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Author(s)
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Pollination mutualism between *Alocasia macrorrhizos* (Araceae) and two taxonomically undescribed *Colocasiomyia* species (Diptera: Drosophilidae) in Sabah, Borneo

Kohei Takenaka Takano\(^1\)*, Rimi Repin\(^2\), Dartin Maryati Bte Mohamed\(^3\) and Masanori J. Toda\(^4\)

\(^1\)Graduate School of Environmental Earth Science, Hokkaido University, Japan; \(^2\)Sabah Parks, Malaysia; \(^3\)Institute for Tropical Biology and Conservation, Universiti Malaysia Sabah, Malaysia; and \(^4\)Institute of Low Temperature Science, Hokkaido University, Japan

**Short title:** Pollination mutualism between *Colocasiomyia* flies and *Alocasia*

**Corresponding author:** Kohei Takenaka Takano,

Present address: Research Institute for Humanity and Nature, Kyoto 603-8047, Japan

koheit@chikyu.ac.jp

Phone/Fax: +81-95-819-7063/7064

**Key words:** *Colocasiomyia* sp.1 aff. *sulawesiana*; *Colocasiomyia* sp.2 aff. *sulawesiana*; Kota Kinabalu; life history; pistilicolous species; pollination experiment.
ABSTRACT

Two taxonomically undescribed Colocasiomyia species were discovered from inflorescences of Alocasia macrorrhizos in Kota Kinabalu City, Sabah, Borneo, Malaysia. The aims of this study were to investigate the reproductive ecology of the flies and the plant, ascertain the importance of the flies as pollinators and examine the intimate association between flowering events and life history of the flies. We conducted sampling, observations and field pollination experiments. The flies were attracted by the odour of female-phase inflorescences in the early morning on the first day of anthesis. They fed, mated and oviposited in the inflorescences for one day. On the second day, the flies, covered with pollen grains, left the male-phase inflorescences for the next female-phase inflorescences. The immatures of both fly species hatched, developed and pupated within the infructescences without damaging the fruits and developed adults emerged when the mature infructescences dehisced. The flowering events and fly behaviours were well synchronized. In the field pollination experiments, the inflorescences bagged with a fine mesh (insect exclusion) produced almost no fruits, whereas those bagged with a coarse mesh (bee exclusion) produced as many fruits as the open-pollinated controls. These results indicate that the flies are the most efficient and specialized pollinators for their host, A. macrorrhizos. These flies, in return, depend on A. macrorrhizos for food and habitat through most of their life cycle. This study provides a deeper insight into the less recognized, highly intimate pollination mutualism between Araceae plants and Colocasiomyia flies.
INTRODUCTION

Araceae plants have an unique and characteristic inflorescence made up of a spadix and a spathe. All the inflorescence morphologies observed in the family Araceae can be seen as variations around this same theme (Bown, 2000). In spite of their constant inflorescence design, Araceae plants have developed a great diversity of pollination systems (Gibernau, 2003; Vogel, 2000) partly because of the evolution of unisexual flowers that has allowed the secondary development of sterile flowers and then floral function specialisation such as barrier, odour emission, thermogenesis and food-reward (Gibernau, 2003; Mayo et al., 1997). Development of an enclosing spathe (i.e., a floral chamber) with secondary appearance of a constriction allows the capture of insects in contact with the flowers (Gibernau, 2003; Lack and Diaz, 2001). In species with a floral chamber, once pollinators have been attracted during the female phase, they are then kept within the floral chamber by trap mechanisms or rewards (food, mating partners, shelter from light and etc.) until the end of the anthesis (i.e., pollen release) hours or a few days later (Gibernau, 2003; Lack and Diaz, 2001).

The flowers of several plant families serve as breeding places for pollinator insects. In many cases, the host plants are pollinated by saprophagous flies or beetles and the decaying floral parts (e.g. corollas or male flowers) in turn serve as food for these insects (Sakai, 2002). Some Araceae plants also serve their inflorescences as reproductive sites whereas others mimic the laying site (i.e., faeces, mushrooms and dead animal) of the pollinator flies (Gibernau, 2003; Seymour et al., 2003a).

The members of the genus *Colocasiomyia* de Meijere, 1914 (Diptera: Drosophilidae), which currently consists of about 70 species, are found only on the flowers of the Araceae, Arecaceae and Magnoliaceae (Sultana et al., 2006; Takenaka, 2006). In the case of some species, oviposition and larval development take place on the host inflorescences (Carson and Okada, 1980; Toda and Okada, 1983) and the flies serve as major species-specific pollinators.
The close association between certain species-groups within the genus *Colocasiomyia* and certain host taxa suggest that these insects have had long evolutionary relationships with their host plants (Sultana et al., 2006).

Sharing of a single aroid inflorescence by a pair of fly species with partial niche separation is a widely observed ecological trait of *Colocasiomyia* flies; a pistilicolous (*pistil*: female flower, -*colous*: a suffix originated from a Latin word that means *inhabiting in*) species uses the female inflorescence for oviposition and larval development, whereas stamenicolous (i.e. inhabiting in male flowers) species uses mostly the male inflorescence (Carson and Okada, 1980). Different pairs of fly species have been found on different aroid host species or from different geographic regions (Carson and Okada, 1980; Honda-Yafuso, 1983; Okada, 1975; Okada, 1980; Okada, 1986; Okada and Yafuso, 1989; Toda and Okada, 1983; Yafuso and Okada, 1990). Further, additional *Colocasiomyia* species are being continuously discovered (Sultana, 2002; Sultana et al., 2006; Takenaka, 2006; Takenaka et al., 2006; Toda and Lakim, 2011) and show different patterns of species coexistence: some monopolize an inflorescence whereas others coexist with up to seven other species (Takenaka, 2006; Takenaka et al., 2006; Toda and Lakim, 2011).

There is high host specificity within the *Colocasiomyia crisata* species group; the flies reproduce exclusively on inflorescences of the genera *Colocasia*, *Alocasia* and *Steudnera* (Araceae) and each fly species is usually associated with just one or two host species (Carson and Okada, 1980; Miyake and Yafuso, 2005; Takenaka, 2006; Toda and Lakim, 2011).

*Alocasia macrorrhizos* (L.) G. Don is the most widely distributed species of *Alocasia* and is visited by different *Colocasiomyia* species in different regions (Okada and Yafuso, 1989; Sultana et al., 2006; Takenaka, 2006; Toda and Lakim, 2011; Yafuso and Okada, 1990). In 1999, two previously undescribed *Colocasiomyia* species were discovered to be coexisting in
the inflorescences of *A. macrorrhizos* in Sabah, Malaysian Borneo (Toda and Lakim, 2011). In 2004, we revisited the same locality with the aims to determine (i) the flowering ecology of the host plant, (ii) the association between flowering events and fly behaviour, (iii) the importance of the flies as pollinators in comparison with other flower visitors and (iv) the reproductive habits of the flies. Finally, we characterise the unique pollination mutualism between Araceae plants and *Colocasiomyia* flies.

**MATERIALS AND METHODS**

*Host plant*

*Alocasia macrorrhizos* is a perennial herb with a thick erect stem and is found along the edges and open gaps of forests as well as along the roadside across Indo-Malesia and Oceania (Fig. 1). Its natural origin is unknown but is likely to be within Southeast Asia. In some Pacific regions, it is traditionally cultivated as a starchy stem crop and has a long history of human utilization and dispersal (Hay, 1998; Hay et al., 1995; Hay and Wise, 1991). Synflorescences (paired inflorescences) bloom one by one as a pair per leaf and such pairs occur alongside each other in a continuous sequence (Fig. 2). From the bottom to the top the spadix has a female zone, a sterile mid-zone, a male zone and a sterile appendix (Fig. 3). A cream-coloured spathe covers the spadix and forms a floral chamber, which constricts around the sterile mid-zone. The upper spathe and spadix begin to decay soon after pollen releasing phase (Fig. 1O), wither and then drop off (Fig. 1Q). An infructescence of an old and big plant usually contains more than 100 fruits or berries and each fruit contains one to five seeds. Voucher specimens were deposited in the herbaria of the Institute for Tropical Biology and Conservation, Universiti Malaysia Sabah (specimen KT-303) and the Headquarters of Kinabalu Park (specimens KT-303 and 305).
**Study site**

Sampling, observations and experiments were conducted at a vacant lot in Kota Kinabalu City (5°58′32″N, 116°04′29″E, 15 m above sea level), Sabah, Borneo, Malaysia. Patches of *A. macrorrhizos* are common in open areas throughout the city and the plant blooms perpetually. The patch at our study site was established by the landowner several years previously and is pruned for control approximately every three months.

**Ecological observation and sampling**

We observed the inflorescences of 23 ramets and the insect visitors between 1000 and 1400 daily from 14 July to 1 August 2004 and conducted continuous observation from 0440 to 2220 from 22 to 25 July 2004.

We defined the stages in the flowering and fruiting sequence as follows:

- **Stage I**—emergence from the leaf sheath to just before opening of the spathe (Fig. 1B, 1C);
- **Stage II**—pollen-receiving phase (i.e. female phase) for a one-day period, with open spathe and *Colocasiomyia* flies (Fig. 1E–1G);
- **Stage III**—pollen-releasing phase (i.e. male phase), during which the flies are covered in pollen (Fig. 1J, 1K);
- **Stage IV**—early swelling of the infructescence and initial decay of the upper spathe and spadix, from just after the pollen-release phase (Fig. 1O);
- **Stage V**—late swelling of the infructescence, from withering and fall of the upper spathe and spadix until dehiscence of the lower spathe (Fig. 1R); and
- **Stage VI**—appearance of red mature fruits in the infructescence after dehiscence (Fig. 1U, 1V).

In total, 288 inflorescences and infructescences of the 23 ramets were marked and the
flowering and fruiting sequence was followed for each.

**Flower visitors**

*Colocasiomyia* flies often gathered in the upper (Fig. 1E, 1F) and lower (Fig. 1G) parts, separately, of the spathe chamber (Fig. 3). We collected visiting insects from the upper part by direct aspiration. Thereafter, the entire inflorescence was covered with a plastic bag and detached from the plant. Insects remaining in the lower part were aspirated from inside the bag. The insects were initially preserved in Kahle’s fluid (distilled water, 95% ethanol, formaldehyde and glacial acetic acid in a 28:17:6:2 ratio) and later stored in 70% ethanol. *Colocasiomyia* flies were identified at the species level, and other insects were identified at the genus, family or order level. The *Colocasiomyia* species composition and sex ratio in the upper and the lower parts of each inflorescence were compared using Fisher’s exact test with JMP 7 software (SAS Institute, USA). Voucher specimens of all insect taxa have been kept for future reference.

**Immature stages of Colocasiomyia in the infructescences**

We examined five inflorescences and 22 infructescences at different stages in the laboratory. The spadix was cut into nine sections (Fig. 3); the male and female zones were divided into three sections of equal length and the sterile mid-zone was divided into two sections of equal length, respectively. The total number of immature individuals and adults, awaiting eclosion in puparia, was counted under a stereomicroscope for each section and species. The diagnostic morphological characteristics of immatures of the two taxonomically undescribed *Colocasiomyia* species and another diptera species were listed in Supplemental File 1. The developmental stages were identified as follows: egg, first-instar larva, second- and third-instar larva, pupa, puparium before eclosion and empty puparium after eclosion. The
distribution data on eggs and first-instar larvae were summed for each *Colocasiomyia* species (Supplemental File 2) and then compared by using the chi-square test with JMP 7 software.

**Age estimation of Colocasiomyia immatures by the infructescences**

The *Colocasiomyia* flies laid eggs on an inflorescence in Stage II, which lasts for only one day. Therefore, we could estimate the age of the fly progenies that develop in the inflorescence as being equal to the period following Stage II; we recorded this period from the beginning of Stage II to the harvesting date of each infructescence.

For developing infructescences in Stages IV and V, we estimated the time since egg deposition by tracing back the infructescence sequence because the inflorescences bloom at a regular interval. We named the interval between the first and the second inflorescences within a synflorescence as the ‘intra-synflorescence interval’ and the interval between the second inflorescence of a preceding (i.e. older) synflorescence and the first inflorescence of the following (i.e. younger) synflorescence as the ‘inter-synflorescence interval’ (Fig. 2). Then, $P_i$, which was defined as the number of days from the beginning of Stage II to the harvesting of infructescence $i$, is calculated as follows:

$$P_i = (D_h - D_j) + (A_{\text{intra}} \times \text{the number of intra-synflorescence intervals between infructescences } i \text{ and } j) + (A_{\text{inter}} \times \text{the number of inter-synflorescence intervals between infructescences } i \text{ and } j)$$  
(Formula 1)

where infructescence $j$ is the first inflorescence, within a ramet, that opened the spathe after we started the observation; $D_j$ is the date of the beginning of Stage II of inflorescence $j$; $D_h$ is the harvesting date of infructescence $i$; $A_{\text{intra}}$ is the average duration of the intra-synflorescence intervals; and $A_{\text{inter}}$ is the average duration of the inter-synflorescence
intervals.

To investigate the time of *Colocasiomyia* adult emergence from the infructescences, we covered some infructescences before dehiscence with nylon stockings.

**Field pollination experiments**

Three inflorescences for each of 13 ramets were treated to examine the importance of each insect group as pollinators: one inflorescence was tagged (open-pollinated control), another was bagged with a coarse mesh (2 mm grid, soap bag) to exclude large insect visitors (Fig. 1D) and the third was bagged with a fine-mesh nylon stocking to exclude *Colocasiomyia* and all other insect visitors. The treatments were performed from 14 to 26 July 2004 and the bags were removed after flowering. Usually, *Alocasia* fruits mature in 45–90 days. However, we collected the developing infructescences before maturation, on 12 August 2004, and preserved them in Kahle’s fluid, because of the schedule of the field trip.

The numbers of developing and undeveloped fruits were counted for each infructescence. We discriminated between the developing and the undeveloped fruits by examining whether a fruit had one or more developing seeds, using a stereomicroscope. The fruit fertility rate was estimated by dividing the number of developing fruits by the total number of fruits (both developing and undeveloped) for each infructescence. We compared the fertility rates between the control treatment and the other treatments by the Wilcoxon matched-pairs signed-ranks test using JMP 7.

**RESULTS**

**Flowering and fructing sequence**

The mean total number of inflorescences and infructescences per ramet was 12.5 (s.d. = 2.6, maximum = 18, \( n = 23 \)). The mean durations of the stages in the flowering and fructing
sequence were 1.3 days in Stage II (s.d. = 0.4, n = 32), 1.0 day in Stage III (s.d. = 0.0, n = 29) and 5.0 days in Stage IV (s.d. = 1.0, n = 14). None of the flowering periods from Stage II (female phase) to Stage III (male phase) overlapped between inflorescences within a ramet, presumably to avoid geitonogamy. The mean intra-synflorescence interval ($A_{\text{intra}}$) was 4.1 days (s.d. = 0.7, n = 18) and the mean inter-synflorescence interval ($A_{\text{inter}}$) was 8.3 days (s.d. = 1.2, n = 16). Thus, Formula 1 becomes

$$P_i = (D_h - D_j) + (4.1 \times \text{number of intra-synflorescence intervals between infructescences } i \text{ and } j) + (8.3 \times \text{number of inter-synflorescence intervals between infructescences } i \text{ and } j).$$

(Formula 2)

Using Formula 2, we estimated the ages of the immatures of the *Colocasiomyia* flies within the harvested infructescences (Supplemental File 3).

*Flowering events and behaviours of the Colocasiomyia flies*

Two fly species, *Colocasiomyia* sp.1 aff. *sulawesiana* and *Colocasiomyia* sp.2 aff. *sulawesiana* (hereafter abbreviated as Sp. 1 and Sp. 2, respectively), were the predominant visitors (Supplemental File 3). We often observed many pollen grains attached on bodies of collected flies of both Sp. 1 and Sp. 2 (Fig. 1K). Their behaviours corresponded well to the flowering events as follows.

Early on the first morning, the spathe opened and presented a narrow slit, and the floral chamber emitted a strong odour. The *Colocasiomyia* flies were attracted to the spathe and then entered through the slit around sunrise, between 0550 and 0620 (Fig. 1D). The flies remained in the chamber for one day to feed, mate (Fig. 1E–1G) and deposit eggs between pistils or staminodes (Fig. 1H, 1I). Some flies often congregated inside the upper spathe
chamber (Fig. 1E, 1F), whereas others swarmed to the lower part (Fig. 1G). The species composition of the collected flies differed significantly between the upper and the lower parts in all the inflorescences examined: Sp. 1 was more abundant in the lower part and Sp. 2 was in the upper part, respectively (Supplemental File 3). The sex ratio, however, was not significantly different between the upper and the lower parts for either species, except for Sp. 2 in one inflorescence (Supplemental File 3).

After the one-day female phase (Stage II), the male phase (Stage III) began with pollen release before 0440 on the second morning. Pollen release continued till 0600, when the constriction of the spathe (Fig. 3) began to tighten (Fig. 1J). Colocasiomyia flies escaped from the lower chamber as it closed by crawling up the spadix in a shower of pollen (Fig. 1J). The flies flew away, presumably to enter female-phase inflorescences on the nearby ramets. These floral events occurred every morning, with the Colocasiomyia flies migrating from one inflorescence to another and staying overnight in each temporary habitat.

**Other visitors to A. macrorrhizos inflorescences**

Many stingless bees—*Trigona (Tetragonula) fuscobalteata* Cameron, 1908 and *Trigona (Tetragonula) laeviceps* Smith, 1857 (Hymenoptera: Apidae)—collected pollen before sunrise and in the daytime (Fig. 1M). Many of them visited the inflorescences only after pollen release, when the female zone was no longer accessible due to spathe closure. They investigated female-phase inflorescences at times and rarely entered the lower part of the spathe chamber, where the pistils are located.

In the morning between 0440 and 0700, honeybees (*Apis cerana* Fabricius, 1793; Hymenoptera: Apidae) also visited the inflorescences that were releasing pollen (Fig. 1L).

Adult flies, which were tentatively identified as *?Atherigona* sp. (Diptera: Muscidae), repeatedly visited the inflorescences and young infructescences of Stages II–IV in the
daytime (Supplemental File 3, Fig. 1N), but not during pollen release in the early morning. They walked around the spadices that had been covered with pollen grains for a couple of days after pollen release and moved to the lower part of female-phase inflorescences sometimes. They used the inflorescences as reproductive sites and presumably, as feeding sites (Fig. 1N). Eggs were laid on the sterile mid-zone and male zone (Supplemental File 2). Second- and third-instar larvae of the species fed on decaying tissue of the male zone and appendices (Supplemental File 2). Several pupae were found from the appendices to the male zones (Supplemental File 2). Some larvae and pupae were collected together with the decaying appendices and reared on the appendices at ambient room temperature until they became identifiable adults.

Two adult females of *Neurochaeta mcalpinei* Woodley, 1982 (Diptera: Neurochaetidae) and one adult female of *Stenomicra (Podocera) australis* Malloch, 1927 (Diptera: Periscelididae) were collected (Supplemental File 3), but their behaviours were not observed. Parasitoid wasps (Hymenoptera) were found in two inflorescences (Supplemental File 3), and several individuals were found developing in *Colocasiomyia* puparia (Supplemental File 4). Earwigs belonging to the species *Chelisoches morio* (Fabricius, 1775) (Dermaptera: Chelisochidae), were often present at the bottom of the spathe chambers. One rove beetle (Coleoptera: Staphylinidae) and one collembolan were also collected (Supplemental File 3).

**Field pollination experiments**

The fruit fertility rate of the control, bee-excluded and *Colocasiomyia*-excluded inflorescences was $0.89 \pm 0.13$ (mean $\pm$ s.d., $n = 13$), $0.85 \pm 0.19$ ($n = 13$) and $0.002 \pm 0.007$ ($n = 13$), respectively (Fig. 4). The inflorescences bagged with the coarse mesh were visited by *Colocasiomyia* flies but not larger insects, and produced as many fruits as the open-pollinated controls ($Z = -5.5$, $P = 0.367$, one-tailed Wilcoxon test for comparisons with
the control) (Fig. 4). However, excluding *Colocasiomyia* and all or most of the other insects with the fine mesh reduced the seed production almost completely ($Z = -45.5, P < 0.0001$); only one of 13 inflorescences produced three fruits, although each inflorescence possessed more than 100 pistils.

**Immature stages of Colocasiomyia on the host inflorescences**

Eggs and young larvae of Sp. 1 and Sp. 2 were found on the pistils of the female zone and the staminodes of the sterile mid-zone (Supplemental File 2 and Fig. 1H, 1I). Their distributions in the infructescences were bimodal, with a larger peak in the lower part of the female zone and a smaller peak in the lower part of the sterile region (subtotal of Supplemental File 2), and significantly different between the species ($\chi^2 = 407.5, df = 8, P < 0.0001$; the numbers of individuals on Sections 4 and 5 were summed because of the small value).

Larvae of both *Colocasiomyia* species were found in the lower part of the male zone and throughout the female zone in Stage III. The larvae were subsequently found only in the female zone during Stage IV (Supplemental File 2), when the infructescence is bathed in its own secretion (Fig. 1P). The *Colocasiomyia* larvae developed in this secretion and seemed to feed on something from it (e.g. the secretion itself or bacteria or yeasts proliferating in the secretion). During the later part of Stage V, when the inside of the infructescences became drier (Fig. 1R), the larvae pupated in spaces between the fruits and the inner side of the spathe tube, especially where cavities formed around aborted fruits (Fig. 1S). The distributions of the larvae and puparia in the infructescences after Stage III were not significantly different between the species (data not shown).

We observed new adults of Sp. 1 and Sp. 2 emerging from a matured infructescence just after dehiscence. On the first day of dehiscence, the surrounding spathe tube began to open at the top (Fig. 1U), and a dozen new adults were observed in the stocking cover. On the second
day, the spathe split further and more than 100 new adults emerged. Only three uneclosed pupae were found among the abundant empty puparia (Supplemental File 4, ramet 222–inflorescence 1). The age of both the flies and the infructescence, estimated by using Formula 2, was approximately 74 days (Supplemental File 4, ramet 222–inflorescence 1).

According to the estimated ages of the infructescences, the youngest infructescence in which empty puparia were found was 62.3 days after anthesis (ramet 224–infructescence 3 at Stage VI, for both species) and the oldest one in which living puparia with a developed adult body were found was 89.0 days (ramet 232–infructescence 1 at Stage V, for both species).

**DISCUSSION**

*Flowering events and behaviour of the Colocasiomyia flies*

The flowering events and the behaviours of the *Colocasiomyia* flies were well synchronised. The *Colocasiomyia* flies pollinated their host in a sophisticated and effective manner, as has been reported for other pollination mutualisms between *Colocasiomyia* flies and their species-specific host plants in the Araceae (Carson and Okada, 1980; Kramadibrata and Hambali, 1983; Mori and Okada, 2001; Takenaka, 2006; Takenaka et al., 2006; see also Cleghorn, 1913; Toda and Okada, 1983; Yafuso, 1993).

Ivancic *et al.* (2005) studied inflorescence heating (thermogenesis) of *A. macrorrhizos* in Vanuatu and reported that the average maximum temperature ± s.e.m. of the appendix reached 43.9 ± 0.6 °C (*n* = 59 inflorescences; the average ambient air temperature was 22.4 ± 0.5 °C) between 0545 and 0645 on the first morning of the anthesis. The function of the inflorescence thermogenesis in Araceae is generally agreed to be to volatilise odour compounds for pollinator attraction (Mayo et al., 1997). The time of flower visiting by the *Colocasiomyia* flies observed in the present study well corresponded to the time of the peak temperature reported in Ivancic *et al.* (2005). Seymour *et al.* (2003b) reported that floral heat
of *Philodendron solimoensis* (Araceae) in French Guiana serves as a direct reward for a pollinating large scarab beetle, *Cyclocephala colasi* (Coleoptera: Scarabaeidae). It is unclear, however, whether floral heat of *A. macrorrhizos* serves as a direct reward for the *Colocasiomyia* flies.

**Other visitors to the inflorescences**

The common visitors were stingless bees, but their role as pollinators (if any) seems to be minor, because they rarely accessed the pistils. Honeybees visited the inflorescences only when pollen was actively released in the early morning. Even if they were able to access the female-phase inflorescences, they are too large to enter the lower part of the spathe chamber and thus would not contribute to cross-pollination. *?Atherigona* species often visited the inflorescences but not strictly in synchrony with the flowering events. The species does not seem to serve as an effective pollinator.

Two female flies of *Neurochaeta mcalpinei* were collected only once (Supplemental File 3). McAlpine (1987) suggested that the members of *Neurochaeta* have a morphology (flattened body shape) and behaviour (running backwards) that appear to be adapted to host plants in the families Araceae, Musaceae, Pandanaceae and Zingiberaceae. These plants shelter the flies in narrow cavities: the axils of bracts, the sheath hollows of petioles and spathe cavities. One female individual belonging to the species *Stenomicra australis* was also collected (Supplemental File 3). McAlpine (1987) noted that flies belonging to *Stenomicra* often share the same habitat as *Neurochaeta*. All of these flies may utilise *Alocasia* plants as preferred hosts, but there is no evidence that they are effective pollinators.

Earwigs (*Chelisoches morio*) were often seen at the bottom of the spathe chambers and appeared to prey on the eggs and larvae of *Colocasiomyia* and *?Atherigona* species. Terry (1905) reported the predatory habits of *C. morio* on leafhoppers; this earwig is omnivorous.
but seems to prefer an insect-based diet (Tenbrink and Hara, 2006). Kamimura (2001) observed the nymphs of another earwig species, *Forficula hiromasai* Nishikawa, 1970 (Dermaptera: Forficulidae), on inflorescences of *Arisaema serratum* and *Arisaema thunbergii* (Araceae) and found pollen grains of these plants in the nymph guts. Thus, earwigs may depend on aroid hosts for food (pollen and prey) and habitat (spathe chamber) throughout their life cycle. From our observations, however, there is no indication that earwigs contribute to cross-pollination.

Rove beetles (Coleoptera: Staphylinidae) also visited the inflorescences (Supplemental File 3). They were often seen at the inflorescences of *Alocasia, Colocasia* and particularly, *Schismatoglottis* in the family Araceae (KTT and MJT, unpubl. res.), presumably targeting eggs and larvae of *Colocasiomyia* or other insects breeding at the site. In this study, the number of rove beetles was very small and they did not seem to function as pollinators.

**Pollination by Colocasiomyia flies**

The inflorescences bagged with a fine mesh produced almost no fruits (Fig. 4). Ivancic *et al.* (2005) concluded that *A. macrorrhizos* is predominantly self-incompatible. Our bagging experiment with a coarse mesh suggested that the main pollinators could only be small insects that can pass through 2 mm grids. From our observations of the insect visitors and the two bagging experiments, we conclude that the *Colocasiomyia* flies (Sp. 1 and Sp. 2) were the main and possibly only effective pollinators for *A. macrorrhizos* at the study site.

**Reproductive traits of Colocasiomyia species**

Usually, a pistilicolous and a stamenicolous species of the *Colocasiomyia cristata* group are found within the same inflorescence of *Colocasia* or *Alocasia* but show different traits in niche choice, morphology and life history (Okada, 1986). Our results contradicted this pattern.
because both species reproduced in the female zone and showed only slight differences.

**Niche choice for oviposition.** Both Sp. 1 and Sp. 2 exhibited the pistilicolous habit of oviposition in the female zone (Supplemental File 2), and subsequent development until adult eclosion occurred within the infructescence (Supplemental File 4). This is the first reported observation of two pistilicolous *Colocasiomyia* species coexisting within a single inflorescence and infructescence. However, some level of niche segregation was still observed because Sp. 2 adults congregated mainly in the upper part of the spathe chamber while those of Sp. 1 gathered in the lower part (Supplemental File 3).

**Morphological characteristics.** Stamenicolous *Colocasiomyia* species generally have narrower ovipositors, which are presumably an adaptation for laying eggs in the narrower spaces between stamens. Pistilicolous species have wider ovipositors and lay their eggs in the wider spaces between pistils. The ovipositor was wide (pistilicolous type) in both species, although Sp. 2 had a longer ovipositor compared with Sp. 1 (Takenaka, 2006).

The following traits of Sp. 2 suggest a stamenicolous tendency: a longer ovipositor, the congregation of adults at the upper part of the spathe chamber and the distinct second peak at the intermediate region of the inflorescence in the egg distribution. The differences between the species may reflect microniche differentiation through reaction to larval food resources or against predators and parasitoid wasps.

**Life history from egg to eclosion.** The life-history traits of both *Colocasiomyia* species reported here strongly indicate intimate adaptation to their host plant. Immatures of stamenicolous *Colocasiomyia* generally leave the inflorescence with decaying tissue when the upper spadix and spathe wither and fall, or ‘pop out’ of the spadix to pupate on the ground.
(a larval behaviour commonly observed in a number of species of Diptera) (Yafuso, 1993), whereas pistilicolous species spend the whole period from egg to eclosion on their host plants. For Sp. 1 and Sp. 2, the time from oviposition to eclosion was estimated to be less than 62 days in the shortest example and more than 89 days in the longest example. These periods are remarkably long when compared with those of other Colocasiomyia species. The respective periods under laboratory conditions are approximately 18 and 30 days for C. stamenicola and C. pistilicola (Carson and Okada, 1980), and two and three weeks for C. alocasiae and C. xenalocasiae (Yafuso, 1999). In the field, C. alocasiae and C. xenalocasiae seem to require more time (KTT, MJT and M. Yafuso, unpubl. res.), but the periods are still considerably shorter than those reported here. A hole often develops at the apex of Alocasia infructescences before dehiscence, because the top part of the spindle of the spadix decays as the infructescence ripens. In contrast, infructescences of A. macrorrhizos are completely sealed (Fig. 1T) until dehiscence (Fig. 1U), because the spindle of the spadix remains. Sp. 1 and Sp. 2 may find it difficult to exit before the infructescences dehisce, possibly resulting in the protracted egg-to-eclosion period. Thus, the floral life history of the hosts may explain the variation in the length of the egg-to-eclosion period of Colocasiomyia species.

In both species, the protracted egg-to-eclosion period was largely due to the relatively long developmental periods in the third instar and pupa as well as the prolonged residence in the puparium, after metamorphosis to the adult form (Appendices 3, 4). These flies seem to adjust their developmental stages to the conditions of their host plant infructescence. The soakage in the infructescences decreases as the fruits ripen and the decreasing wetness may be the cue for the larvae to pupate. Adults that have completed metamorphosis within puparia then appear to wait for dehiscence of the infructescence, before leaving the puparium and the plant.

Most adult flies emerged at the same time when the host infructescence dehisced. Rapid
and simultaneous departure after dehiscence may be necessary because the mature fruit are frequently eaten by animals (possibly birds or squirrels) after dehiscence and sometimes before dehiscence (Fig. 1W). Exposed puparia (Fig. 1V) may also be attractive for other insects such as ants, which are often observed on the plant. Changes in certain physical conditions such as the light intensity or air composition inside the spathe tube may be cues for eclosion. Experimental investigations such as making a hole in the spathe tube before dehiscence might help to reveal such cues for the eclosion of Colocasiomyia species.

**Characteristics of this pollination mutualism**

The two Colocasiomyia species were host specific pollinators and depended their reproduction on inflorescence and infructescence of their host plant. In this sense, this pollination system is comparable to obligate pollination mutualisms in which fig–fig wasp and yucca–yucca moth systems are the best-documented examples (Janzen, 1979; Pellmyr et al., 1996; Powell, 1992; Wiebes, 1979). In typical obligate pollination mutualisms, plants have a dilemma to sacrifice ovules or developing seeds in return for pollination. In contrast, larvae of the Colocasiomyia species do not damage fruits, therefore the host plant does not have the dilemma.

Moreover, no adults of both the Colocasiomyia species were caught by intensive net sweeping either around the host plants or at other adjacent sites. This suggests that both fly species spend most of their lifetime within the inflorescences of the single host species except for brief periods of adult migration. It also suggests that Alocasia macrorrhizos provides most of necessary resources for its pollinators to survive: food for immatures and adults, place for reproduction (mating and oviposition sites) and shelter for egg, larvae and adults. Year-round availability of inflorescences may secure continuous reproduction of the flies. However, many of Araceae host plants of other Colocasiomyia flies have limited flowering seasons. How
Colocasiomyia flies survive when host flowers are not available has been unknown.

More than 70 Alocasia species, about 10 Colocasia species and about 7 Steudnera species are distributed in the Oriental and Papuan regions (Mayo et al., 1997). Given the high host specificity of the Colocasiomyia cristata group (Sultana et al., 2006), there seems to be many undiscovered pollination mutualisms between Araceae plants and Colocasiomyia flies. Comparative studies of these presumable pollination systems would shed more light on the evolution of the highly intimate pollination mutualisms (Takano et al., 2011).
ACKNOWLEDGEMENTS

We thank Mitsuru Hotta and Peter C. Boyce for identification of A. macrorrhizos; Teruyoshi Nagamitsu (Trigona and Apis), Fabian Haas and Masaru Nishikawa (Chelisoches) for insect identification; Masaaki Suwa for suggestions on ?Atherigona. Neurochaeta and Stenomicra species were officially identified by Masahiro Sueyoshi of the Kyusyu Research Centre, Forestry and Forest Products Research Institute (18 FFPRI No. 665). We also thank members of the Institute for Tropical Biology and Conservation, Universiti Malaysia Sabah; Sabah Parks; the Bornean Biodiversity and Ecosystems Conservation, JICA; Mr. Josef, the owner of the vacant lot in Kota Kinabalu; and the Economic Planning Unit of the Malaysian Government for permission and arrangements to conduct the field research. We are grateful to Naoko Takano, Peter J. Matthews, Daniel Impoinvil and Celeste Donato for their critical reading of the manuscript. This work was supported by Grants-in-Aid for Scientific Research from the Japan Society for the Promotion of Science [15255006] and the 21st Century Centre of Excellence Program [E-01] of the Ministry of Education, Culture, Sports, Science and Technology, Japan.
REFERENCES


Kramadibrata, K. and Hambali, G.G. (1983) [The roles of some insects in pollination of *Colocasia esculenta* var. *esculenta* and *C. gigantea*]. Berita Biologi, 2(1), 143-146 (in Indonesian).


Yafuso, M. (1999) Larval performance and niche separation between two synhospitalic species of *Colocasiomyia* (Diptera, Drosophilidae). Memoirs of the Faculty of
Science, Kyoto University. Series of Biology, **16**, 119-126.
FIGURE LEGENDS

**Fig. 1.** Flowering and fruiting sequences of *Alocasia macrorrhizos* associated with *Colocasiomyia* flies and other flower visitors in Kota Kinabalu. (A) A ramet of *A. macrorrhizos*. (B) A bud of a synflorescence (paired inflorescences). (C) Pairs of buds (Stage I) and developing infructescences (Stages IV and V). (D) *Colocasiomyia* flies just arrived on a female-phase inflorescence (Stage II) in the morning twilight. (E, F) *Colocasiomyia* flies gathered in the upper and (G) the lower parts (dissected spathe tube) of the spathe chamber. Eggs of (H) *Colocasiomyia* sp.1 aff. *sulawesiana* and (I) *Colocasiomyia* sp.2 aff. *sulawesiana* laid in the spaces between pistillate flowers. (J, K, L) An inflorescence releasing pollen (Stage III) early in the morning. (J, K) *Colocasiomyia* flies crawling up the spadix being dusted with pollen grains and (L) honeybees collecting pollen. (M) Stingless bees collecting pollen deposited on the upper spathe chamber after pollen release (Stages III and IV). (N) *Atherigona* species on an inflorescence (Stage IV). (O) Young infructescence at Stage IV. (P) A dissected infructescence filled with secretion, in which *Colocasiomyia* larvae developed. (Q) Developing infructescences at Stage V. (R, S) A dissected infructescence with *Colocasiomyia* pupae in the spaces between fruits and the spathe tube. (T) The top of an infructescence tightly enclosed with the spindle and the spathe tube (dissected). (U) An infructescence starting to dehisce at Stage VI. (V) A dissected infructescence with red mature fruits and *Colocasiomyia* puparia. (W) Remains of an infructescence several days after dehiscence.

**Fig. 2.** Schematic of the flowering sequence within a ramet (dashed arrow, left) and estimation of the period from the day of spathe opening to the day of collection of the infructescence. In this example, Formula 1 becomes $P_i = (D_h - D_j) + (4.1 \times 1) + (8.3 \times 2) = (D_h - D_j) + 20.7$. $P_i$ was calculated in this manner for each collected infructescence to
estimate the age of the *Colocasiomyia* immatures inside.

**Fig. 3.** Schematic of a spadix (left), an inflorescence with the lower spathe removed (centre) and an intact inflorescence (right) of *Alocasia macrorrhizos*. The numbered sections correspond to those in Supplemental File 2. Scale bar = 10 cm. Interestingly, the appendix of some inflorescences became pinkish at the end of flowering.

**Fig. 4.** Comparison of the fruit fertility rate among the open-pollinated control, bagging with coarse mesh (bee exclusion) and bagging with fine mesh (*Colocasiomyia* exclusion) treatments (*n* = 13 for each treatment). The bold horizontal line shows the median fertility rate. The bottom and top of each box show the 25th and 75th percentiles (i.e. the first and the third quartiles), respectively. The vertical dashed lines are either the maximum value or 1.5 times the interquartile range of the data, whichever is smaller. Points more than 1.5 times the interquartile range above the third quartile and those more than 1.5 times the interquartile range below the first quartile are plotted individually. The *P* values were obtained by one-tailed Wilcoxon test in each comparison with the control.
Flowering and fruiting sequences of *Alocasia macrorrhizos* associated with *Colocasiomyia* flies and other flower visitors in Kota Kinabalu. (A) A ramet of *A. macrorrhizos*. (B) A bud of a synflorescence (paired inflorescences). (C) Pairs of buds (Stage I) and developing infructescences (Stages IV and V). (D) *Colocasiomyia* flies just arrived on a female-phase inflorescence (Stage II) in the morning twilight. (E, F) *Colocasiomyia* flies gathered in the upper and (G) the lower parts (dissected spathe tube) of the spathe chamber. Eggs of (H) *Colocasiomyia* sp.1 aff. *sulawesiana* and (I) *Colocasiomyia* sp.2 aff. *sulawesiana* laid in the spaces between pistillate flowers. (J, K, L) An inflorescence releasing pollen (Stage III) early in the morning. (J, K) *Colocasiomyia* flies crawling up the spadix being dusted with pollen grains and (L) honeybees collecting pollen. (M) Stingless bees collecting pollen deposited on the upper spathe chamber after pollen release (Stages III and IV). (N) *Atherigona* species on an inflorescence (Stage IV). (O) Young infructescence at Stage IV. (P) A dissected infructescence filled with secretion, in which *Colocasiomyia* larvae developed. (Q) Developing infructescences at Stage V. (R, S) A dissected infructescence with *Colocasiomyia* pupae in the spaces between fruits and the spathe tube. (T) The top of an infructescence tightly enclosed with the spindle and the spathe tube (dissected). (U) An infructescence starting to dehisce at Stage VI. (V) A dissected infructescence with red mature fruits and *Colocasiomyia* puparia. (W) Remains of an infructescence several days after dehiscence (to be continued).

133x84mm (300 x 300 DPI)
Fig. 1O-W (reduced-size, continued)
(Takano et al.)

Continued.
226x397mm (300 x 300 DPI)
Schematic of the flowering sequence within a ramet (dashed arrow, left) and estimation of the period from the day of spathe opening to the day of collection of the infructescence. In this example, Formula 1 becomes

$$P_i = (D_h - D_j) + (4.1 \times 1) + (8.3 \times 2) = (D_h - D_j) + 20.7.$$  

$P_i$ was calculated in this manner for each collected infructescence to estimate the age of the *Colocasiomyia* immatures inside.

Fig. 2.  
(Takano et al.)

81x47mm (300 x 300 DPI)
Schematic of a spadix (left), an inflorescence with the lower spathe removed (centre) and an intact inflorescence (right) of *Alocasia macrorrhizos*. The numbered sections correspond to those in Supplemental File 2. Scale bar = 10 cm. Interestingly, the appendix of some inflorescences became pinkish at the end of flowering.

111x155mm (300 x 300 DPI)
Comparison of the fruit fertility rate among the open-pollinated control, bagging with coarse mesh (bee exclusion) and bagging with fine mesh (Colocasiomyia exclusion) treatments (n = 13 for each treatment). The bold horizontal line shows the median fertility rate. The bottom and top of each box show the 25th and 75th percentiles (i.e. the first and the third quartiles), respectively. The vertical dashed lines are either the maximum value or 1.5 times the interquartile range of the data, whichever is smaller. Points more than 1.5 times the interquartile range above the third quartile and those more than 1.5 times the interquartile range below the first quartile are plotted individually. The P values were obtained by one-tailed Wilcoxon test in each comparison with the control.
### Supplemental File 1. Diagnostic morphological characteristics of immatures of *Colocasiomyia* and *Atherigona* species collected from *Alocasia macrorrhizos* inflorescences and infructescences in Kota Kinabalu, Sabah, Borneo, Malaysia.

<table>
<thead>
<tr>
<th>Immature stage</th>
<th>Species</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Egg</td>
<td>Sp. 1</td>
<td>Larger than Sp. 2; relatively straight (gently curved in Sp. 2) along the axis, gently convex (not convex and narrower) ventrally. Surface mat (glossy) white; terminal disk spherical (tabular) with a short filament. Hatching earlier than Sp. 2, from the date of oviposition to the next day.</td>
</tr>
<tr>
<td></td>
<td>Sp. 2</td>
<td>Smallest among the three species; hatching a half to one day later than Sp. 1.</td>
</tr>
<tr>
<td></td>
<td>?<em>Atherigona</em> sp.</td>
<td>Biggest among the three species; surface with two conspicuous cristae along the axis, mat white; afloat on water and hardly sink; hatching about concurrently with or later than Sp. 2.</td>
</tr>
<tr>
<td>First instar larva</td>
<td>Sp. 1</td>
<td>Body more slender than ?<em>Atherigona</em> sp., obviously larger than Sp. 2 at hatching. Caudal abdominal segments elongated, with posterior spiracles with long divergent stalks; spicules on body surface stouter and hooked more strongly than in Sp. 2; anterior spiracle almost retracted into the body. Wiggling in the water.</td>
</tr>
<tr>
<td></td>
<td>Sp. 2</td>
<td>General appearance resembling Sp. 1; spicules on body surface slightly weaker; mouth hook thinner but curving more strongly than in Sp. 1.</td>
</tr>
<tr>
<td></td>
<td>?<em>Atherigona</em> sp.</td>
<td>Somewhat roundish overall; mouth hook stout and short; caudal abdominal segment terminated abruptly without elongation; posterior spiracles short and separated each other.</td>
</tr>
<tr>
<td>Puparium</td>
<td>Sp. 1</td>
<td>Generally larger Sp. 2; puparial surface glossy and darker (mat and pale in Sp. 2). Anterior spiracles short, but the apical end of the stalk rather distinctive (ambiguous) and the spiracles looking ternate often (not often). Relative length of circular opening of puparium compared to puparial size relatively larger (smaller). Divergent stalks of posterior spiracles as long as or longer (shorter) than one third of that of caudal abdominal segments.</td>
</tr>
<tr>
<td></td>
<td>Sp. 2</td>
<td>Spicules on surface weaker and sparser than in Sp. 1.</td>
</tr>
<tr>
<td></td>
<td>?<em>Atherigona</em> sp.</td>
<td>Much larger than those of <em>Colocasiomyia</em> puparium.</td>
</tr>
</tbody>
</table>
### Supplemental File 2

The numbers of individuals in each developmental stage found in different parts of each inflorescence (stage II and III) or young infructescence (stage IV) of *Alocasia macrorrhizos* in Kota Kinabalu.

<table>
<thead>
<tr>
<th>Flowering / fruiting stage</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>Sub total*²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Host ramet No.</td>
<td>201</td>
<td>234</td>
<td>241</td>
<td>242</td>
</tr>
<tr>
<td>Inflorescence / infructescence No.</td>
<td>5</td>
<td>8</td>
<td>11</td>
<td>9</td>
</tr>
<tr>
<td>Recoded number of days from anthesis</td>
<td>0-1</td>
<td>0-1</td>
<td>0-1</td>
<td>0-1</td>
</tr>
</tbody>
</table>

*Sp. 1 Section in a inflorescence or infructescence*  
<table>
<thead>
<tr>
<th>Sp. 1</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>5</td>
<td>10</td>
<td>1</td>
<td>2</td>
<td>5</td>
<td>1</td>
</tr>
</tbody>
</table>

*Sp. 2 Section in a inflorescence or infructescence*  
<table>
<thead>
<tr>
<th>Sp. 2</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3</td>
<td>5</td>
<td>26</td>
<td>2</td>
<td>16</td>
<td>25</td>
</tr>
</tbody>
</table>

*?Atherigona sp. Section in a inflorescence or infructescence*  
<table>
<thead>
<tr>
<th>?Atherigona sp.</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>1</td>
<td>2</td>
<td>7</td>
<td>1</td>
<td>5</td>
</tr>
</tbody>
</table>

Notes: *n*, eggs; *n*₁, first instar larvae; *n*₂, second and third instar larvae; and (*n*₃), pupae. *The numbered sections are shown in Fig. 3. *²Significantly different between the *Colocasiomyia* species by chi-squared test (*χ² = 407.5, df = 8, *P* < 0.0001).
Supplemental File 3. The numbers of adult *Colocasiomyia* flies and other insects collected from each stage-II (female phase) inflorescence of *Alocasia macrorrhizos* at Kota Kinabalu.

<table>
<thead>
<tr>
<th>Host ramet No.</th>
<th>Inflorescence No.</th>
<th>205</th>
<th>202</th>
<th>201</th>
<th>234</th>
<th>241</th>
<th>242</th>
<th>227</th>
<th>308</th>
<th>224</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>U</td>
<td>L</td>
<td>U</td>
<td>L</td>
<td>U</td>
<td>L</td>
<td>U</td>
<td>L</td>
<td>U</td>
</tr>
<tr>
<td>Part in a inflorescence&lt;sup&gt;1&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sp. 1</td>
<td>♀</td>
<td>2</td>
<td>26</td>
<td>2</td>
<td>8</td>
<td>0</td>
<td>22</td>
<td>2</td>
<td>50</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>♂</td>
<td>3</td>
<td>34</td>
<td>0</td>
<td>3</td>
<td>0</td>
<td>19</td>
<td>1</td>
<td>48</td>
<td>21</td>
</tr>
<tr>
<td>Sp. 2</td>
<td>♀</td>
<td>24</td>
<td>0</td>
<td>4</td>
<td>2</td>
<td>13</td>
<td>2</td>
<td>9</td>
<td>1</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>♂</td>
<td>28</td>
<td>7</td>
<td>11</td>
<td>5</td>
<td>13</td>
<td>2</td>
<td>17</td>
<td>2</td>
<td>33</td>
</tr>
<tr>
<td>Difference in species composition&lt;sup&gt;2&lt;/sup&gt;</td>
<td></td>
<td>***</td>
<td>**</td>
<td>***</td>
<td>***</td>
<td>***</td>
<td>*</td>
<td>***</td>
<td>***</td>
<td>***</td>
</tr>
<tr>
<td>Difference in sex ratio&lt;sup&gt;2&lt;/sup&gt;</td>
<td>sp. 1</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td></td>
<td>sp. 2</td>
<td>*</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
</tbody>
</table>

* Neurochaeta mcalpinei* (Diptera: Neurochaetidae)  
♀ 2

* Stenomicra australis* (Diptera: Periscelididae)  
♀ 1

?Atherigona sp. (Diptera: Muscidae)  
1

* Trigona fuscobalteata* (Hymenoptera: Apidae)  
1

Parasitoid wasp (Hymenoptera)  
2 2

Staphylinidae species (Coleoptera)  
1

Collembola species  
1

Notes: <sup>1</sup> U = upper part, L = lower part (Fig. 3).  
<sup>2</sup> Difference in the number of each *Colocasiomyia* species in the upper and the lower parts of each inflorescence by Fisher's exact tests.  
* = P < 0.05, ** = P < 0.01, *** = P < 0.0001, n.s. = not significant.
Supplemental File 4. The numbers of *Colocasiomyia* individuals at different developmental stages on each infructescences (stages V-VI) of *Alocasia macrorrhizos* at Kota Kinabalu.

<table>
<thead>
<tr>
<th>Fruiting stage</th>
<th>V</th>
<th>VI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Host ramet No.</td>
<td>223, 237, 222, 241</td>
<td>225, 230, 228, 236</td>
</tr>
<tr>
<td>Infructescence No.</td>
<td>11^s, 10^s, 6, 6</td>
<td>3, 3, 1, 2, 4, 4</td>
</tr>
<tr>
<td>Recoded (*) or estimated (= P_i) number of days from spathal opening</td>
<td>16^s, 16^s, 47.5, 47.8</td>
<td>60.2, 64.2, 64.2, 74.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sp. 1</th>
<th>Third instar larva</th>
<th>1</th>
<th>10</th>
<th>77</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Puparium with undeveloped adult body</td>
<td>1</td>
<td>16</td>
<td>54</td>
</tr>
<tr>
<td></td>
<td>Puparium with developed adult body (parasited*)</td>
<td>33 (2*)</td>
<td>9</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td>Empty puparium after eclosion</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Dead pupa</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dead adult failed in eclosion</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sp. 2</th>
<th>Third instar larva</th>
<th>5</th>
<th>32</th>
<th>73</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Puparium with undeveloped adult body</td>
<td>2 (14*)</td>
<td>17</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Puparium with developed adult body (parasited*)</td>
<td>(12*)</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Empty puparium after eclosion (parasited*)</td>
<td>(1*)</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>

Note: *^s Used for coarse-meshed bagging experiment.