

Article

Effectiveness of *Diachasmimorpha longicaudata* in Killing *Ceratitis capitata* Larvae Infesting Commercial Fruits in Dryland Agroecosystems of Western Argentina

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Abstract: *Ceratitis capitata* (Wiedemann) (medfly) strongly affects Argentinean fruit production and export. Augmentative biological control using the exotic parasitoid *Diachasmimorpha longicaudata* (Ashmead) is currently applied to this problem. The ability to find and parasitize medfly larvae on a wide diversity of fruit host species is a key issue that needs to be analyzed. This research assessed the effect of the physical features of fruit on the preference of foraging *D. longicaudata* females and the influence of varying release density on parasitoid performance as a pest mortality factor in three fruit species. Trials were performed inside field cages under semi-arid environmental conditions in Argentina's central-western fruit-growing region. Sweet orange, peach, and fig were tested. The fruits were inoculated with third-instar larvae of the Vienna-8 temperature-sensitive lethal medfly strain. Naïve, 5 d-old mated *D. longicaudata* females were released in cages at 20, 40, 80, and 160 parasitoid densities. The highest levels of medfly mortality and parasitoid emergence were recorded in fig and peach, although *D. longicaudata* also induced mortality in orange, a fruit with few physical features favorable to parasitism. The medfly mortality in all fruit host species significantly increased with an increased number of parasitoid females released into the field cages. *Diachasmimorpha longicaudata* has high potential as a medfly biocontrol agent.

Keywords: medfly; host fruits; semi-arid environmental; larval parasitoid performance; biological control; augmentative parasitoid release



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1. Introduction

Ceratitis capitata (Wiedemann) (Diptera: Tephritidae) is an invasive dipterous species native to sub-Saharan Africa, although it is widespread throughout the African continent [1]. Commonly known as the Mediterranean fruit fly or just medfly, it is currently regarded as a cosmopolitan species due to its worldwide dispersal, mainly resulting from the increase in the global fruit trade [2]. The medfly is a highly polyphagous pest species that infests healthy, commercially valuable fruit [3]. The pattern of host relationships is mainly related

to the fruits available locally in the region where the medfly occurs [4]. The economic damage caused by the medfly is related to its biological cycle [5]. After mating, females lay eggs inside healthy fruit, causing additional damage to the epidermis and allowing fruit-rotting organisms such as insects and/or pathogens to access them. Eggs give rise to larvae that pass through three instars while feeding on the fruit flesh, which leads to the softening and rotting of the fruit. The mature larvae leave the fruit to pupate by burying themselves in the soil or within or beneath the fallen fruit. The adult emerges from the puparium and searches for a host plant to access water, food, and shelter, and the female searches for an optimal environment for mating and oviposition. From the first record of medfly infesting commercial peaches in Argentina in 1905 [6], this pest has been characterized by its rapid range expansion throughout all Argentinian fruit-growing regions, covering latitudes from 22° to 56° S. However, fruit-producing areas of the Patagonian Region (southern Argentina) and the Central and Southern Oases of Mendoza from the Cuyo region (southwestern Argentina) are currently fruit fly free areas [7]. The medfly has been reported to infest 58 commercial and wild, exotic, and indigenous fruit species grown in fruit-producing regions throughout Argentina [8,9]. Given that Argentina is a relevant world producer of fresh fruits and vegetables due to its diverse ecosystems [10], the widespread distribution of the medfly is a critical constraint for the country's fruit-growing industry. In this regard, *C. capitata* is one of the main fruit pests that strongly affects the production, marketing, and export of fruits, which has a negative socioeconomic impact on Argentinean fruit systems [11]. This pest causes economic losses either by direct damage to fruits, such as the presence of larvae inside fruit or oviposition activity of females, or by indirect losses, which involve quarantine treatments or other measures that increase marketing costs in response to export restrictions imposed by importing countries [8]. Therefore, given the adverse effects of the medfly on the Argentinian economy, since 1994, the federal government has put into operation the National Fruit Fly Control and Eradication Program (PROCEM, Spanish acronym) [12], which currently involves area-wide integrated medfly management approaches. The strategy applied through PROCEM has been based on the combined use of the sterile insect technique (SIT), cultural and air/ground chemical controls, trapping systems, a quarantine protection system, and the implementation of a phytosanitary emergency schedule for pest outbreaks in medfly free or low-prevalence areas [13]. However, from 2014 to the present, augmentative biological control has been used by releasing parasitoids in fruit crops by the PROCEM in the semi-arid fruit production valleys of the province of San Juan (central-western Argentina, Cuyo region) as a complementary tool in an environmentally friendly way [14].

The implementation of augmentative biological control against fruit flies in Argentina was achieved through the establishment and optimization of the mass rearing of the South Asian-native, larval parasitoid *Diachasmimorpha longicaudata* (Ashmead) (Hymenoptera: Braconidae) [15]. This braconid parasitoid species belongs to a guild of synovigenic, solitary, koinobiont, endoparasitoids that attack late-instar larvae of tephritid fruit flies [14]. The *D. longicaudata* female forages on infested fruit with host larvae and oviposits by drilling through the pericarp from the exterior [15]. This exotic parasitoid is currently reared using cobalt-60 irradiated medfly larvae of the temperature-sensitive lethal (=tsl) genetic sexing Vienna-8 strain at the San Juan mass-rearing biofactory [16]. Results from a recent augmentative release of *D. longicaudata* showed a significant decrease in the wild medfly population inhabiting a commercial multi-fruit farm in the province of San Juan [14]. Such results, added to the previous ones reported by Sanchez et al. [15], revealed a medfly population control between 40 and 70%. Such evidence encourages the use of *D. longicaudata* against medfly in the fruit-growing oases of San Juan. Furthermore, the need to promote sustainable and eco-friendly agricultural strategies in arid areas in a progressively warming and drying world [17] highlights the need to focus on this issue. Likewise, the lack of resident parasitoid species that attack the pest in the region [18] reinforces the use of biologically plastic imported species, i.e., those species capable of adapting to different environmental conditions, such as the braconid *D. longicaudata* [19].

Earlier studies [14,15] showed how *D. longicaudata* mass reared at the San Juan biofactory can perform effectively as a biocontrol agent under semi-arid environmental conditions in the ecologically isolated fruit-growing valleys of San Juan. In those areas, the climatic characteristics of arid lands, such as high daytime and low overnight temperatures, drought, strong winds, low relative humidity, sparse vegetation, and low alternative host densities, are considerably attenuated. Another essential factor that must be addressed involves host species diversity in the fruit-growing region and their influence on parasitoid effectiveness in finding and parasitizing host larvae. Large fruits, such as citrus, can limit parasitoid foraging behavior on fruit-infesting Tephritidae [20–22]. Therefore, the host fruit's physical features have important practical implications for implementing medfly biological control in any fruit-producing region.

Several authors highlighted the ability of *D. longicaudata* to forage successfully on host larvae over a wide range of non-crop and crop fruit species [19,23–27]. Given this, it is hypothesized that the *D. longicaudata* female is particularly efficient at finding and killing medfly larvae infesting diverse commercial host fruit species ranging from orange, fig, and peach in the irrigated fruit-growing areas of San Juan. In addition, it is also hypothesized that the parasitoid female becomes significantly more efficient at increased release densities. This assumption is based on several open-field augmentative release trials using *D. longicaudata* for suppressing pest tephritid fruit fly populations worldwide [23,27–31]. Therefore, this research aims to evaluate the performance of *D. longicaudata* females as a medfly mortality factor in three main host fruit species. For this purpose, the physical features of the fruit species and the potential effect of variation in the parasitoid release densities were assessed. Thus, this study discusses the findings of using *D. longicaudata* mass reared at the San Juan biofactory as a biological control agent against the medfly in one of Argentina's most important fruit-growing regions, recognized for its semi-arid climate marked by hot and cool extreme temperatures.

2. Materials and Methods

2.1. Insect Rearing Procedures

Diachasmimorpha longicaudata adult females were reared from parasitized 90 Gy-irradiated medfly larvae of the Vienna-8 tsl strain at the Parasitoid Rearing Laboratory of the San Juan mass-rearing biofactory. This laboratory belongs to the Plant, Animal, and Food Health Bureau of the government of the San Juan province, located in the central-western fruit-growing region of Argentina. Parasitoids were reared using 60 cm × 60 cm × 30 cm rectangular iron-framed screen-covered cages at 25 ± 1 °C, 60 ± 5% RH, 12:10 (L:D), and were fed with honey and water ad libitum every other day. Medfly tsl larvae were reared at the Fruit Fly Rearing Laboratory of the San Juan biofactory based on Caceres' [32] protocol. Medfly larvae radiation was performed in an IMO-1 mobile Gamacell irradiator with a Co-60 source of γ irradiation, located at the San Juan biofactory, which belongs to the National Atomic Energy Commission from Argentina.

2.2. Test Site and Weather Conditions

The trials were performed inside field cages under uncontrolled environmental conditions at a 1000-m² experimental field of the Plant, Animal, and Food Health Bureau (31°31'00" S, 68°36'00" W, 710 m), located in the Rivadavia District, Tulum Valley, San Juan. A windbreak curtain of poplar trees (*Populus alba* L., Salicaceae) surrounds the experimental plot, supplying natural shade within the site. According to Köppen's classification, the San Juan lowlands, in which the main fruit-producing valleys are found, possess a BWwka climate, i.e., a desert climate with a high concentration of rainfall in the summer, with a mean annual temperature below 18 °C [33]. Mean relative humidity (=RH), mean air temperature (=AT), and precipitation were recorded during the study using a digital weather station (LUFT[®], model WS80, Shenzhen, China). Both HR and AT and accumulated rainfall recorded during the trial dates are shown in Table 1.

Table 1. Minimum (=Min), maximum (=Max), and average temperature and relative humidity, and accumulated rainfall recorded during trial dates in March 2019 and 2020 (late summer/early autumn).

Study Year	Temperature (°C)			Relative Humidity (%)			Precipitation (mm)
	Min.	Max.	Average Monthly	Min.	Max.	Average Monthly	Accumulated Rainfall
2019	9.9	36.3	21.1	10	86	56.2	0.2
2020	10.9	36.5	22.8	23	89	57.1	0.3

2.3. Tested Host Fruit Species and Fruit Handling

Three fruit species were tested: *Citrus sinensis* (L.) Osbeck (“sweet orange”, Salustiana cultivar), *Prunus persica* (L.) Batsch (“peach”, Hesse cultivar), and *Ficus carica* L. (“fig”, Brown Turkey cultivar). The fruits were purchased at the market and, once brought to the laboratory, were washed and disinfected with a sodium benzoate and nipagin methylparaben solution. Before tests, each fruit was weighed, and rind thickness, flesh depth, and fruit diameter were measured, and the surface area ($=4\pi r^2$) over which the female parasitoid was to forage was also calculated. Fruits of similar size and ripeness were selected for the trials. Each washed fruit was artificially inoculated with non-irradiated 5 d-old (early third-instars) larvae of the tsl medfly strain. Host larval density inoculated on fruit remained constant in all tests. Larval density per fruit species depended on the fruit surface area of each fruit species; therefore, a significantly equal larval density was used relative to fruit surface area. Thus, each orange or peach was inoculated with 130 medfly larvae to match larval proportions per fruit species, while the fig was inoculated with 100 host larvae. Inoculation was carried out by cutting the top of the fruit using a sterilized scalpel; half of the pulp was removed, and the fruit was filled with host larvae. They occupied 50% of the fruit volume. In the case of the peach, the stone was removed. Once the fruit was inoculated, the small upper cover (lid) and the large lower portion (body) were attached with a 2.5 cm-wide Parafilm “M[®]” sheets (Pechiney Plastic Packaging, Chicago, IL, USA). Each infested fruit accounted for one oviposition unit.

2.4. Experimental Setup and Procedure

The tests were performed under natural conditions and with variation in parasitoid female release densities. Trials were carried out inside three 3.5 m × 3.0 m (diameter × height) nylon field cages throughout March 2019 and March 2020, corresponding to late summer and early autumn. Five 0.5 m × 2.0 m (diameter × height) voile mesh-covered tube cages were placed inside each major field cage. Each tube cage had a 1.5 m long zipper along its middle part to enter the cage. The tube cages were placed at 20 cm distance from each other inside the major field cage (Figure 1). Each field cage was protected from solar radiation by a 130 g/m² black shade net and from rain by a 100- μ translucent high-density polyethylene cover (AGROREDES[®], Buenos Aires, Argentina). Both coverings were 40 cm above the cage, allowing natural light and preventing rainwater. A potted small orange, peach, or fig tree was placed inside each tube cage according to the fruit species of the treatment to simulate a natural environment. Fruit trees were 1–1.3 m high, 2–3 yrs-old, had 10–20 leaves, and were purchased from a commercial fruit plant nursery in San Juan City. For each tube cage, three fruits inoculated with host larvae were placed. The fruit species used varied according to the treatment. The fruits were placed equidistant in a circle only in the upper section, at 1.5 m from the cage roof. The fruits were hung from the top using a 0.5 mm nylon fishing line attached to a metal clip inserted into the Parafilm strip wrapped around the fruit. Immediately below the fruit was placed into a 10 cm × 2 cm (diameter × height) galvanized wire basket with a double voile covering, with a poplar shaving at the bottom as a pupation substrate. The device was held with four nylon fishing lines tied to the main rope from which the fruit hung. Each major field cage accounted for one treatment as follows: treatment #1 involved only peach (=T1), treatment #2 sweet orange (=T2), treatment #3 fig (=T3). Each treatment was divided into four sub-treatments

with different *D. longicaudata* release densities and one control. Sub-treatment locations were randomized to avoid the edge effects. All sub-treatments and the control were simultaneously performed. In sub-treatment “A”, 20 parasitoid females were released ($=T_A$), sub-treatment “B” involved 40 parasitoid females ($=T_B$), in sub-treatment “C”, 80 parasitoids were released ($=T_C$), and in sub-treatment “D”, 160 parasitoids were released ($=T_D$) (Figure 1). Three controls were tested as follows: controls #1, #2, and #3 involved only peaches, oranges, and figs infested with medfly larvae, respectively, but no parasitoids were released ($=C_1$, C_2 , and C_3 , respectively) (Figure 1). The host–parasitoid ratio per sub-treatment involving sweet orange and peach was 6.5, 3.3, 1.6, and 0.8 host larvae per 1 parasitoid female released, whereas in the fig sub-treatment it was 5, 2.5, 1.3, and 0.6 host larvae per 1 parasitoid female. The trials were conducted with 5 d-old mated *D. longicaudata* females, which had not previously been exposed to host larvae, infested fruit, or both. The parasitoids tested were from an artificial rearing of 65 generations. The parasitoids were released into experimental tube cages on leaves of the potted fruit tree using 2000 mL transparent plastic vials. Once the parasitoid females were released, they were allowed to forage infested fruit for 48 h starting at 9 a.m. In this study, only fruits located in the highest section of the cage were used. Previous studies showed a similar ability of *D. longicaudata* females to forage on infested fruit located on the ground or in the tree canopy [34,35]. Once foraging time was over, the fruit, the pupation substrate, and parasitoids were removed from each tube cage. At the laboratory, fruits were dissected to remove living host larvae, and puparia were recovered from each basket. All hosts were placed in 10 cm × 8 cm (diameter × height) disposable plastic cups with sterilized poplar shavings at the bottom until insect emergence. These devices were kept at 26 ± 1 °C, $75 \pm 5\%$ RH, and in darkness. Once the insects emerged, the number and sex of the parasitoids and the number of adult flies were recorded from each sub-treatment and control test. Based on these data, the biological parameters were determined as follows: (1) the host-killing parasitoid ability, (2) the parasitoid reproductive success, and (3) the parasitoid offspring sex ratio. The first parameter is equivalent to the parasitoid performance as a pest killer, and it was calculated using Abbott’s percent-corrected host mortality index [36]. This index allows the determination of the host-killing parasitoid female capacity by the interaction between host emergence from the treatment and that from the control. The parasitoid reproductive success was estimated as the number of emerged parasitoid offspring divided by the number of exposed larvae inside the fruit × 100. The sexual ratio of parasitoid offspring was calculated as the percentage of emerged females divided into all recovered adult parasitoids. Each sub-treatment and the related controls were replicated 10 times in both 2019 and 2020. Each replicate involved new inoculated fruit and pupation substrate on each basket and a new parasitoid female cohort.

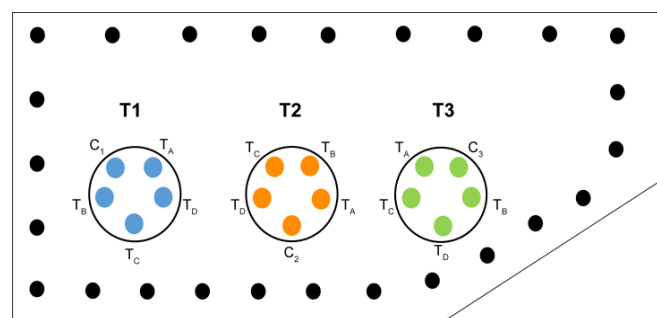


Figure 1. Location of the three major field cages and their five internal tube cages at the experimental trial yard from the Plant, Animal, and Food Health Bureau, government of the San Juan province, Rivadavia district, San Juan province, Argentina. Specifications: large colorless circle = main experimental field cage; small blue, orange, and green circles = inner tube cages; small black circles = poplar tree windbreak. Treatments: T1 = infested peaches; T2 = infested oranges fruit; T3 = infested figs. Sub-treatments: T_A = 20 released female parasitoids (RFP), T_B = 40 RFP, T_C = 80 RFP, T_D = 160 RFP. Control tests: C_1 = control test from T1; C_2 = control from T2; C_3 = control from T3.

2.5. Data Analysis

The host-killing, parasitoid reproductive success, and the parasitoid offspring sex ratio data were analyzed as nested models, with three-factor levels (year, fruit, and number of released parasitoids), implementing Generalized Linear Mixed Models (GLMM) performed with the function “lme” from “nlme” [37,38] packages of R [39]. The final model was implemented using only significant factors at $\alpha = 0.05$. For paired data, the comparison of host killing, reproductive success, and sex ratio throughout the experiment and within treatments, without considering factors, was made using a Wilcoxon signed rank test with continuity correction, with the “wilcox.test” function from the “stats” based packages of R [39]. Comparisons of the fruit features were made using the Kruskal–Wallis rank sum test, using the “kruskal.test” function from the “stats” based packages of R [39]. The R-4.4.1 software was used for statistical analysis [39]. The box plot figures show the median (horizontal line inside the box), interquartile range Q1–Q3 (bottom and top ends of the box), range (minimum: Q0, maximum: Q4; both ends of the whisker on the vertical line outside the box), and outliers (red balls).

3. Results

3.1. Tested Host Fruit Species

The physical features between the tested medfly host fruit species were significantly different (weight, $\chi^2 = 79.12$, $df = 2$, $p < 0.01$; diameter, $\chi^2 = 74.50$, $df = 2$, $p < 0.01$; rind thickness, $\chi^2 = 79.12$, $df = 2$, $p < 0.01$; pulp depth, $\chi^2 = 79.12$, $df = 2$, $p < 0.01$; surface area, $\chi^2 = 62.79$, $df = 2$, $p < 0.01$). Sweet orange had a significantly greater weight, diameter, rind thickness, pulp depth, and external surface area than peach and fig (Table 2). Peach differed significantly from fig in all physical traits, while the host density/cm² of the fruit surface used in the trials was significantly similar for all fruit species (Table 2).

Table 2. Physical features of host fruits used in this study, *Citrus sinensis* (sweet orange), *Prunus persica* (peach), and *Ficus carica* (fig), and density of *Ceratitis capitata* larvae per cm² of fruit surface area.

Host Fruits	Fruit Physical Features (n = 30) (Mean ± SE)					Medfly Larvae	
	Weight (g)	Diameter (cm)	Rind Thickness (cm)	Pulp Depth (cm)	Surface Area (cm ²)	Density * (cm ²) (Mean ± SE)	Larvae per Fruit
Sweet orange	200.2 ± 4.8 a	7.1 ± 0.2 a	4.6 ± 0.2 a	3.1 ± 0.1 a	44.0 ± 8.0 a	3.08 ± 0.06 a	130
Peach	156.9 ± 1.5 b	6.6 ± 0.1 b	0.3 ± 0.1 b	2.5 ± 0.1 b	41.9 ± 0.4 b	3.10 ± 0.03 a	130
Fig	57.2 ± 0.8 c	5.1 ± 0.2 c	0.1 ± 0.1 c	2.0 ± 0.1 c	32.3 ± 1.1 c	3.11 ± 0.01 a	100

* Inoculated host larvae ratio per cm² of fruit surface area (density). Means followed by different letters within the same column are significantly different (Kruskal–Wallis rank sum test, $p = 0.05$).

3.2. Host-Killing Parasitoid Capacity

The medfly killing parasitoid capacity parameter did not vary significantly between the two study years ($Estimate = 1.20$, $SE = 0.94$, $df = 709$, $Z = 1.27$, $p = 0.20$). Therefore, the data from each testing year were pooled for analysis. The results for each tested host fruit species by parasitoid release density are shown in Figure 2. The parasitoid females killed between 54 and 88% of the host larvae in the fig, between 52 and 83% in the peach, and between 34 and 62% in the sweet orange. When all of the treatments were pooled, the host-killing parasitoid capacity of the tested fruit species was significantly higher in the fig (71% marginal mean), followed by the peach (67%), and the lowest in the sweet orange (46%) (Table S1). Overall, the host-killing parasitoid capacity became significantly higher as parasitoid release density increased, regardless of the tested fruit species. The marginal means of the host mortality percentage pooling of all of the fruit species were ~47, 53, 69, and 78 for treatments #1, #2, #3, and #4, respectively (Table S2).

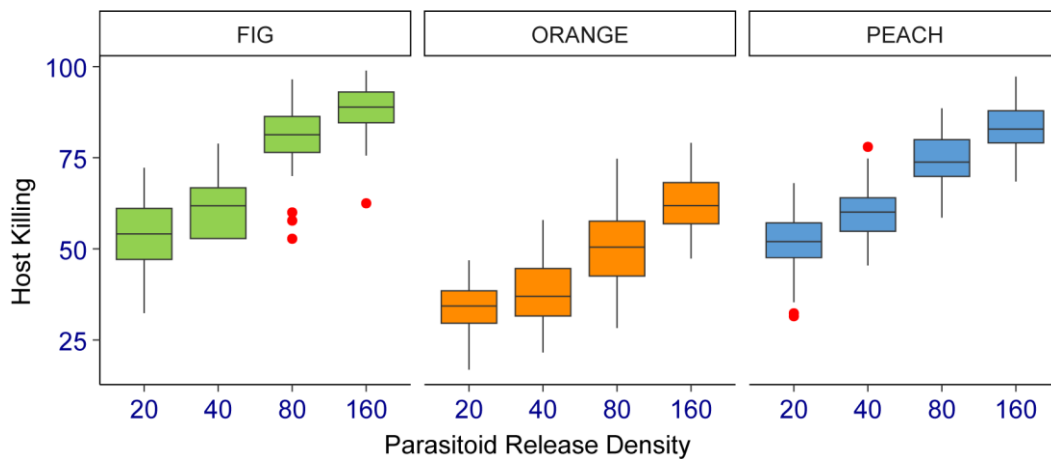


Figure 2. Percentage of the medfly killing ability of *Diachasmimorpha longicaudata* on three fruit host species (fig, sweet orange, and peach) at four parasitoid release densities (20, 40, 80, and 160 females) under field cage conditions in Tulum fruit-growing valley, San Juan, central-western Argentina.

3.3. Parasitoid Reproductive Success

In the case of parasitoid reproductive success, there is a small but significant difference between the two years studied ($Estimate = 0.81$, $SE = 0.34$, $df = 709$, $Z = -2.39$, $p = 0.02$). Thus, the nested model was performed for this parameter at three levels (studying year, fruit species, and parasitoid release density). The results of the parasitoid reproductive success for each host fruit species by parasitoid release density are shown in Figure 3. The percentage of offspring produced by *D. longicaudata* females from fig, peach, and sweet orange varied between 19 and 30%, 20 and 28%, and between 11 and 21%, respectively. Both fig and peach were the two host fruit species with the most substantial influence on increasing parasitoid offspring when pooling treatments (25 and 24% marginal means for fig and peach, respectively). In contrast, the sweet orange had the least impact on the parasitoid reproductive success by a significant margin (16% marginal mean) (Table S3). When the treatments were pooled, the parasitoid reproductive success significantly increased as parasitoid release density increased. The marginal means of the parasitoid offspring percentages were 17, 19, 24, and 27 for treatments #1, #2, #3, and #4, respectively (Table S4).

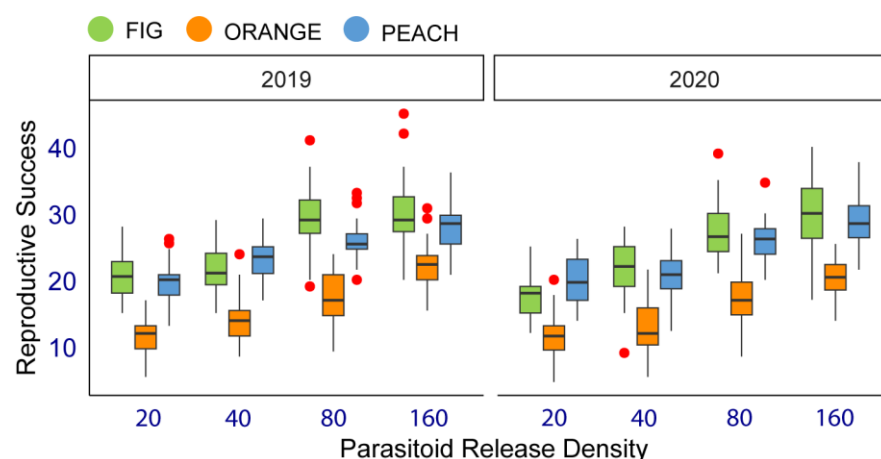


Figure 3. Percentage of offspring produced by *Diachasmimorpha longicaudata* (parasitoid reproductive success) when parasitizing medfly larvae infested three fruit host species (fig, sweet orange, and peach) at four parasitoid release densities (20, 40, 80, and 160 females) in studying years 2019 and 2020 under field cage conditions in Tulum fruit-growing valley, San Juan, central-western Argentina.

3.4. Parasitoid Offspring Sex Ratio

The parasitoid offspring sex ratio did not vary significantly between the two study years ($Estimate = 0.28$, $SE = 0.80$, $Z = 0.35$, $p = 0.72$). Therefore, the data from each testing year were pooled for analysis. The results of the emerged parasitoid females for each host fruit species by parasitoid release density are shown in Figure 4. *Diachasmimorpha longicaudata* exhibited around a 1:1 female–male sex ratio, as the percentage of females emerged from medfly puparia recovered from all three host fruit species ranged from 49 to 51%. There were no significant differences between the marginal means of parasitoid offspring sex ratio when both tested factors, host fruit species (Table S5), and parasitoid release densities (Table S6) were compared.

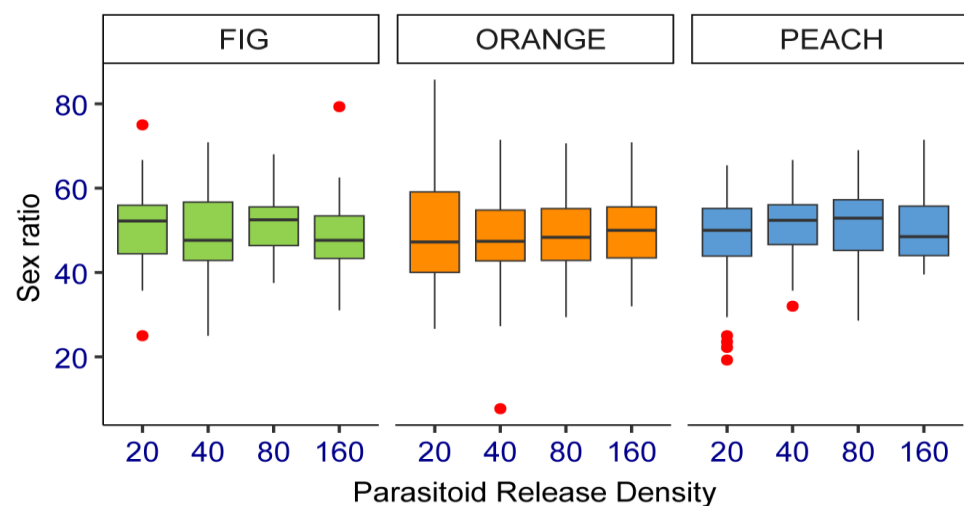


Figure 4. Percentage of emerged *Diachasmimorpha longicaudata* females from medfly puparia recovered from three fruit host species (fig, sweet orange, and peach) at four parasitoid release densities (20, 40, 80, and 160 females) under field cage conditions in Tulum fruit-growing valley, San Juan, central-western Argentina.

4. Discussion

Augmentative biological control using parasitoids has become a relevant complementary tool with other eco-friendly techniques for medfly control [27,31]. Therefore, parasitoid species selected as potentially effective medfly biocontrol agents entail a prior and exhaustive assessment of their host foraging and killing abilities in different host fruits and environments. Understanding these biological aspects is strategically essential for developing and implementing a parasitoid mass-release biological control schedule. In this framework, the current study reports the ability of the introduced parasitoid *D. longicaudata* to kill the invasive *C. capitata* infesting main host fruit species through field cage-simulated augmentative releases. Consequently, the results highlighted three major findings. Firstly, *D. longicaudata* females successfully parasitized host larvae on all three tested medfly multiplying fruit species. Secondly, the highest levels of both medfly mortality and parasitoid offspring were recorded in those fruit species with physical features highly favorable to parasitism, such as fig and peach. Thirdly, the medfly mortality recorded on all tested fruit host species significantly increased as the number of parasitoid females released into field cages increased.

The first finding highlighted the ability of *D. longicaudata* to successfully find and parasitize medfly larvae on the three most important host fruits occurring in the central-western fruit-producing region of Argentina. Such information verifies previous host fruit surveys, which found medfly larvae parasitized by *D. longicaudata* on figs and sweet oranges (unknown cultivars) [18] and on peaches (Elegant Lady variety) [15], following releases of this parasitoid throughout different fruit-growing valleys of San Juan. The fig, peach, and sweet orange are characterized mainly by their physical differences and are

widespread in all the irrigated fruit-growing valleys of the San Juan province. Figs and peaches are not only grown commercially in central-western Argentina, but they are also common fruit trees, like sweet orange cultivars, in backyard and home orchards throughout urban and rural areas for domestic consumption, or for making jams or jellies for local trade [14]. The performance of *D. longicaudata* on medfly larvae infesting the main pest multiplying hosts is highly relevant given the temporal overlapping availability of orange, fig, and peach during the summer and early autumn (December–March) [40]. This scenario enables medfly population growth in the fruit-growing region of Cuyo, mostly in urban areas. Thus, it is essential to apply integrated management strategies against the medfly that minimize the environmental impact involving safety techniques for human health and, at the same time, ensure maximum sustainability. In this context, the most advisable approach is biological control.

With regard to the second issue, the host fruit species had a strong influence on the ability of parasitoids to kill host larvae. Medfly mortality was appreciably higher on fig than on the other two tested fruit species, regardless of testing different parasitoid release densities in field cages. However, host mortality on peach clearly surpassed that on sweet orange. Particular physical characteristics of the fruit, e.g., rind thickness and flesh depth, appear to have been the main influences on the differences in medfly mortality rates found in the trials. A small fruit size, thin peel, and shallow flesh are often linked with a higher level of natural parasitism in a host fruit–frugivorous tephritid larva–parasitoid trophic relationship [21,22,26,41,42].

Based on the above, not surprisingly, the highest levels of medfly mortality due to the *D. longicaudata* ovipositional activity were found in figs. The fruit of this medfly host species showed very suitable features for making high-level parasitoid effectiveness in killing the pest possible. In this regard, the thin skin, which is about 50- and 3-fold thinner than those of orange and peach, respectively, with a flesh 1.6- and 1.3-fold shallower than those of orange and peach, respectively, were apparently the most advantageous physical features for parasitism. Such an assertion is also supported by the host larval density/cm² of the fruit surface area ratio, which was significantly similar among the three host fruit species, suggesting no influence on the parasitoid foraging activity on the fruits. A similar host density per cm² of fruit surface area balanced out the differences in fruit size, mainly characterized by fruit weight, diameter, and external fruit surface area. The physical features of the fig being propitious to parasitism may also be associated with the markedly high reproductive success of *D. longicaudata* females. However, the parasitoid offspring rate found in fig was not different from that recorded in peach but was highly different from orange. The *D. longicaudata* female also exhibited a high ability to seek, locate, and kill medfly larvae on peach, although the parasitoid performance was lower than that recorded in fig. Peach is a host fruit species that also has suitable physical properties to facilitate parasitoid ovipositional activity on frugivorous dipteran larvae [43]. The orange was the fruit species with both the lowest medfly mortality and parasitoid offspring rates compared to the other two-tested host fruit species. The lower values of both parameters may be mainly due to the thicker rind and deeper flesh, which further limit parasitism [21]. Excessive host larval crowding inside the fruit may also cause larval mortality [44]. The trials in the current study used a larval density based on maximum natural infestation levels of fig recorded in fruit-growing valleys of San Juan province. Such infestation rates ranged from 80 to 120 larvae per fruit, as fig is one of the fruit host species most highly infested by medfly [18]. The host larval densities of 100 larvae per fig and the equivalent of 130 larvae per peach and per orange tested in the trials are high fruit infestation levels, which may have caused additional medfly mortality.

In the current study, the medfly mortality in sweet oranges caused by *D. longicaudata* is highly relevant. *Citrus sinensis*, like other citrus species, provides a bridge for medfly during the cold season, the late autumn and winter, when peaches and figs are unavailable in central-western Argentina. Although a previous study showed that *D. longicaudata* females may be more attracted to non-infested peach volatiles than those of orange [45],

chemical stimuli likely did not play a major role in the medfly killing parasitoid capacity in the current study. The above is due to *D. longicaudata* females strongly preferring infested fruit based on chemical cues from the tephritid larvae feeding on rotting fruit substrate, regardless of host fruit species [46,47]. Therefore, any effect of volatiles emanating from a particular fruit species on the attraction of *D. longicaudata* females was likely homogenized due to the presence of medfly larvae in all fruit exposed to parasitoids.

The third finding showed a similar pattern of increasing both host mortality and parasitoid offspring in the three fruit species when the ratio of released *D. longicaudata* females to medfly larvae was gradually increased. Thus, the higher host-killing capacity and reproductive success achieved by *D. longicaudata* occurred at the highest ratios of parasitoid females released into the tube cages, i.e., 80 and 160. In this regard, 160 parasitoids released per experimental tube cage compared to 20 and 40 released parasitoids substantially increased medfly mortality 1.4- to 1.7-fold in peach, 1.5- to 1.6-fold in fig, and 1.6- to 1.9-fold in orange, taking into account the two-year study period. When comparing the two maximum parasitoid release densities (80 vs. 160), there was less difference between them than with the two lower parasitoid densities. However, there was a substantial increase of 7–8% in fig, 8–9% in peaches, and 9–14% in oranges towards 160 parasitoids released per tube cage. The last data are relevant because the increase to 160 parasitoid females released in the trials involving orange yielded the maximum effectiveness of the *D. longicaudata* in killing the medfly larvae infesting this citrus species. Similarly, the number of offspring produced by *D. longicaudata* increased at higher parasitoid female densities, reaching their maximum value at 160 parasitoids/cage tube density. However, the parasitoid reproductive success in fig and peach was substantially similar and higher than in orange, most likely due to the physical properties that favor parasitism in both fruit species. Such data from field cage natural conditions, i.e., restricting the parasitoid dispersal and host search area, suggest that medfly control may improve substantially with an increase in parasitoid release density per hectare in an infested fruit crop. Previous augmentative releases of *D. longicaudata* on fruit farms in the irrigated valleys of San Juan used parasitoid release densities averaging between 1600 and 2300 parasitoids/hectare with a sex ratio of 0.8–1.1 females–male [14,15]. Sulfite paper bags holding 500–600 parasitoids were used in those open-field parasitoid releases. These release bags were opened in sectors located along line transects within the farm. Both studies (14–15) recorded a decrease in the medfly population at the *D. longicaudata* release site, providing evidence of the effectiveness of augmentative biological control on farms with little or no conventional chemical treatment. However, based on the results of the current study, in which the maximum release density of *D. longicaudata* was the most successful in controlling medfly, the host-killing parasitoid capacity at higher open-field parasitoid release densities needs to be tested in future studies. Therefore, release densities above 2300 parasitoids/hectare for a more effective medfly control may be assessed.

5. Conclusions

The current study supports *D. longicaudata* as a biocontrol agent able to produce high mortality to medfly in all three primary host fruit species, such as fig, peach, and orange, which are key hosts responsible for growing and supporting medfly populations in all of the irrigated fruit-growing valleys of central-western Argentina. Particularly, high parasitoid release densities were found to be more effective in controlling a greater proportion of the medfly population. The capacity of *D. longicaudata* to kill medfly larvae infesting a *Citrus* species with more challenging physical characteristics for parasitism, like orange, is a potentially valuable trait when using this exotic parasitoid in open-field largescale releases. The results of the current study are critical for designing an area-wide medfly management approach in irrigated semi-arid fruit-producing areas of Argentina.

Supplementary Materials: The following supporting information can be downloaded at <https://www.mdpi.com/article/10.3390/agronomy14102418/s1>: Table S1: Host-killing parasitoid ca-

capacity_fruit species; Table S2: Host-killing parasitoid capacity_treatments; Table S3: Parasitoid reproductive success_fruit species; Table S4: Parasitoid reproductive success_treatments; Table S5: Parasitoid offspring sex ratio_fruit species; Table S6: Parasitoid offspring sex ratio_treatments; Database_Statistical Analysis S7: Overviews of statistical models applied for tested biological parameters_HK_PRS_POSR.

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