taxon in this study by grouping either with the colotrechnine pteromalid Colotrechnus Thomson or within Chalcididae instead of within the otherwise monophyletic Eurytomidae.

The goal of the current study is to use UCEs and an expanded global taxonomic sampling in order to (1) test the monophyly of Eurytomidae, the four subfamilies, and major genera; (2) infer divergence times of major groups using molecular dating analyses; and (3) investigate how phytophagy and entomophagy have evolved over the history of the family.

MATERIALS AND METHODS

Taxonomic sampling

The dataset comprises 208 individuals with 197 ingroups chosen to cover the taxonomic breadth within Eurytomidae. UCE data were newly generated for 151 ingroups, while data for 57 specimens were acquired from previously published studies, including 10 Chalcididae and one Chalcedectidae as outgroups (Blaimer et al., 2023; Branstetter, Danforth, et al., 2017; Cruaud et al., 2021, 2024; Zhang et al., 2022). The ingroup taxa represent 48 genera, as well as a number of specimens that could not be placed to genus within the current classification (Table S1, File S1). The identification of the specimens was confirmed by authors considered to be the world authorities on Eurytomidae (GD and MWG). Twenty-one known genera are missing from our sampling, but these genera constitute only 49 species among the >1600 total described species from the family (File S1). In many cases, these missing genera are monotypic and known only from their type specimens. A few other missing genera are doubtful according to original descriptions. Nevertheless, the some regions are

under-sampled, especially the tropical realms (Neotropical, Afrotropical and Indomalayan regions) mostly because of the lack of suitable material for the sequencing.

Generation of the UCE dataset

The UCE library preparation (N = 151) was conducted in the Laboratories of Analytical Biology at the Smithsonian Institution's National Museum of Natural History (NMNH, Washington DC, USA) and the Centre de Biologie pour la Gestion des Populations (Biology Centre for Population Management, CBGP, Montpellier, France). The protocol largely follows the standard pipeline for capturing and enriching UCE loci from Hymenoptera (Branstetter, Longino, et al., 2017; Cruaud et al., 2019; Zhang et al., 2019). Briefly, DNA was extracted using the DNeasy Blood and Tissue Kit, with individuals either destructively processed during extraction or left intact. The Kapa Hyper Prep library preparation kit (Kapa Biosystems, Wilmington, MA) was used along with TruSeg universal adapter stubs and 8-bp dual indexes (Glenn et al., 2019), combined with sheared genomic DNA and amplified using polymerase chain reaction (PCR). We followed the myBaits V4 or V5 protocol (Arbor Biosciences, Ann Arbor, MI) for target enrichment of the pooled DNA libraries (4-12 samples per pool) using either the Hymenoptera 1.5Kv1 (Faircloth et al., 2015) or the 2.5Kv2P kit (Branstetter, Longino, et al., 2017), with probe/target hybridization at 65°C for 24 h. The combined library was sequenced on an Illumina HiSeq 2500 (150-bp paired-end) at Huntsman Cancer Institute (Salt Lake City, UT, USA) or Admera Health (South Plainfield, NJ, USA), or on an Illumina MiSeq (300-bp paired-end) at UMR AGAP (Montpellier, France). Voucher specimens are stored either at NMNH, CBGP, or Taiwan Agricultural Research Institute (TARI, Taichung, Taiwan).

UCE processing and matrix assembly

All downstream analyses were conducted on the Smithsonian High Performance Cluster (SI/HPC, https://doi.org/10.25572/SIHPC). We used the Phyluce v 1.7.4 pipeline (Faircloth, 2015) to process newly generated raw reads and published genome/UCE assemblies mined from previous studies. Adapters were trimmed using illumiprocessor or trimmomatic (Bolger et al., 2014; Faircloth, 2013) and assembled using spades v 3.14.0 (Prjibelski et al., 2020). The assemblies were aligned using MAFFT v 7.490 (Katoh et al., 2002; Katoh & Toh, 2008) and trimmed using GBlocks (Castresana, 2000) with the following settings: b1 = 0.5, b2 = 0.5, b3 = 12, b4 = 7. Additionally, we used Spruceup v 2024.7.22 with a 99% Weibull distribution or manual cutoff of select samples to remove any potentially misaligned regions as they can produce exaggerated branch lengths (Borowiec, 2019a). The manual cutoffs, samples and their values are: (Aximopsis_sp1_USN-MENT01322402, 0.4; Aximopsis_sp2_USNMENT01322403, 0.4; Bruchodape_sp_USNMENT01322364, 0.4; Bruchophagus_abscedus_USN MENT01525852, 0.35; Bruchophagus_gibbus_USNMENT01322388,

0.25; Bruchophagus_phlei_USNMENT01322376, 0.4; Bruchophagus_ roddi_USNMENT01322386, 0.3; Burksoma_scimitar_USNMENT013 22361, 0.5; Eudoxinna_sp_USNMENT01322401, 0.4; Eurytoma_erythrinae_USNMENT01938343, 0.4; Eurytoma_obtusiventris_USN-MENT01322387, 0.4; Heimbra_opaca_USNMENT01339599, 0.6; New_Genus_USNMENT01322400, 0.5; Philolema_latrodecti_USN-MENT01322374, 0.4; and Phylloxeroxenus_sp3_USNMENT01322411, 0.5). Additionally, DNA barcode sequences were bioinformatically extracted from UCE contigs using phyluce_assembly_match_ contigs to barcodes.

Phylogenetic analyses

We selected the 50% complete matrix (1605 loci) and 70% complete matrix (962 loci) which represent loci that are present in >50% and ≥70% of the taxa, respectively. The 70% matrix is our preferred dataset and is used for other downstream analyses, and while the 50% consisted of a larger number of genes, it also had higher levels of missing data (for results of all 50% analyses see Figures S5-S8). Phylogenetic summary statistics were calculated using AMAS v 0.98 (Borowiec, 2016). We conducted phylogenetic analyses under the maximum-likelihood criterion with IQ-TREE v 2.3.1 with the default 100 iterations (Minh, Schmidt, et al., 2020). The best model for each locus was selected using ModelFinder v2.0 (Kalyaanamoorthy et al., 2017) with the IQ-TREE command '-MFP + MERGE'. To assess support, we performed 1000 Ultrafast Bootstrap (UFBoot, Hoang et al., 2018), along with '-bnni' to reduce the risk of overestimation, and a Shimodaira-Hasegawa approximate likelihood-rate test (SHaLRT, Guindon et al., 2010) with 1000 replicates. Only nodes with support values of UFB ≥ 95 and SH-aLRT ≥ 80 were considered robust. We also inferred gene and site concordance factors (gCF and sCF) following Minh, Hahn, and Lanfear (2020) resulting in the trees 50p_CONCORD and 70p_CONCORD.

In order to minimize the effects of phylogenetic heterogeneity, we also filtered the matrices with the symmetry test of stationarity, reversibility and homogeneity for sequence alignments (Naser-Khdour et al., 2019) and conducted separate analyses using the same settings as above, resulting in the trees 50p_SYM and 70p_SYM. The matrices were also analysed using partitions based on Sliding-Window Site Characteristics of Site Entropy (SWSC-EN, Tagliacollo & Lanfear, 2018) and partitioned using the rcluster algorithm in PartitionFinder2 via RAxML using default settings (Lanfear et al., 2014, 2016; Stamatakis, 2006) resulting in the trees 50p_SWSC and 70p_SWSC. In order to ensure the global optimum is reached while also limiting the computational time, we conducted five separate runs of the 70p_SWSC analysis in IQ-TREE to ensure the log likelihood, Robinson-Foulds (RF) distance and topology are comparable among runs (Liu et al., 2024). We also extracted the protein-coding genes using the custom scripts provided by Borowiec (2019b) using the protein FASTA file from assembled genomes of Nasonia vitripennis (Walker) as reference. We analysed the resulting concatenated matrix of amino acids using the posterior mean site frequency (PMSF)

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method in order to ameliorate long-branch attraction artefacts (Wang et al., 2018). The 50p_SWSC and 70p_SWSC were used as guide trees with 60-component mixture models (Wang et al., 2018), resulting in trees 50p_PMSF_C60 and 70p_PMSF_C60, respectively.

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Species trees under the multispecies coalescent (MSC) model were inferred using ASTRAL-III v5.7.8 (Zhang et al., 2018) for both matrices. Gene trees of each locus were estimated in IQ-TREE, and the best models of nucleotide substitution were selected using ModelFinder. We collapsed all branches in the gene trees with <10% UFB using the 'nw_ed' command included in Newick Utilities (Junier & Zdobnov, 2010), as this step has been shown to improve accuracy in MSC analysis (Zhang et al., 2018). Support was assessed by annotating the MSC tree with local posterior probabilities (LPPs, Sayyari & Mirarab, 2016), with ≥0.95 considered as strong support, resulting in trees 50p_BS10_ASTRAL 70p_BS10_ASTRAL.

Resulting raw tree outputs were modified using R packages ggtree (Yu et al., 2017), treeio (Wang et al., 2020) and phytools (Revell, 2024), along with Inkscape v1.4.

Divergence dating

We estimated the divergence times using MCMCTree using PAML v4.9j (Yang, 2007), using approximate likelihood calculation and ML estimation of branch lengths and the 70p SWSC tree as the fixed input along with the 70% matrix. Due to the lack of reliable published eurytomid fossil records, we instead used secondary calibration based on divergence dates from previous studies (Blaimer et al., 2023; Cruaud et al., 2024). The soft minimum bound for the common ancestor of Eurytomidae + Chalcididae was set at 92 Ma, with Chalcididae and Eurytomidae set at 73-86 Ma and 74-84 Ma, respectively (Blaimer et al., 2023; Cruaud et al., 2024). Uniform distributions were used as calibration densities. We conducted the analysis with a 50,000-generation burn-in and 100,000 additional generations using independent rates under the HKY model to strike a balance between complexity and computational efficiency, and convergence was assessed using Tracer v1.7.1 (Rambaut et al., 2018) to ensure ESS values were >200 for all categories. The resulting tree was visualised using MCMCtreeR (Puttick, 2019) in R v4.3.3 (R Core Team, 2024).

Historical biogeography

Historical biogeography was estimated using BioGeoBEARS v1.1.1 (Matzke, 2014, 2018) with the MCMCTree result as input. Coding of occurrences was based on collecting labels and expert knowledge and scored as present/absent in six biogeographic regions (Neotropical, Nearctic, Afrotropical, Palearctic, Indomalayan and Australasian) (Table S3). Dispersal-Extinction-Cladogenesis (DEC; Ree & Smith, 2008), BAYAREALIKE and DIVALIKE (Ronquist, 1997) models were used with and without the jump parameter for founder events (+J; Matzke, 2014), and maximum range size allowed was set to five.

Model selection was performed based on Akaike Information Criterion (AIC_c) (Matzke, 2022). We included constraints on three different time periods (0–23 Ma/Neogene + Quaternary; 66–23 Ma/Paleogene; and 100–66 Ma/Late Cretaceous) with different dispersal rate scalers following Cruaud et al. (2024) (Table S5a).

Ancestral state reconstruction of feeding behaviour

We reconstructed the ancestral states of feeding behaviour using the R package CorHMM v2.8 (Beaulieu et al., 2013), using the 70p_SWSC phylogeny as input after pruning the outgroups. We coded '0' for entomophagous, '1' for phytophagous, and '?' for unknown (Table S3). The host data was based on a combination of sample information, literature and unpublished data (Table S3). We performed reconstructions under the 'equal rates' model (ER), 'symmetrical rate' (SYM) and 'all rates different' model (ARD) and compared the fit of these models with the corrected AICc.

RESULTS AND DISCUSSION

Phylogenetic relationships of Eurytomidae

Our study is the most comprehensively sampled phylogenomic study of Eurytomidae to date, highlighting the utility of museomics on poorly studied but diverse lineages of small parasitic wasps. The number of UCE loci recovered ranged from 228 to 2145 (Table S1). The 50% complete UCE matrix consisted of 565,920 bp of data, of which 414,715 (73.3%) are variable sites, and 324,331 (57.3%) are parsimony informative. The 70% complete UCE matrix consisted of 340,005 bp of data, of which 255,163 (75%) are variable sites, and 204,108 (60%) are parsimony informative. The amount of missing data for the 50% and 70% matrices was 38.82% and 32.04%, respectively (Table S2). We also managed to extract 137 COI fragments >200 bp from the UCE assemblies, and while these fragments cannot be used to accurately infer family-level relationships, they can still improve the publicly available sequences of Eurytomidae for future identifications.

The monophyly of Eurytomidae and its four subfamilies was recovered in all of our analyses (Figures 1 and S1–S8), which is consistent with Cruaud et al. (2024) and somewhat consistent with previous morphological studies (Gates, 2008; Lotfalizadeh et al., 2007). The topologies of the 50% and 70% matrices are mostly identical among the different analyses, with slight shifts among the relationships among genera within the *Chryseida* clade of eurytomines (Figure 1, Figures S1 and S5). Larger differences were observed between the protein-coding-only analyses (Figures S3 and S7) and between the MSC analyses using ASTRAL (Figures S4 and S8). Here we summarize our main findings based on the preferred tree of the 70% matrix partitioned by SWSC (70p_SWSC), as it limits the impact of missing data and model violations while also highlighting major differences among analyses (for full comparisons see Figures S1–S8). The five separate analyses of the 70p_SWSC had log likelihoods ranging



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from -8137442.20 to -8138023.72, and with zero RF distance among them and no topological differences:

Rileyinae is recovered to be the sister to all other eurytomids. Among the two genera within this subfamily represented by more than one species, *Neorileya* was monophyletic while *Rileya* Ashmead was not.

Buresiinae was recovered as the sister group to Heimbrinae and Eurytominae, and within this subfamily *Macrorileya* Ashmead was recovered as a monophyletic group in the ML analyses and sister to *Buresium* Bouček, which was represented by a single specimen. *Macrorileya_sp_USNMENT01558252* is recovered as the sister group to the rest of eurytomines in the MSC analysis with strong support of LPP = 1, as opposed to being grouped with the rest of *Macrorileya*.

Heimbrinae was only represented by *H. opaca* in our study, but it was recovered as the sister to Eurytominae in all analyses except in the 70p_PMSF_C60 tree, where it was recovered as the sister to Rileyinae. We used the same sample of *Heimbra_opaca_USN-MENT01139599* as in Blaimer et al. (2023) as the sole representative for Heimbrinae. While it was recovered outside of Eurytomidae in that earlier study, it was placed as the sister to the subfamily Eurytominae in all of our analyses, similar to Cruaud et al. (2024). The difference in topology with the Blaimer et al. (2023) tree could be the result of filtering of misaligned regions or better taxonomic sampling.

The subfamily Eurytominae is the largest of the four subfamilies and represents the bulk of our taxonomic sampling. Unsurprisingly, most of the larger genera where we included multiple species were not recovered as monophyletic: Aximopsis Ashmead, Bruchophagus Ashmead, Chryseida Spinola, Eurytoma, Gibsonoma Narendran, Isosomodes Ashmead, Prodecatoma Ashmead, Phylloxeroxenus Ashmead and Tetramesa Walker. Bephratelloides Girault, Bephratoides Brues, Bruchodape Burks, Exeurytoma Burks, Paradecatoma Masi, Philolema sensu largo Cameron, Sycophila Walker and Systole Walker were monophyletic. The relationships among genera within the large subfamily Eurytominae are somewhat consistent with the morphological study by Lotfalizadeh et al. (2007); we provide more detailed taxonomic discussion in the Supplemental File. Our study highlights the need to revise the generic concepts for Eurytominae, as many genera are para- or polyphyletic. While this is a well-known problem within the subfamily, the high level of morphological homoplasy and interpretation of difficult morphological characters made it extremely challenging to accomplish (Lotfalizadeh et al., 2007). With the aid of the UCE phylogeny as a backbone, refinement of Eurytominae taxonomy can take place on a smaller scale with more extensive taxonomic sampling of certain clades of interest (Zhang et al., 2021). The establishment of multiple new genera is needed to accommodate outliers that do not fit the current generic limits (e.g., Campos-Moreno et al., 2022; Gates & Cascante-Marin, 2004; Gates & Delvare, 2008), preferably in combination with UCE and/or multilocus data such as COI.

The base of the Eurytominae consists of a series of phytophagous genera: (1) The genus *Ausystole* Bouček is sister to all other Eurytominae, (2) followed by the genus *Aranedra*, (3) then the *Proseurytoma* clade consisting of *Austrodecatoma* Girault, *Risbecoma* Subba Rao, *Proseurytoma* Kieffer and an undescribed genus feeding on Myrtaceae.

The next clade consists of the genus Foutsia Burks, then a paraphyletic assemblage of a clade that includes Phylloxeroxenus, Prodecatoma and Tenuipetiolus Bugbee. This is followed by the large Chryseida clade that consists of a mix of some 'Aximopsis' (including the newly described microgastrine braconid hyperparasitoid 'Aximopsis' gabrielae Zhang, Gates & Campos) and 'Eurytoma' species (parasitoid 'Eurytoma' obtusiventris Gahan, phytophagous 'Eurytoma' werauhia Gates and New Genus Guadeloupe USNMENT01322395), then Eudoxinna Walker and an undescribed genus assigned to the 'erythroapsis group' New Genus USNMENT01322400. This is followed by ((Bephrata Cameron + Isosomodes) + Khamul Gates), which are sister to (Bephratoides + Bruchodape), and all of which are sister to a clade with Axima Walker, the remaining Aximopsis, Chryseida and the monotypic Burksoma scimitar Subba Rao. In the PMSF analysis, Bephratoides sp2 USNMENT01339598 was instead recovered as the sister to Eudoxinna, while Khamul was an isolated branch.

'Eurytoma' timaspidis (Mayr) (aspila species group) and Eurytoma_sp14_USNMENT01525759 (verticillata species group) were recovered as isolated branches and are broadly separated from the bulk of Eurytoma including its type species Eurytoma abrotani (Panzer). This is followed by the Sycophila clade, which consists of ('Bruchophagus' feeding on Citrus + Mangoma Subba Rao) + ('Eurytoma' in the salicis and pistaciae species group + Gibsonoma), which issister to (Sycophila + Ficomila Bouček).

The monotypic Syceurytoma ficus Bouček is recovered as an isolated lineage, followed by the Systole clade which consists of the phytophagous Systole, a monophyletic genus. Its sister group Tetramesa is largely monophyletic with the exception of Tetramesa phragmitis (Erdös), which is sister to all of the other Tetramesa plus the monotypic Aiolomorphus rhopaloides Walker and Cathilaria opuntiae (Muesebeck). As all these genera gall twigs or flowers of Poaceae, the four described Cathilaria species (C. certa Zerova, C. globiventris (Zerova), C. opuntiae and C. rigidae Zerova) and A. rhopaloides are synonymized within Tetramesa syn. nov., which is consistent with their biology and no longer renders Tetramesa paraphyletic.

The Plutarchia clade includes the genus Plutarchia Girault, unplaced Eurytominae_Taiwan_MWG189, Prodecatomidea Risbec and two species of Paradecatoma, followed by the clade/genus Philolema s.l.

The 'Eurytoma' dentata clade consists of an unnamed genus reared from Magnolia L., Eurytoma erythrinae Gates & Delvare, Eurytoma laserpitii Mayr, Eurytoma dentata Mayr and unplaced Eurytominae_Taiwan_MWG187.

FIGURE 1 Phylogram of Eurytomidae based on the 70% UCE matrix, inferred under the SWSC-EN partitioning, partition merging and ModelFinder model selection, generated using IQ-TREE (70p_SWSC). Black dots at nodes indicate support values of UFB \geq 95 and SH-aLRT \geq 80, values below that are omitted. Scale bar in substitutions per site. Habitus photos from top to bottom: *Rileya pallidipes* (Michael Gates), *Macrorileya* sp. (Michael Gates), *Heimbra opaca* (Craig Brabant) and *Eurytoma abrotani* (Gérard Delvare).

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The small Endobia clade (Fronsoma Narendran + Endobia Erdös) is sister to the Bruchophagus clade, which consists of the rest of Bruchophagus, unplaced Eurytominae_Tanzania_JRAS08586_0101 and Exeurytoma. In the 70p_PMSF_C60 tree, Bruchophagus roddi (Gussakovskiy) was recovered as an isolated lineage outside of these two clades, whereas in the 70p_PMSF_C60 tree, Fronsoma was recovered in this position. The Bruchophagus bajarii, gibbus, phlei and squamea species groups defined by Lotfalizadeh et al. (2007), along with the former genus Nikanoria (Nikol'skaya) were recovered within Bruchophagus. Additionally, a set of Bruchophagus species reared from seeds of Asphodelaceae sharing the same characteristics as the genus Parabruchophagus Zerova (B. lecomtei Delvare, B. gijswijti Askew & Ribes, B. abscedus Askew, B. insulare Delvare, B. asphodelinae Askew & Stojanova) is recovered within the genus. Therefore, all five described species included in Parabruchophagus (P. kazakhstanicus Zerova, P. nikolskaji Zerova, P. rasnitsyni Zerova, P. saxatilis Zerova and P. tauricus Zerova), which also develop in the seeds of Asphodelaceae (Eremurus M.Bieb.), are transferred to Bruchophagus syn. nov. Finally, Exeurytoma, which is represented by E. anatolica Cam and an undescribed species Exeurvtoma sp2 USNMENT01525808, is recovered as the sister to the Bruchophagus gibbus species group; all three described Exeurytoma species (E. anatolica, E. caraganae Burks and E. kebanensis Doğanlar) are therefore synonymized within Bruchophagus syn. nov. as both groups develop within leguminous pods.

The final grouping consists of the clade/genus Bephratelloides Girault, followed by Axanthosoma clade (unplaced Eurytominae_-Taiwan_MWG191 and Axanthosoma Girault), the newly described Kavayva Zhang, Silvestre & Gates and finally the rest of Eurytoma sensu stricto (including unplaced Eurytominae_Taiwan_MWG191) + the monotypic Masneroma angulifera Bouček. In the 70p_PMSF_C60 tree, *Eurytoma_sp11_USNMENT01525819* was instead recovered outside of the Kavayva/Eurytoma s. s./Masneroma clade. The following Eurytoma s.s. species groups from Lotfalizadeh et al. (2007) were recovered within our analysis: robusta, compressa, amygdali, morio, fumipennis, rosae/abrotani + appendigaster groups.

Divergence dating and historical biogeography

Based on our divergence dating analysis (Figures 2 and S9, Table S4), the crown age of Eurytomidae is \sim 83.3 Ma [95% equal tail credibility interval (CI) 81.1–84.9 Ma], with Rileyinae estimated at \sim 71.8 Ma (64.9–80.0 Ma); followed by Buresiinae at \sim 66.6 Ma (55.6–75.6 Ma). The subfamilies Eurytominae and Heimbrinae diverged at \sim 75.5 Ma (71.5–80.2 Ma), while crown Eurytominae diverged at \sim 70.9 Ma (66.6–75.8 Ma).

DIVALIKE+J was selected as the best model, with a weighted AICc of 0.48 (Table S5b). The historical biogeographic analyses

recovered a South American origin for Eurytomidae (Figures 2 and S10). This result aligns well with the paleogeographic configurations during the Late Cretaceous and early Cenozoic, where the three continents were still relatively connected through intermittent land bridges or close continental proximities. Rileyinae, Heimbrinae and Eurytominae are proposed as having a South American origin, while Buresiinae was recovered with an Afrotropical origin. A major divide within Eurytominae occurred approximately 66 Ma around the K-Pg mass extinction, splitting the group into mostly Old World and New World clades. The timing of this divergence coincides with the rapid radiation of angiosperms, which began earlier in the Late Cretaceous (~100-70 Ma) but saw major ecological restructuring after the K-Pg extinction event. Angiosperms provided new ecological opportunities and likely contributed to the diversification of Eurytominae, which are associated with plant-host interactions, particularly seeds and gallforming species (Peris & Condamine, 2024). This angiosperm-driven expansion is consistent with the Eurytomidae diversification patterns in both the Old and New World.

As the divergence dates of Eurytomidae were secondarily derived from previous studies (Blaimer et al., 2023; Cruaud et al., 2024), it is not surprising that the dates recovered in our study were similar to previous studies, estimating an age of \sim 83 Ma for crown-group Eurytomidae. However, the ages of the split between Heimbrinae and Eurytominae and the crown age of Eurytominae are estimated to be much older in our study (\sim 75.5 and \sim 70.9 Ma, respectively), when compared to Cruaud et al. (2024) (\sim 66.3 and \sim 33.5 Ma, respectively). This could be the result of more comprehensive taxonomic sampling in our study, or the lack of fossil calibrations within Eurytominae. Nevertheless, these results align with the biogeographic history of the Late Cretaceous and early Paleogene, during which South America and Africa were in the final stages of separating (\sim 100-80 Ma), and the Atlantic Ocean was widening. The mostly Neotropical origin of Eurytomidae is also consistent with Cruaud et al. (2024). In the Paleogene, Eurytominae split into two major clades, one predominantly found in the Americas (Nearctic + Neotropical), while the other has moved into the Old World (Palearctic, Afrotropical, Indomalayan, Australasian) with intermittent shifts back into the Nearctic. This transcontinental expansion likely reflects a combination of tectonic events, the development of tropical forests in the Paleogene and the global dispersal of angiosperms, which facilitated the spread of plantassociated species (Benton et al., 2022; Peris & Condamine, 2024). The role of plate tectonics in facilitating these shifts-such as the northward drift of India and the eventual collision with Asia-may have contributed to the diversification and establishment of the group in the Indomalayan and Australasian regions.

It is worth pointing out that while the Mid-Jurassic (169–162 Ma) is a common estimated age of origin for Chalcidoidea (Blaimer

FIGURE 2 Chronogram of Eurytomidae inferred in MCMCTree under likelihood approximation, root age with hard bound at 91.5 Ma and independent rates clock using fixed consensus topology of 70p_SWSC. Age range in 95% Highest Posterior Density (HPD) is shown as blue bars. Ancestral biogeographic reconstruction using the DIVALIKE+J model, with the geographic regions as Neotropical (S), Nearctic (N), Afrotropical (A), Palearctic (P), Indomalayan (O) and Australasian (U). Pie charts at nodes indicate likely geographic region assignments.



FIGURE 3 Ancestral state reconstruction of Eurytomidae biology based on the 70p_SWSC matrix, inferred using the ARD model. Green, unknown biology; orange, parasitoid; purple, phytophagous.

et al., 2023; Cruaud et al., 2024), there is a 30 Ma gap between this age and the oldest known chalcidoid fossil from the Early Cretaceous (~128 Ma); thus, more detailed studies are needed to confirm the validity of the divergence dates (Rasplus et al., 2025; Zhang et al., 2025). Eurytomidae is very rare in the fossil record, and to date, the oldest confirmed fossil is an undescribed species from the French Oise fossil (55.8–48.6 Ma) (Brasero & Martin, 2009), which likely represents an extinct lineage as it does not match any of the four extant

subfamilies. Additionally, three more confirmed Eurytomidae fossils have been described: one from the Green River formation in Wyoming, USA (50.3–46.2 Ma), and two from the Florissant Shale in Colorado, USA (37.2–33.9 Ma) (Rasplus et al., 2025). All three of these species have been assigned to *Eurytoma*, but this will need to be reexamined given the polyphyletic nature of the genus. Nevertheless, the rapid diversification of angiosperms and other phytophagous insects during the Cretaceous, coupled with the continued breakup of Pangea, likely enabled Eurytomidae to achieve a cosmopolitan distribution and develop such diverse host ranges, warranting further study.

Evolution of secondary phytophagy

We tested three different models of ancestral state reconstruction, with the ARD model receiving the lowest AIC_c score (Figure 3, Table S6). The ancestral state of Eurytomidae is entomophagy, which is observed in Rileyinae, Buresiinae and Heimbrinae. Phytophagy is likely the ancestral state of the subfamily Eurytominae, as the earliest branching lineages (e.g., *Ausystole, Aranedra* and *Proseurytoma*) are all phytophagous. Entomophagy has evolved at least seven times within Eurytominae, including genera such as *Aximopsis, Chryseida, Eurytoma s.s.*, *Philolema s.l.*, *Sycophila* and *Tenuipetiolus*.

When mapping the phytophagous clades on the tree, the propensity for secondary phytophagy appears a number of times according to our ancestral state reconstruction analysis. Phytophagy (including seed feeders and gall formers) occurs in the earliest branching clades of eurytomines in our study, which are mostly Neotropical. However, it is difficult to assess the true biology of many of the eurytomine species as their larvae can be parasitoids that attack their hosts in concealed habitats (galls, twigs or fruit), making it difficult to establish the trophic relationships. In addition, it is known that larvae of some species start their development as parasitoids, and once the host is consumed, continue their growth by feeding on plant tissues (La Salle, 2005). Finally, some species, such as Sycophila developing at the expense of an epichrysomallid in sycones of figs, are known to be inguilines (Lotfalizadeh et al., 2024). Thus, masked phytophagy may be widespread, and detailed studies are needed to determine the feeding habits for many groups of eurytomine wasps.

The adaptation from an entomophagous ancestor to phytophagy indeed presupposes that the larvae are able to develop with a completely different diet and have the relevant esterases for this purpose. The switch to full herbivory could be facilitated by the intermediate entomophytophagy lifestyle, as the larvae were already spending much of their life embedded in plant tissues (La Salle, 2005; Tooker & Giron, 2020). Eurytomine species are capable of galling different parts of the plants (roots, twigs, flowers or maturing fruit); it is possible that the larva exudes chemicals mimicking plant hormones for inducing a rapid cell division. Gall induction has evolved multiple times within Chalcidoidea, including within Eurytomidae itself (Cruaud et al., 2024; La Salle, 2005; Tooker & Giron, 2020). To date, no comparative studies have been done on gall induction across Chalcidoidea and/or Hymenoptera, so it is unknown how similar the gall induction process by Eurytomidae is compared to the relatively better studied gall wasps and allies (superfamily Cynipoidea). The galls of chalcidoids, including eurytomids, are relatively simple and do not have many defensive traits compared to galls induced by cynipoids, yet they occur across a much wider variety of host plants ranging from monocot to dicot (La Salle, 2005). Similarly, many eurytomid species are capable of feeding on a variety of seeds, some of which have toxic

chemicals (e.g., Rosaceae, Fabaceae), but little is known whether they can sequester these toxic chemicals for their protection against seed eaters, and whether the detoxification of different chemistries of the host plant has led to the diversification of eurytomids.

Blaimer et al. (2023) examined the link between innovations and diversification in Hymenoptera, including secondary phytophagy. The authors recover a positive diversification rate in the Cynipidae and the Eurytominae, both lineages that have repeated evolution of secondary phytophagy. It is thus possible that the plasticity of Eurytominae larvae for the diet provides an advantage by allowing new niches to be explored, increasing their radiation rate and diversity. A large number of undescribed eurytomines have been recorded from various plants in tropical forest habitats (GD, unpublished data). Based on these observations, we can anticipate that the actual number of species is much higher than currently recognized. The predominantly endophytic lifestyle has no doubt led to a very successful global distribution and rapid diversification of eurytomine wasps, as shown by the short branch lengths in our analyses. Future studies with more detailed taxon sampling of eurytomines could focus on, for example, whether sister group pairs with different life history strategies and/or hosts with dissimilar chemicals have different diversification rates.

AUTHOR CONTRIBUTIONS

Y. Miles Zhang: Investigation; writing - original draft; methodology; validation; writing - review and editing; visualization; software; formal analysis; data curation. Gérard Delvare: Conceptualization; investigation; writing - review and editing; visualization; validation; data curation; resources; writing - original draft. Bonnie B. Blaimer: Conceptualization; investigation; writing - review and editing; visualization; methodology; validation; software; project administration; data curation. Astrid Cruaud: Investigation; writing - review and editing; resources; funding acquisition; data curation; methodology; validation; software. Jean-Yves Rasplus: Conceptualization; investigation; funding acquisition; writing - review and editing; resources; project administration; supervision; data curation. Seán G. Brady: Writing - review and editing; conceptualization; resources; project administration; supervision; funding acquisition. Michael W. Gates: Conceptualization; investigation; funding acquisition; writing - review and editing; validation; project administration; resources; supervision; data curation; writing - original draft.

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CONFLICT OF INTEREST STATEMENT

The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

Clean FASTQ files for each sample newly generated in this study are uploaded onto the SRA database under Bioproject PRJNA1217593, *COI* barcodes are uploaded onto GenBank under accession numbers PV031733-PV031868. All UCE matrices and codes used for analyses are uploaded onto Dryad https://doi.org/10.5061/dryad.3bk3j9kx3.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

File S1. Taxonomic discussion.

Figure S1. Raw output tree for the 70% UCE matrix, inferred under the SWSC-EN partitioning, partition merging and ModelFinder model selection, generated using IQ-TREE (70p_SWSC).

Figure S2. Raw output tree for the 70% UCE matrix, inferred under MFP + MERGE and filtered using SRH test, with concordance factors, generated using IQ-TREE (70p_CONCORD).

Figure S3. Raw output tree for the 70% protein-coding-gene matrix, inferred under PMSF Class 60 model in IQ-TREE with 70p_SWSC as input guide tree, generated using IQ-TREE (70p_PMSF_C60).

Figure S4. Raw output multispecies coalescent tree for the 70% UCE matrix, with individual best gene trees generated in IQ-TREE, boot-strap <10 collapsed, final species tree generated using ASTRAL-III (70p_BS10_ASTRAL).

Figure S5. Raw output tree for the 50% UCE matrix, inferred under the SWSC-EN partitioning, partition merging and ModelFinder model selection, generated using IQ-TREE (50p_SWSC).

Figure S6. Raw output tree for the 50% UCE matrix, inferred under MFP + MERGE and filtered using SRH test, with concordance factors, generated using IQ-TREE (50p_CONCORD).

Figure S7. Raw output tree for the 50% protein-coding-gene matrix, inferred under PMSF Class 60 model in IQ-TREE with 50p_SWSC as input guide tree, generated using IQ-TREE (50p_PMSF_C60).

Figure S8. Raw output multispecies coalescent tree for the 50% UCE matrix, with individual best gene trees generated in IQ-TREE,

bootstrap <10 collapsed, final species tree generated using ASTRAL-III (50p_BS10_ASTRAL).

Figure S9. Output for MCMCtree divergence dating analysis, node numbers and 95% HPD are displayed. Modified using MCMCtreeR.

Figure S10. Raw output from BioGeoBEARS for DIVALIKE+J model.

Table S1. Voucher information and UCE stats.

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Table S2. Matrix summary output from AMAS.

 Table S3. Biogeography and feeding habits.

 Table S4. Node ages for divergence dating.

 Table S5. (A) Dispersal rate scaler for BioGeoBEARS, (B) Model selection results for biogeography.

Table S6. Model selection results for ancestral state reconstruction.

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