A place to grow? Host choice and larval performance of *Microplitis similis* in the host *Spodoptera litura*

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Abstract

Host selection is a key stage in the lifecycle of parasitoids, and is critical to both their function in control and to the maintenance of their population. The solitary endoparasitoid *Microplitis similis* (Hymenoptera: Braconidae), is a potential biological control agent of *Spodoptera litura* larvae (Lepidoptera: Noctuidae). In this study, we examined the preference *M. similis* exhibits for different instars of the host, host instar effects on parasitoid development and the weight gain and food consumption of different instars of parasitized larvae. In no-choice tests, parasitization rates were highest in second and early third instar larvae, and no fourth or fifth instar hosts were parasitized. When provided with a choice of first to late third instars host larvae, *M. similis* preferred to parasitize early third instar host larvae (41 %) with a selection coefficient of 0.37. All morphometric features of wasp offspring increased with increasing age of the host at parasitization. A lower proportion of females emerged from first instar larvae than any other instar. Parasitized *S. litura* larvae showed a pronounced reduction in food consumption and weight gain. *Microplitis similis* may have the potential to significantly suppress population growth and the damage caused by *S. litura*.

**Keywords** *Microplitis similis, Spodoptera litura* (Fabricius), Host age preference, Weight gain, Food consumption

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Introduction

The oriental leafworm moth, *Spodoptera litura* (Fabricius) (Lepidoptera: Noctuidae) has a worldwide distribution, can migrate long distances, and has a high reproductive rate (Shad et al. 2012). *Spodoptera litura* is a polyphagous insect pest which has been
recorded from over 100 families and causes extensive damage to tobacco, cotton, soybean and cruciferous vegetables (Tuan et al. 2014). Oriental leafworm was recently reported to cause economic losses of about $44.7 million to soybean in the Kota region of Rajasthan (Dhaliwal et al. 2010). Conventional control of noctuid pests chiefly relies on chemical pesticides, and the frequent application of insecticide has led to the rapid evolution of insecticide resistance in the field (Ahmad et al. 2007; Su et al. 2012).

Biological control is an important component of successful Integrated Pest Management (IPM) programs. The use and enhancement of natural enemies of the insect pests plays an important role in IPM, and parasitoids are considered to be important biological control agents against Lepidopteran pests (Mills 2009; Barzman et al. 2015). He and Liu (2002) list hymenopterous parasitoids of S. litura from China, including 29 primary parasitoids and 11 hyperparasitoids. The genus Microplitis contains 200 species (Ernandez-Triana et al. 2015) and M. similis Lyle (Hymenoptera: Braconidae) is a solitary endoparasitoid, with hosts including Agrotis ypsilon (Rottemberg), Spodoptera exigua (Hübner) and S. litura (Fabricius) (Shepard and Barrion 1998; Ranjith et al. 2015). Microplitis similis was first recorded in India (Wilkinson 1930) and identified as a vector of Heliothis virescens ascovirus 3h only recently (Li et al. 2016).

Host species much influence the behavior of parasitoid wasps (Kapranas and Tena 2015) and their development within the host (Mawela et al. 2013). In addition, the instar of host larvae has received widespread attention, and often influences adult oviposition choice and the performance of immatures (Hu and Vinson 2000; Nussbaumer and Schopf 2000; Mironidis and Savopoulou-Soultani 2008). Once an egg has been laid in the host, many parasitoid larvae grow by feeding on silk glands
and haemolymph (Jackson et al. 1978), resulting in a reduction of food intake and weight gain of host larvae when compared to the non-parasitized larvae (Mironidis and Savopoulou-Soultani 2009).

In this paper, we describe the host instar preference of *M. similis*, the effects of *S. litura* larval stage on the development of *M. similis* offspring and weight gain and food consumption of parasitized hosts. This study provides knowledge for assessing the biological control potential of *M. similis* against *S. litura*, and for finding the most suitable conditions for mass production of the parasitoid.

**Materials and Methods**

**Study insects**

*Spodopera litura* larvae were reared on pinto bean-based diets until pre-pupation at $30 \pm 1 \, ^{\circ}\text{C}$, $60 \pm 5 \% \text{ RH}$, with a photoperiod of 14:10 (L: D) h (Burton 1970; Li et al. 2015). First to fifth *S. litura* instars were used for these experiments.

A colony of *M. similis* wasps was established in the laboratory in May 2013. Cocoons of *M. similis* were collected from wild second and third instar *S. litura* larvae in a vegetable field near Hunan Agricultural University, Changsha City, Hunan Province, China. The parasitoids were reared on *S. litura* larvae with a 40 % honey solution as food at $30 \, ^{\circ}\text{C}$ (Li et al. 2015). For this experiment, *M. similis* adults were cultured with 30 % honey solution at $30 \pm 1 \, ^{\circ}\text{C}$, $60 \pm 5 \% \text{ RH}$, with a photoperiod of 14:10 (L: D). All experiments were conducted under the same environmental conditions described above. The species identification of *M. similis* was referenced to Li et al. (2015). Molecular identification based on the barcoding *col* sequence following Huang’s methods (Huang et al. 2011) was performed based on the *col* sequence of *M. similis* (GenBank Access Number: KX077544) and the
Microplitis co1 sequences in NCBI, the phylogenetic relationship of M. similis is shown in supplementary figures. There is a growing literature demonstrating that co1 will reliably discriminate a diverse range of taxa at the species level. Therefore, the identification of M. similis was also conducted on the basis of phylogeny using DNA barcoding.

**Instar Preferences**

Experimental studies were conducted on four larval stages (third instar larvae were split into early and late stages after He et al. (2014)) of S. litura using no-choice and choice tests in environmental chambers. In no-choice tests, twenty individual larvae of the same instar and one mated female wasp were kept in a plastic tube (10 cm length × 3 cm diameter, n = 30 per treatment) containing 30 % honey solution and a piece of sweet potato leaf. After 24 h exposure, the host larvae were placed in a plastic case (4.5 × 12.0 × 7.0 cm) and provided with the standard diet. The larvae were cultured in the environment chambers, and checked daily until they pupated or died. Host larvae that died before parasitoid cocooning or pupation were dissected, to ascertain whether they contained parasitoid eggs or larvae. The number of parasitoid cocoons was recorded. Successful parasitization was defined by a host larvae yielding at least one parasitoid, and parasitism was calculated as number of successful parasitized larvae/number of larvae tested × 100.

In choice tests, high densities of fourth and fifth instar S. litura larvae exhibited cannibalistic behavior, which resulted in high mortality. Therefore, only first to late third instars larvae were provided for the choice test. Eighty hosts (20 of each stage) were exposed to 10 parasitoids in a plastic cage (6 ×18 × 12 cm). After 24 h exposure, the host larvae were separated and cultured in an environment chamber under the same conditions as described above for rearing. The host larvae were checked and
recorded daily as described above. This experiment was replicated 5 times. Selection coefficient of *M. similis* was calculated as follows (Cock 1978; Chu et al. 2014).

\[
\text{Selection Coefficient} = \frac{R_i}{\sum_{i=1}^{m} R_i^i}
\]

Where \( R_i \) = the percentage parasitism during host instars \( i \); \( m \) = number of host instars tested.

**Host instar effects on the development of *M. similis***

First to late third instar *S. litura* were tested for their effect on the development and body size of *M. similis*. Adult parasitoid were taken from the culture as described in the no-choice tests. The development times from egg to cocoon, cocoon to adult, emergence percentage and sex ratio were recorded for each treatment. Adult parasitoids which emerged were then checked daily until the host died. Adult body length, front wing length and hind-tibia length of each parasitoid were measured under Motic Digital Microscope Mutual System (He et al. 2014).

**Food consumption and weight gain of parasitized hosts**

The first, second, early third and late third instar *S. litura* were exposed individually to two mated female wasps in a plastic tube (as above). Once each larvae received an oviposition, it was immediately removed. These parasitized host larvae were examined for the body weight and food consumption each day until death or pre-pupation. These data were compared with corresponding data from non-parasitized larvae, which were reared separately from the fifth day. A piece of sweet potato leaf was placed in a test tube without larvae to estimate the percentage of moisture lost daily from a leaf in the environment chambers. The parasitized/unparasitized larvae were weighed on an analytical balance (Mettler Toledo, China). The following formula was used to calculate the food consumption (Chu et al. 2014).
Proportion of daily moisture loss = \( \frac{\text{fresh weight after dehydrating (mg)}}{\text{initial fresh weight of leaf (mg)}} \)

Food consumption (mg) = initial fresh weight of leaf (mg) \times \text{Proportion of daily moisture loss} - \text{fresh weight after feeding (mg)}.

**Statistical analysis**

All data analyses were performed using SPSS 16.0 for Windows (SPSS Inc®, MA, version 16.0), and checked for normality and homoscedasticity (Qiu et al. 2013). When the data did not fit a normal distribution, the parameters, such as host parasitism, emergence percentage and female rate were first transformed by \( \text{arcsin} \, x \), and the parameters, such as the developmental durations were first transformed by \( \text{log}_{10} \, x \). The body length and front wings length were first transformed by \( \text{lngamma} \, x \). The Duncan’s multiple range test was used as a one way-analysis of variance test (\( \alpha = 0.05 \)) in comparing the means between treatments.

For weight gain and food consumption of parasitized hosts data, a two way-analysis of variance test (\( \alpha = 0.01 \)) was conducted to show the impact of both instar and level parasitism (parasitized and non-parasitized) on the change of weight gain and food consumption of parasitized hosts (McLoud 2011).

**Results**

**Instar preference**

*Microplitis similis* did not parasitize fourth and fifth instars larvae, which possess strong defensive behavioral reactions. Parasitism in no choice experiments was highest in second and third instar, and lowest in first instar (\( F = 49.01; \text{df} = 3, 117; P < 0.05; \text{Fig. 1} \)). When provided with a choice of first to late third instars host larvae (Fig. 2), early third and second instars *S. litura* were parasitized at the highest rates,
followed by first and late third instars ($F = 7.43; df = 3, 20; P < 0.05$). There was also significant variation in the selection coefficient among instars ($F = 26.39; df = 3, 20; P < 0.05$), being highest in the early third instar at and lowest in the late third instar (Fig. 3).

**Effects of *S. litura* stage on the development of *M. similis* offspring**

Egg-cocoon development of *M. similis* in first instar *S. litura* was 7.06 ± 0.06 days, and increased with successive instars to 8.10 ± 0.08 days in the late third instar of *S. litura* ($F = 41.22; df = 3, 504; P < 0.05$) (Table 1). Cocoon-adult development in late third instar larvae was significantly longer than in other host instars ($F = 38.91; df = 3, 504; P < 0.05$). It just took 4.13 ± 0.06 days for *M. similis* to develop from cocoon-adult in the first instar host. The development duration of the parasitoid from egg to adult emergence was longest in the late third instar larvae, followed by the early third instar, second instar and finally first instar ($F = 110.78; df = 3, 504; P < 0.05$). The stage of the host parasitized had no effect on the percentage of parasitoids successfully emerging ($F = 0.493; df = 3, 93; P = 0.688$).

The female rate was lower for adults emerging from first instar larvae than from other host instars ($F = 3.90; df = 3, 35; P < 0.05$). The female rate did not differ between adults emerging from the older instars ($F = 3.90; df = 3, 35; P = 0.248$). In the aspect of morphometric features, parasitoid body length, fore wing length, and hind tibia length were affected by all host instars (Table 2). All the above morphometric features increased with the increase of host instar age at parasitism. All of the morphometric features (body length, fore wing length, hind tibia length), were greatest in late third instar larvae (body length: $F = 102.33; df = 3, 231; P < 0.05$;
fore wing length: $F = 79.10; \text{df} = 3, 231; P < 0.05$; hind tibia length: $F = 42.52; \text{df} = 3, 231; P < 0.05$).

**Weight gain and food consumption of parasitized hosts**

Parasitized *S. litura* showed a pronounced reduction in weight gain (ANOVA: level of parasitism, $F = 77.30; \text{df} = 1, 255; P < 0.01$; host instars, $F = 130.07; \text{df} = 3, 255; P < 0.01$, interaction: $F = 543.40; \text{df} = 1, 255; P < 0.01$) (Fig. 4). All instar of *S. litura* larvae were affected, although this was most marked in the late third instar, which were the largest when unparasitized. The daily food consumption of parasitized hosts showed a clear reduction in the food consumed (ANOVA: level of parasitism, $F = 86.72; \text{df} = 1, 256; P < 0.01$; host instar, $F = 91.92; \text{df} = 3, 256; P < 0.01$, interaction: $F = 548.45; \text{df} = 1, 255; P < 0.01$) (Fig. 5). All instar of *S. litura* larvae were affected, and this was most notable in the late third instar, which had the highest rate of food consumption when unparasitized. The maximum food consumption of the host after parasitism was dependent on host instar of *S. litura* into which the parasite had oviposited.

**Discussion**

We found that female *Microplitis similis* were more likely to parasitize second and early third instars *S. litura* larvae in both choice and no-choice tests, and immature *M. similis* grew well in second and early third instars *S. litura* larvae. *Microplitis similis* had a lower success rate parasitizing late third instar larvae, most likely as an effect of the older instar larvae having a strong defensive behavior (Ameri et al. 2014). Early third instar hosts ranked higher than other instars in terms of suitability, yielding
offspring of greater body size, although they required a slightly longer development time. Parasitized *S. litura* larvae exhibited a decreased rate of weight gain and daily food consumption in comparison to unparasitized larvae. In summary, *M. similis* females selected the best host stage for the development of their offspring, and had the beneficial effect of greatly reducing consumption by host larvae. Thus, it is apparent that enhancing *M. similis* can contribute as a component of an IPM strategy for *S. litura* in economic crops.

Host evaluation and selection by hymenopteran parasitoids is a key event, because high host suitability is critical to the growth and development of parasitoid larvae (Vinson 1990; Murillo et al. 2013). Under natural conditions, adult females accept or reject host larvae for oviposition, and the success rates of parasitism are low when inoculating small and large hosts due to early host death and parasitoid injury, respectively (Wei et al. 2014). In our study, we identified the preferred host instar range of *M. similis* to be first to late third instars *S. litura* larvae. When *M. similis* sought to parasitize the fourth and fifth instars larvae, the larvae demonstrated strong defensive behaviors. When the parasitoid used antennae to evaluate fourth and fifth instars larvae, the larvae would twist the head and attack the wasp, resulting in wounding and even death. Selectivity by parasitoids, in terms of host instar, has been demonstrated by He et al. (2014) who found that *Euplectrus laphygmae* (Ferrière) also could parasitize first to late third instars *S. litura* larvae, but parasitized second and third instar hosts at a higher rate in no-choice tests. However, *E. laphygmae* prefer to parasitize early third *S. litura* larvae when offered a choice of second, early third and late third.

In koinobionts, parasitoid larvae have two kinds of feeding strategies, tissue-feeding strategies and hemolymph-feeding (Harvey and Malcicka 2016). The
The vast majority of Microgastrinae parasitoid larvae consume most host tissues, including in the genera *Microgaster, Apanteles* and *Dolichogenidea* (Harvey and Malcicka 2016). The larva of *Hyposoter ebeninus* consumed the entire *Pieris rapae* larvae (Gauld and Bolton 1988). On the other hand, microgastrines in the genera *Microplitis* and *Cotesia* are hemolymph feeders (Malcicka and Harvey 2015). Different hosts provide quantitative and qualitative variation in nutrition, which consequently influence the growth and ultimately the morphometric features of parasitoid offspring from different host instars or host species (Harvey 2000; Harvey and Strand 2002; Mironidis and Savopoulou-Soultani 2009). In general, larger hosts yield more offspring (Harvey 2000; Stockermans and Hardy 2013) and female offspring often hatch from large high quality hosts and male offspring from low quality hosts (Charnov et al. 1981; Pekas et al. 2016). Strand et al. (1988) also reported that *Microplitis demolitor* achieved larger body size in the larger later instars of *Heliothis virescens* larvae, in a manner similar to that found here for *M. similis* developing in *S. litura*. The female *M. similis* laid few fertilized eggs in small early instar larvae. The low female ratio obtained from first instar *Microplitis tuberculifer* larvae (Chu et al. 2014) is similar also to this study.

Harvey et al. (2010) reported that the intensity of resourced-related constraints on parasitoid development also varies from one parasitoid species to another. In koinobiont parasitoids, the host represents a potentially dynamic resource, koinobiont parasitoids attack hosts that continue feeding and growing during immature parasitoid development (Harvey 2005; Harvey et al. 2010). In our study, larvae of *S. litura* parasitized by *M. similis* continue feeding and growing until the mature parasitoid larva emerged from the body of the host, and the parasitized herbivores showed greatly reduced weight gain and food consumption compared to unparasitized larvae,
whatever the instar parasitized. This result is similar to larvae of *Mythimna separata* parasitized by *M. mediator* (Li et al. 2006). In terms of host regulation, early instar hosts provide insufficient nutrients to koinobiont parasitoids, so parasitoids do not immediately inhibit lepidopteran host growth until the host becomes larger. In contrast, koinobiont parasitoids attack later instar lepidopteran hosts greatly reducing host growth (Harvey et al. 1994, 2014). In this study, significant decreases in weight gain of the first instar *S. litura* larvae parasitized by *M. similis* began on the fifth day, whereas late third instar host larvae showed a significant decrease in weight gain only two days after being parasitized by *M. similis*. Very similar results were obtained for *Hyposoter didymator* parasitizing *Helicoverpa armigera* (Mironidis and Savopoulou-Soultani 2008). Changes in the food consumption of *S. litura* parasitized by *M. similis* began on different days after parasitism in different host instars, a result similar to that found for *M. tuberculifer* on *S. exigua* (Chu et al. 2014). The influence of a parasitoid on host feeding is of key importance to the role of parasitoids in IPM strategies, and from a plant protection perspective it is clearly an advantage if parasitization results in a larvae reducing or halting feeding. In that *M. similis* results in a progressive reduction in feeding over a moderate period, it is less favorable than a species which rapidly causes the host to stop feeding.

We found that *M. similis* parasitized second and early third instar larvae at a higher rate than other larvae, and this was optimal for the parasitoid larvae to develop and survive. Most parasitoids can successfully complete their life cycle in different instar host larvae, but with different performance in different instars. This is important for IPM of the pest, but also is important information for mass-rearing of *M. similis*. In the current study, we found the most suitable host stage for the growth of *M. similis* to be early third instar. However, environmental factors, such as temperature, humidity
and photoperiod, as well as the host species, body size, and instar would influence parasitoid’s mass production. The details to optimize rearing of *M. similis* will require further study, but the control of *S. litura* with parasitic natural enemies can potentially provide a component of an overall IPM strategy.

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Tables

Table 1. Mean (±SE) developmental time, percentage of emergence and female rate of *M. similis* in different developmental stage of *S. litura*

Table 2. The effect of host stages on body size of *M. similis* (M±SE)

Figure legends

Figure 1. Mean (±SE) parasitism of *M. similis* parasitizing *S. litura* in no-choice tests. Note: The x axis values represent 1 = first instar larvae, 2 = second instar larvae, 3E = early third instar larvae, 3L = late third instar larvae, respectively. The same letters above the same bars represent no significant differences in parasitism of the different host instars (Duncan test, *P* < 0.05)

Figure 2. Mean (+ SE) parasitism of *S. litura* larvae of various instars when parasitized by *M. similis* in choice tests. Note: The x axis values represent 1 = first instar larvae, 2 = second instar larvae, 3E = early third instar larvae, 3L = late third instar larvae, respectively. The same letters above the same bars represent no significant differences in parasitism of the different host instars (Duncan test, *P* < 0.05)

Figure 3. Selection coefficient of *S. litura* larvae of various instars when parasitized by *M. similis* in choice tests. Note: The x axis values represent 1 = first instar larvae, 2 = second instar larvae, 3E = early third instar larvae, 3L = late third instar larvae, respectively. The same letters above the bars represent no significant differences in selection coefficient of the different host instars (Duncan test, *P* < 0.05)

Figure 4. Weight gain by hosts parasitized in different larval instars and un-parasitized hosts. (a) Weight of parasitized and un-parasitized first instar *S. exigua*, (b) Weight of parasitized and non-parasitized second instar *S. exigua*, (c) Weight of parasitized and non-parasitized early third instar *S. exigua*, and (d) Weight of parasitized and non-parasitized late third instar *S. exigua*, Note:
1-d-old *S. litura* larvae were not parasitized, so they were used at 2 days old. Control: Un-parasitized larvae. Weight gain of parasitized / non-parasitized 1-d-old to 4-d-old larvae was measured together, therefore the larvae (1-d-old to 4-d-old) values are mean, and the other values are mean + SE.

**Figure 5.** Daily food consumption parasitized in different larval instars and un-parasitized hosts. *(a)* Daily food consumption of parasitized and non-parasitized first instar *S. exigua*, *(b)* Daily food consumption of parasitized and non-parasitized second instar *S. exigua*, *(c)* Daily food consumption of parasitized and non-parasitized early third instar *S. exigua*, and *(d)* Daily food consumption of parasitized and non-parasitized late third instar *S. exigua*, Note: 1-d-old *S. litura* larvae were not parasitized, so they were used at 2 days old. Control = non-parasitized larvae. Daily food consumption of parasitized / non-parasitized 1-d-old to 4-d-old larvae was measured together, therefore the larvae (1-d-old to 4-d-old) values are mean, and the other values are mean + SE.