



# Revisiting the host use and phylogeny of Colastomion Baker (Hymenoptera, Braconidae, Rogadinae), with a new host record from Japan

Sakagami, Kota  
Shimizu, So  
Fujie, Shunpei  
Maeto, Kaoru

---

## (Citation)

Journal of Hymenoptera Research, 77:175-186

## (Issue Date)

2020-06-29

## (Resource Type)

journal article

## (Version)

Version of Record

## (Rights)

Copyright Kota Sakagami et al.

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

## (URL)

<https://hdl.handle.net/20.500.14094/90009299>



# Revisiting the host use and phylogeny of *Colastomion* Baker (Hymenoptera, Braconidae, Rogadinae), with a new host record from Japan

Kota Sakagami<sup>1,2\*</sup>, So Shimizu<sup>1,2,3\*</sup>, Shunpei Fujie<sup>4</sup>, Kaoru Maeto<sup>1</sup>

**1** Laboratory of Insect Biodiversity and Ecosystem Science, Graduate School of Agricultural Science, Kobe University, Rokkodaicho 1–1, Nada, Kobe, Hyogo 657–8501, Japan **2** Research Fellow (DC), Japan Society for the Promotion of Science, Tokyo, Japan **3** Department of Life Sciences, the Natural History Museum, Cromwell Road, London SW7 5BD, UK **4** Osaka Museum of Natural History, Nagaikoen 1–23, Higashisumiyoshi, Osaka 546–0034, Japan

Corresponding author: Kota Sakagami ([kota.sakagami1@gmail.com](mailto:kota.sakagami1@gmail.com))

Academic editor: J. Fernandez-Triana | Received 31 March 2020 | Accepted 8 June 2020 | Published 29 June 2020

<http://zoobank.org/F1CDBD46-5D8D-42A4-AAA9-0F88A4D78505>

**Citation:** Sakagami K, Shimizu S, Fujie S, Maeto K (2020) Revisiting the host use and phylogeny of *Colastomion* Baker (Hymenoptera, Braconidae, Rogadinae), with a new host record from Japan. Journal of Hymenoptera Research 77: 175–186. <https://doi.org/10.3897/jhr.77.52774>

## Abstract

We report the solitary parasitism by *Colastomion formosanum* (Watanabe) (Hymenoptera, Braconidae, Rogadinae) on the larva of *Nevrina procopia* (Stoll) (Lepidoptera, Crambidae) feeding on *Turpinia ternata* Nakai (Staphyleaceae) in Amami Ōshima Is., Japan. This is the first host record for the genus *Colastomion* Baker outside of Papua New Guinea. We have also inferred the phylogenetic relationships of *Colastomion* species using Bayesian and maximum likelihood approaches, based on the mitochondrial cytochrome oxidase 1 gene. The results indicate two major clades—solitary and gregarious parasitoids—within *Colastomion*. *Colastomion formosanum* belongs to the clade of solitary parasitoids that specifically parasitize the crambid subfamily Spilomelinae. Plant-host-parasitoid associations and the evolutionary scenario of the host use of *Colastomion* are discussed.

## Keywords

*Colastomion formosanum*, Crambidae, endoparasitoid, mtCO1, mummy, *Nevrina procopia*

\* Contributed equally as the first authors.

## Introduction

The family Braconidae is one of the largest lineages in Hymenoptera, containing 21,221 valid species worldwide (Yu et al. 2016). Braconid wasps are parasitoids of various insects (e.g. Lepidoptera, Coleoptera, and Hymenoptera), including important agricultural pests (Wharton 1993; Quicke 2015; Maeto 2018). Their amazing diversity has resulted from the complicated biological interactions with host insects as well as contrasting lifestyles, e.g. solitary vs. gregarious, ecto- vs. endoparasitic, or idio- vs. koinobiont (Shaw 1988). Understanding their biology and phylogeny lends us a greater appreciation of the evolutionary pattern of host-parasitoid systems. Further, from a practical standpoint, the study of Braconidae can also be useful to leverage their parasitic abilities for biological pest control (Quicke 2015).

The subfamily Rogadinae Förster *sensu stricto* contains diverse koinobiont endoparasitoids of Lepidoptera and comprises 63 genera and 1,243 species that are distributed across all zoogeographical regions, except for polar regions (Quicke 2015; Yu et al. 2016). One of the marked biological features of the rogadine parasitoids is the mummification of their host larvae (Chen and He 1997; Zaldívar-Riverón et al. 2008).

*Colastomion* Baker is an uncommon genus of Rogadinae and comprises 15 species that occur throughout Papua New Guinea, southern East Asia, and Africa (Quicke et al. 2012; Yu et al. 2016). Crambid larvae have been reported to be hosts of *Colastomion* only in Papua New Guinea (Quicke et al. 2012). While the host larvae of *Colastomion* are typically mummified like other rogadine parasitoids, the mummy has been illustrated for only one species of *Colastomion* to date (Zaldívar-Riverón et al. 2008). Also, although the phylogenetic placement of *Colastomion* within Rogadinae has been largely established (Zaldívar-Riverón et al. 2008), the phylogenetic relationships within the genus has not been firmly estimated (Quicke et al. 2012).

Recently, the first and second authors (KS and SS) conducted field studies in Amami Ōshima Is., Kagoshima Pref., Japan, and obtained *Colastomion* specimens from host caterpillars and in a light trap. Here, we identified the *Colastomion* specimens as a species and conducted phylogenetic analyses based on the mitochondrial cytochrome oxidase 1 (*COI*). These provide the first host record of *Colastomion* outside of Papua New Guinea, detailed mummy morphology, and the evolutionary scenario of host use within *Colastomion*.

## Materials and methods

### Field collection and rearing

A field study was conducted at Naze-Ōaza-Chinase (28°21'N, 129°26'E, 16 m alt.), Amami Ōshima Is., Kagoshima Pref., southwest Japan on 11 April 2019. The study site was the evergreen broad-leaved forest dominated by *Ficus* spp., *Ligustrum japonicum* Thunb. and *Pittosporum tobira* (Thunb.) W.T.Aiton. KS collected larvae of *Nevrina procopia* (Stoll)

(Lepidoptera, Crambidae, Spilomelinae) as they were feeding on *Turpinia ternata* Nakai (Staphyleaceae). The larvae rolled young leaves roughly and hid themselves with the rolled leaves (Fig. 1). Collected larvae were reared in plastic cases under laboratory conditions (25 °C, 16:8 h light:dark). Emerged insects were killed in a freezer.

Further specimen was collected in a field study at Nishinakama (28°15'47.3"N, 129°24'56.1"E, 120 m alt.), Sumiyô Town, Amami Ôshima Is. on 12 July 2019, using High Intensity Discharge light traps by SS.

### Specimens examined, repositories and identification

The examined specimens of *Colastomion* from Japan are deposited in the Institute for Agro-Environmental Sciences, NARO, Tsukuba, Japan (NIAES) and Osaka Museum of Natural History, Osaka, Japan (OMNH): 1♀, Naze-Ôaza-Chinase, Amami City, Amami Ôshima Is., Kagoshima Pref. (28°21'N, 129°26'E, 16 m alt.), 11.IV.2019, K. Sakagami leg. (by rearing host) (NIAES) [DDBJ-LC485659]; 1♀, Nishinakama, Sumiyô Town, Amami Ôshima Is., Kagoshima Pref. (28°15'47.3"N, 129°24'56.1"E, 120 m alt.), 12.VII.2019, S. Shimizu leg. (light trap) (NIAES) [DDBJ-LC499982]; 1♀, Mt. Yui-dake, Setouchi Town, Amami Ôshima Is., Kagoshima Pref., 24.VIII.2004, H. Makihara leg. (sweeping net) (OMNH); 1♂, Yona, Kunigami Village, Okinawa-hontô Is., Okinawa Pref., 29.VI.2013, S. Fujie leg. (sweeping net) (OMNH).

We identified *Colastomion* specimens based on Watanabe (1932, 1934), Tenma (2002), and Quicke et al. (2012): the specimens were morphologically similar to *C. formosanum*. To make sure that *Colastomion* specimens are *C. formosanum*, all the specimens were compared to photos of the Taiwanese specimens of *C. formosanum* deposited in the Senckenberg Deutsches Entomologisches Institut, Müncheberg, Germany (SDEI), which include the holotype of the species as follows: holotype ♂, Kankau, Changkou (Koshun), 22.IV.1912, H. Stauter leg.; 1♂, Kankau, Changkou (Koshun), VIII.1912, H. Stauter leg.; 1♀, Kankau, Changkou (Koshun), IV.1912, H. Stauter leg. The photos were provided by Taeger (2020), published online (<https://doi.org/10.6084/m9.figshare.11984355.v1>).

### Morphological observation, photo technique and terms

Morphological observation was conducted using a stereoscopic microscope (SZ61, OLYMPUS, Tôkyô, Japan). Multi-focus photographs were taken using a single lens reflex camera (α7II, Sony, Tôkyô, Japan) with micro-lens (LAOWA 25 mm F2.8 2.5–5X ULTRA MACRO, Anhui Changgeng Optics Technology Co., Ltd, Hefei, China and A FE 50mm F2.8 Macro SEL50M28, Sony, Tôkyô, Japan). The photos were captured in RAW format and developed using Adobe Lightroom Creative Cloud. Then, they were stacked using Zerene Stacker and edited in Adobe Illustrator 2019. Morphological terms follow those of Quicke et al. (2012).



**Figure 1.** Leaves of *Turpinia ternata* Nakai rolled by a caterpillar of *Nevrina procopia* (Stoll).



## Molecular technique

Part of the mitochondrial protein-coding cytochrome c oxidase 1 (*COI*) gene, often referred to as “barcoding gene”, of two individuals of *C. formosanum* was sequenced for phylogenetic analysis. DNA was extracted from the right middle leg using the DNeasy Blood and Tissue Kit (Qiagen, Düsseldorf, Germany). For amplification, the following primers were used: LCO1490 (5'-GGTCAACAAATCATAAAGA-TATTGG-3') and HCO2198 (5'-TAAACTTCAGGGTGACCAAAAAATCA-3') (Folmer et al. 1994). Polymerase chain reactions (PCR) were conducted using KOD FX NEO kit (Toyobo, Ōsaka, Japan), and PCR conditions were 94 °C for 2 min as initial denaturation, followed by 35 cycles of denaturation (10 sec at 98 °C), annealing (30 sec at 48 °C), and extension (30 sec at 68 °C), and then a final extension at 72 °C for 10 min. PCR product was purified using Illustra GFX kit (GE Healthcare Life Sciences, Marlborough, USA). The purified product was amplified with the same primers using the BigDye™ Terminator v3.1 Cycle Sequencing kit (Applied Biosystems, Waltham, USA). Cycle sequencing products were purified using the 3.0 M sodium acetate, 95% ethanol, 70% ethanol, and Hi-Di formamide. Cycle sequencing reactions were run on an ABI Prism 3100 Genetic Analyzer (Applied Biosystems, Waltham, USA), and the forward and reverse sequences were assembled using the DNA Dynamo Sequence Analyze Software (Blue Tractor Software, North Wales, UK). Finally, we deposited the obtained sequence to DNA Data Bank of Japan (DDBJ).

## Phylogenetic analyses

In order to exclude the taxon sampling bias, a single sequence for each species was selected from 42 sequences of *COI* of the nine *Colastomion* species from Papua New Guinea and Benin deposited in the DNA databases (Quicke et al. 2012). *Megarhogas maculipennis* Chen & He and *Myocron* sp., closely related genera to *Colastomion* (Zaldívar-Riverón et al. 2008), were selected as outgroups. A total of 10 sequences from ingroup and two from outgroups were used for analyses (see Table 1).

Multiple sequence alignment was conducted in MAFFT v7.409 (Katoh and Toh 2008) using the L-INS-I algorithm. The aligned sequences were checked visually. Subsequently, they were manually optimized for phylogenetic analysis.

Using Bayesian Inference (BI) and maximum likelihood (ML) approaches, phylogenetic analyses were performed. Evolutionary models were determined using jModelTest v2.1.10 (Tanabe 2011). The best-fit models were selected based on the lowest corrected Akaike information criterion (AICc) for ML and the lowest Bayesian Information Criterion (BIC) for BI.

For the ML analysis, we used RAxML v8.2.10 (Stamatakis 2014) with 1,000 bootstrap replications, the codon separate model, and GTRGAMMA as a substitution model.

**Table 1.** GenBank and DNA Data Bank of Japan accession numbers and information for specimens used for the phylogenetic analyses.

Species	Identifier	Locality	Latitude / longitude	Date	Collector	Accession number
<i>Colastomion crambidiphagus</i>	DLJ Quicke	PNG: Madang, Wanang	5.23088S, 145.182E	16.II.2007	local collector	JF963127
<i>Colastomion formosanum</i>	K Maeto	JPN: Amami-Oshima, Kagoshima	28.3500N, 129.433E	11.IV.2019	K. Sakagami	LC485659
<i>Colastomion gregarius</i>	DLJ Quicke	PNG: Madang, Wanang	5.23088S, 145.182E	24.V.2007	local collector	JF963128
<i>Colastomion maclayi</i>	DLJ Quicke	PNG: East Sepik, Yapsiei	4.62825S, 141.097E	27.I.2004	local collector	JF271312
<i>Colastomion madangensis</i>	DLJ Quicke	PNG: Madang, Wanang	5.23088S, 145.182E	24.V.2007	local collector	JX034716
<i>Colastomion masalaii</i>	DLJ Quicke	PNG: West Sepik, Sandaun, Uta	3.38405S, 141.586E	28.VII.2004	local collector	JF271307
<i>Colastomion parotiphagus</i>	DLJ Quicke	PNG: Madang, Wanang	5.23088S, 145.182E	30.V.2007	local collector	JX034711
<i>Colastomion pukpuk</i>	DLJ Quicke	PNG: East Sepik, Wamangu	3.78713S, 143.652E	3.XI.2005	local collector	JF271303
<i>Colastomion wanang</i>	DLJ Quicke	PNG: Madang, Wanang	5.23088S, 145.182E	29.IV.2005	local collector	JF271302
<i>Colastomion</i> sp.	–	BEN	–	–	–	AY935370
<i>Myocron</i> sp.	–	KEN	–	5.V.2005	R Copeland	JN278218
<i>Megarhogas maculipennis</i>	–	THA: Chanta Bari, Pong Nani Ron	–	–	–	EU979615

\*BEN, Benin; JPN, Japan; KEN, Kenya; PNG, Papua New Guinea; THA, Thailand.

For the BI analysis, we used MrBayes v3.2.2 (Ronquist et al. 2012) with two independent runs of a Bayesian Markov chain Monte Carlo (MCMC) analyses of eight chains each, heating at 0.1, as well as random starting trees with trees sampled every 1,000<sup>th</sup> generations for 10,000,000 generations. If the average standard deviation of split frequencies was below 0.01, chain stationarity was checked with Tracer v1.6 (Rambaut and Drummond 2007) and two converged MCMC runs were considered adequate (Ronquist and Huelsenbeck 2003). The anterior half of the generations were discarded as a conservative burn-in and estimates were obtained for the harmonic means of the likelihood scores from the remaining half generations using the sum command. A final check of the convergence of the runs by the value of the potential scale reduction factor was conducted and a majority-rule consensus tree was obtained using the sumt command. The phylogenetic tree was edited using FigTree v1.4.3 (Rambaut 2006–2016) and Adobe Illustrator 2019.

We consider the node to be supported by either the Bayesian posterior probabilities (PP) > 0.95 or the bootstrap (BT) > 80%.

## Results

### Rearing and mummy morphology

One adult female of *Colastomion* emerged from a mummified final instar larva of *N. procopia* on 30 April 2019 (Figs 2, 3). The mummy remained unfixed within rolled leaves. It was mildly hardened, having a posterolateral and irregular, noncircular emergence hole (Fig. 4). Five adults of *N. procopia* emerged from unparasitized pupae, of which two emerged on 30 April, two on 8 May, and one on 9 May 2019.

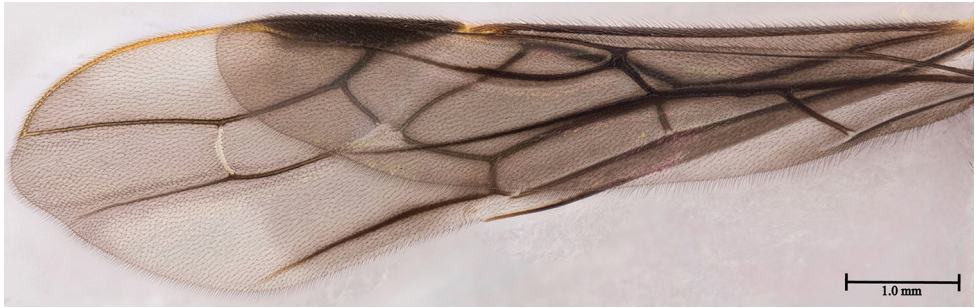


**Figure 2.** Female adult wasp of *Colastomion formosanum* (Watanabe) from Amami Ōshima Is., Japan **A** habitus **B** head, frontal view **C** head and mesosoma, lateral view **D** head and mesosoma, dorsal view **E** propodeum, dorsal view **F** metasoma, dorsal view.

### Identification and adult morphology

All Japanese *Colastomion* specimens were identified as *C. formosanum* because of yellow face, antennae (although apical segments were brown) and legs (although middle and hind coxae and telotarsi were brown), sharply contrasting with brown mesosoma





**Figure 3.** Wings of *Colastomion formosanum* (Watanabe).



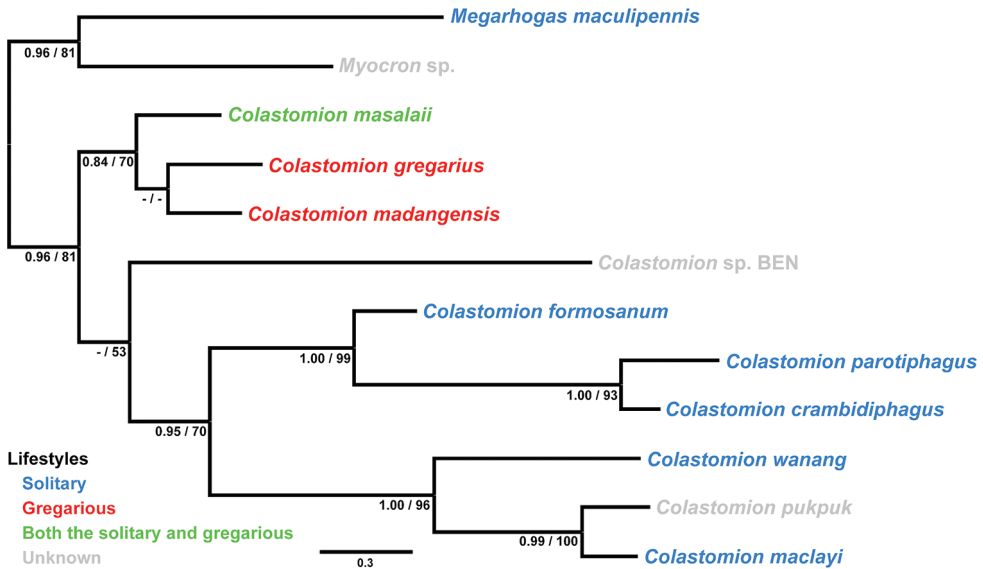
**Figure 4.** A larva of *Nevirina procopia* (Stoll) mummified by *Colastomion formosanum* (Watanabe), lateral view.

and metasoma (Fig. 2A, B); epicnemial area finely strigose, precoxal sulcus shallowly impressed and with some rugae, and pleural sulcus crenulate (Fig. 2C); notauli deep and weakly crenulate (Fig. 2D); propodeum rugose and with complete midlongitudinal carina (Fig. 2E); 1<sup>st</sup> metasomal tergite 1.7–1.8× longer than posteriorly wide; 2<sup>nd</sup> tergite as long as maximally wide (Fig. 2F); pterostigma entirely dark brown; fore wing cu-a postfurcal to 1-M (Fig. 3); and subquadrate vein 2-SC+R of hind wing (Fig. 3).

A female specimen from Taiwan shows a distinct protuberance on the base of the first metasomal tergite (Taeger 2020), whereas it is absent in males (including holotype) from Taiwan (Taeger 2020) and females and a male from Japan. The protuberance will probably be due to ontogenetic deformation in the individual. On the other hand, the sculpture of epicnemial area and precoxal sulcus tends to be stronger in females than in males, which is most likely a sexual variation.

## Phylogeny

The Bayesian majority-rule consensus tree of *Colastomion* obtained from the *COI* sequences is shown in Figure 5. Both the BI and ML topologies were consistent with each other. *Colastomion formosanum* was identified as a sister group of the well-supported clade (*C. crambidiphagus* and *C. parotiphagus*) by both the BT and PP. Monophyly



**Figure 5.** Bayesian majority-rule consensus tree of *Colastomion* species based on *COI*. Posterior probabilities (> 0.80) and bootstrap values (> 50%) are indicated at below of each node (PP / BT). Lifestyles are indicated as shown in Quicke et al. (2012), except for *C. formosanum*.

of the clade of solitary parasitoids (*C. crambidiphagus*, *C. formosanum*, *C. maclayi*, *C. parotiphagus*, and *C. wanang*) plus *C. pukpuk* was good supported only by the PP. On the other hand, the clade of gregarious parasitoids (*C. masalaii*, *C. gregarius*, and *C. madangensis*) was not fully supported.

## Discussion

We have found that *N. procopia* feeding on rolled leaves of *T. ternata* are the host species of *C. formosanum* in Japan. This supports the hypothesis of the host specificity of *Colastomion* to the crambids subfamily Spilomelinae as mentioned by Quicke et al. (2012), whereas the genus *Nevrina* of the tribe Udeini is a new host genus and the Staphyleaceae is a new host plant family for *Colastomion* (Table 2). Interestingly, Quicke et al. (2012) has reported that two solitary species of *Colastomion* use some species of crambid moths on a certain plant: *C. crambidiphagus* parasitizes several species of different tribes of Spilomelinae feeding only on the Convolvulaceae, and *C. parotiphagus* uses various host tribes of moth while mostly feeding on the Rubiaceae. Moreover, all gregarious species (*C. gregarius*, *C. madangensis*, and *C. masalaii*) consistently use Margaroniini on the Moraceae (Table 2). These may indicate that adult wasps search the host larvae by plant cues, such as herbivory-induced plant volatiles (Arimura et al. 2009) or oviposition-induced plant volatiles (Hilker and Fatoaros 2015). It is therefore important to understand the specificity of parasitoids not only in host insects but also in host plants.

**Table 2.** Host tribes (Crambidae: Spilomelinae), host plant families, and lifestyles of *Colastomion*. Sources: Quicke et al. (2012), except for *C. formosanum*. Systematics of host tribes follows Mally et al. (2019).

Species	Host tribe	Host plant family	Lifestyle
<i>C. crambidiphagus</i>	Hydririni, Udeini	Convolvulaceae	Solitary
<i>C. formosanum</i>	Udeini	Staphyleaceae	Solitary
<i>C. gregarius</i>	Margaroniini	Moraceae	Gregarious
<i>C. maclayi</i>	Udeini	Rubiaceae	Solitary
<i>C. madangensis</i>	Margaroniini	Moraceae	Gregarious
<i>C. masalaii</i>	Margaroniini	Moraceae	Solitary/gregarious
<i>C. parotiphagus</i>	Agroterini	Malvaceae	Solitary
ditto	Margaroniini	Rubiaceae	Solitary
ditto	Unidentified	Lauraceae, Ulmaceae	Solitary
<i>C. pulkruk</i>	Unidentified	Rubiaceae	Unknown
<i>C. wanang</i>	Udeini	Myrtaceae, Vitaceae	Solitary

*Colastomion formosanum* (Figs 2, 3) is a solitary endoparasitoid that forms a hard mummy (Fig. 4). Solitary parasitism in *Colastomion* is considered the ancestral relative to gregarious parasitism because *Megarhogas*, which is closely related to *Colastomion* (and is included as an outgroup in our phylogenetic analyses), and most species in Rogadini are solitary (Quicke and Shaw 2005; Zaldívar-Riverón et al. 2008), while the phylogenetic placement of gregarious species is still unresolved (Fig. 5). The formation of a hard mummy as displayed by *C. formosanum* remains unusual within Rogadiinae, although various types of host mummies appear to act as a protective roll during parasitoid metamorphosing (Zaldívar-Riverón et al. 2008; Maeto 2018). The hosts of most rogadoine genera are killed as prepupae within host cocoons, which can protect the parasitoid larvae and pupae, and thus relatively frail mummies are the norm. In contrast, the hosts of genera that form hard mummies are killed in the larval stage, and the hardness of the mummies plays a vitally important protective role. In the case of *C. formosanum*, the host is killed in the larval stage probably because the immature host caterpillar is large enough for the parasitoid. Host stage of mummifying, which may vary according to relative size of host caterpillars to parasitoids, would therefore be relevant to the hardness of mummies.

*Colastomion formosanum* was originally described from Taiwan (as Formosa) (Watanabe 1932), later recorded from Hainan Is., China (Chen and He 1997), and was most recently recorded from Okinawa-hontô Is. and Iriomotejima Is., Japan (Tenma 2002). Our collection of *C. formosanum* from Amami Ôshima Is., Japan, has therefore expanded the northernmost border of the genus *Colastomion* as well as the species. It is most likely that *C. formosanum* has advanced northward from the tropics where species of *Colastomion* have highly diversified (Table 1; Fig. 5), although further investigations are needed to reveal the biogeographical history of *Colastomion*. *C. formosanum* is found only in the subtropical islands of China and Japan, whereas its host moth *N. procopia* is distributed much more broadly, from southern Japan to China, India, Papua New Guinea and western Africa (Swinhoe 1916; Chandra 1994; Nasu et al. 2013; Poltavsky et al. 2018). It would be interesting to know whether *C. formosanum* is truly an insular species of East Asia or is also further distributed in continental parts.

## Acknowledgements

We thank Satoru Matsubara, Masafumi Harada, and Naoto Shimada (Kobe University) for assistance with the field observation of KS. We are grateful to Andreas Taeger (SDEI) for providing photos. This research is partially supported by the Grant-in-Aid for JSPS Fellows Grant Number 18J20333 to SS and the JSPS KAKENHI Grant Number 19H00942 to KM. We also thank Donald L. Quicke for making very helpful comments.

## References

- Arimura GI, Matsui K, Takabayashi J (2009) Chemical and molecular ecology of herbivore-induced plant volatiles: proximate factors and their ultimate functions. *Plant and Cell Physiology* 50: 911–923. <https://doi.org/10.1093/pcp/pcp030>
- Chandra K (1994) Further new records of moths from Andaman and Nicobar Islands. *Journal of Andaman Science Association* 10: 17–24.
- Chen X, He J (1997) Revision of the subfamily Rogadinae (Hymenoptera: Braconidae) from China. *Zoologische Verhandelingen* 308: 1–187.
- Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R (1994) DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. *Molecular Marine Biology and Biotechnology* 3: 294–299.
- Hilker M, Fatouros NE (2015) Plant responses to insect egg deposition. *Annual Review of Entomology* 60: 493–515. <https://doi.org/10.1146/annurev-ento-010814-020620>
- Katoh K, Toh H (2008) Improved accuracy of multiple ncRNA alignment by incorporating structural information into a MAFFT-based framework. *BMC Bioinformatics* 9: 1–212. <https://doi.org/10.1186/1471-2105-9-212>
- Maeto K (2018) Polyphagous koinobiosis: the biology and biocontrol potential of a braconid endoparasitoid of exophytic caterpillars. *Applied Entomology and Zoology* 53: 433–446. <https://doi.org/10.1007/s13355-018-0581-9>
- Mally R, Hayden JE, Neinhuis C, Jordal BH, Nuss, M (2019) The phylogenetic systematics of Spilomelinae and Pyraustinae (Lepidoptera: Pyraloidea: Crambidae) inferred from DNA and morphology. *Arthropod Systematics & Phylogeny* 77: 141–204. <https://doi.org/10.26049/ASP77-1-2019-07>
- Nasu Y, Hirowatari T, Kishida Y (2013) *The Standard of Moths in Japan* 4. Gakken Education Publishing, Tokyo, 552 pp.
- Poltavsky AN, Kravchenko VD, Traore MM, Traore SF, Gergely P, Witt TJ, Sulak H, Beck RH, Junnila A, Revay EE, Doumbia S, Beier JC, Muller GC (2018) The Pyraloidea (Lepidoptera) fauna of the woody savannah belt in Mali, West Africa. *Zootaxa* 4457: 39–69. <https://doi.org/10.11646/zootaxa.4457.1.2>
- Quicke DLJ (2015) *The Braconid and Ichneumonid Parasitoid Wasps: Biology, Systematics, Evolution and Ecology*. Wiley Blackwell, New Jersey, 733 pp. <https://doi.org/10.1002/9781118907085>
- Quicke DLJ, Shaw MR (2005) First host records for the rogaeline genera *Rogasodes* Chen and He and *Canalirogas* van Achterberg and Chen (Hymenoptera: Braconidae) with descrip-



- tion of a new species and survey of mummy types within Rogadinae s. str. *Journal of Natural History* 39: 3525–3542. <https://doi.org/10.1080/00222930500392782>
- Quicke DLJ, Smith MA, Miller S, Hrcek J, Butcher B (2012) *Colastomion* Baker (Braconidae, Rogadinae): nine new species from Papua New Guinea reared from Crambidae. *Journal of Hymenoptera Research* 28: 85–121. <https://doi.org/10.3897/jhr.28.3484>
- Rambaut A (2006–2016) Tree Figure Drawing Tool, version 1.4.3, Institute of Evolutionary Biology, University of Edinburgh [online]. <http://tree.bio.ed.ac.uk/software/figtree/> [accessed 14 July 2019]
- Rambaut A, Drummond AJ (2007) Tracer: MCMC trace analysis tool v1.4.1. <http://tree.bio.ed.ac.uk/software/tracer?/>
- Ronquist F, Huelsenbeck JP (2003) MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* 19: 1572–1574. <https://doi.org/10.1093/bioinformatics/btg180>
- Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA, Huelsenbeck JP (2012) MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. *Systematic Biology* 61: 539–542. <https://doi.org/10.1093/sysbio/sys029>
- Shaw SR (1988) Euphorine phylogeny: the evolution of diversity in host-utilization by parasitoid wasps (Hymenoptera: Braconidae). *Ecological Entomology* 13: 323–335. <https://doi.org/10.1111/j.1365-2311.1988.tb00363.x>
- Stamatakis A (2014) RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics* 30: 1312–1313. <https://doi.org/10.1093/bioinformatics/btu033>
- Swinhoe C (1916) LVI.—New species of butterflies and moths from Australia, Africa, and the Indo-Malayan region. *Journal of Natural History* 18: 480–490. <https://doi.org/10.1080/00222931609486904>
- Taeger A (2020) *Cystomastax formosana* Watanabe, 1932. figshare. Figure. <https://doi.org/10.6084/m9.figshare.11984355.v1>
- Tanabe AS (2011) Kakusan4 and Aminosan: two programs for comparing nonpartitioned, proportional and separate models for combined molecular phylogenetic analyses of multi-locus sequence data. *Molecular Ecology Resources* 11: 914–921. <https://doi.org/10.1111/j.1755-0998.2011.03021.x>
- Tenma K (2002) Newly recorded genera, *Conspinaria* Schulz, 1906 and *Colastomion* Baker, 1917 (Hymenoptera, Braconidae, Rogadinae) from Japan. *Japanese Journal of Systematic Entomology* 8: 185–191.
- Watanabe C (1932) Description of new Species of Genera *Megarhogas*, *Cytomastax* and *Stantonia* (Braconidae) from Fomosa. *Insecta Matsumurana* 6: 184–189.
- Watanabe C (1934) H. Sauter's Formosa-collection: Braconidae. *Insecta Matsumurana* 8: 182–205.
- Wharton RA (1993) Bionomics of the Braconidae. *Annual Review of Entomology* 38: 121–143. <https://doi.org/10.1146/annurev.en.38.010193.001005>
- Yu DS, van Achterberg C, Horstmann K (2016) World Ichneumonoidea 2016 – Taxonomy, Biology, Morphology and Distribution. Taxapad 2016. Vancouver.
- Zaldívar-Riverón A, Shaw MR, Sáez AG, Mori M, Belokobyl'skij SA, Shaw SR, Quicke DLJ (2008) Evolution of the parasitic wasp subfamily Rogadinae (Braconidae): phylogeny and evolution of lepidopteran host ranges and mummy characteristics. *BMC Evolutionary Biology* 8: 1–329. <https://doi.org/10.1186/1471-2148-8-329>