Domestication and mass rearing of the Mediterranean fruit fly, *Ceratitis capitata* (Diptera: Tephritidae) from Argan forest

R. ELAINI1,2, A. MAZIH1, M. ALONSO VALIENTE3, Y. RAHAL2

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Abstract

The Mediterranean fruit fly, *Ceratitis capitata* (Wiedemann) (Diptera: Tephritidae) is one of the major fruit crop pests in Morocco and worldwide. A successful and optimized mass rearing of this pest is a key component of several integrated pest management strategies. In this context, colonies of the Mediterranean fruit fly, *C. capitata* from field-collected pupae from infested argan fruits were established, maintained and studied in laboratory for 21 successive generations. Life history and demographic parameters were monitored over 14 generations. Afterward, the 20th and 21st generation were studied to investigate optimal pupal and eggs production respectively. Therefore, different eggs volumes varying from 0.25 to 3 ml were seeded on 1 kg larval diet and adult cages were populated with different pupae densities from 2000 to 20000. Our study highlights the potential adaptability of the Mediterranean fruit fly to laboratory conditions. In fact, from generation 7 onward all eggs are laid through the cages mesh instead of oviposition device. Egg production, pupal production, eggs eclosion rate, pupae emergence rate and larval development period was improved over different generations. Our data recommends the use of 1.25 to 1.5 ml of eggs per 1 kg diet and 16000 pupae per cage to optimize pupal and eggs production. Further experiments, especially regarding fitness, mating competitiveness and genetic variation, are required to investigate the stability of populations after additional captivity phases in laboratory.

Keywords: Mediterranean fruit fly, *Ceratitis capitata*, Tephritidae, Domestication, mass-rearing, wild strain.

INTRODUCTION

The Mediterranean fruit fly, *Ceratitis capitata*, is one of the major fruit crop pests in Morocco (Mazih, 2008). The fly originated in Africa, but today presents a world-wide geographic distribution, showing great colonizing potential (Diamantidis et al., 2009).

Efficient control of *C. capitata* requires, in addition to combining multiple control methods in an integrated pest management strategy, adequate information regarding the pest in relation to beneficial species, as well as the dynamics of these organisms under varying environmental conditions and factors, all within an economic framework for assessment of costs and benefits (Ditomasso et al., 2017). Mass and laboratory-rearing of insects is a key component of several integrated pest management strategies (Sorensen et al., 2012). Hence, several purposes of mass rearing are known: Production of raw material for secondary product (virus, pheromones, etc), production of hosts for entomophagous species, displacement, parasites and predators, sterile insect techniques (Boller,
1972). Also, laboratory-reared flies can be useful for insect biocconversion of waste (Van Huis, 2013) or for the study of problems concerning adaptation and colonization processes, in a more general biological approach (Souza et al., 1988). In addition, insects have a large potential for sustainably enhancing global food and feed production, and commercial insect production is a rising industry of high economic value (Jensen et al., 2017).

Mass production often focuses on increasing output and deliberate or inadvertent selection can result in traits that optimize rapid growth, shorter development, and reduced or even loss of diapause under artificial conditions (Grayson et al., 2015). Within a mass-rearing facility the most important factors for efficient production of larvae are high egg production, low mortality, and a large number of fertile adults in as little space as possible (Parry et al., 2017).

For many ecologically important traits, the performance of laboratory strains can differ from wild populations and the use of artificial diet may not reflect growth and development on natural diets in some contexts (Grayson et al., 2015). Demographic studies have been completed for several species of tephritids. Population parameters have been measured for wild strains of the Mediterranean fruit fly, C. capitata, several differences between collected flies reared for 12 generations and populations maintained for more than 12 years were observed (Rössler, 1975).

The objectives of this research were to implement a laboratory adapted strain of medfly and to optimize its mass rearing output (eggs and pupae). We therefore initiated a mass rearing of wild strain medfly, collected from infested argan fruits, to reach adapted colony able to be maintained and scaled-up in the laboratory. Life history and demographic parameters of the first generation, considered as wild, and the successive generations were recorded. Further trials carried out on 14th generation, considered as laboratory-adapted, to optimize its mass rearing outputs.

MATERIALS AND METHODS

Laboratory adapted strain implementation:

Wild populations were established using medfly collected from infested argan fruit in and around the Hassan II Institute of Agronomy and Veterinary Medicine, Agadir (30°35’ N, -9°47’ E). Dropped argan fruit was collected from the ground surrounding infested trees and was placed in perforated trays, which allowed emerging larvae to crawl through the holes and pupate in a sandy container below. Pupae were sieved from the sand daily and were maintained in sealed petri-dishes (200 Pupae/petri-dish) at 27°C±2 during 7 days. Wild adults eclosed in insect rearing cages, covered with mesh, with a density of about 16000 flies per cage. This constituted the first generation (G1) or wild type mentioned in the text.

For generations 1 and 2, because of pupae availability, three cages (replicates) were used instead of four cages for the following generations. Eggs collected indifferently from diverse cages constituted the next generation. Therefore, material from different cages was mixed and all cages constituted a single population (Zygouridis et al., 2014). Adult fly colonies were maintained in a controlled environment room (temperatures; 25±1 °C, RH: 60-65%, 12:12 light:dark cycle) and were provided a standard diet of enzymatic yeast hydrolysate and sugar (1:4) together with water ad libitum.

For the first six generations, cages (30 cm × 30 cm × 30 cm) were used. Eggs were collected either in the water trays that are located under the cages or using an oviposition device. Only eggs from oviposition device are seeded in the larval diet to produce pupae. The larval diet used were slightly modified from that developed by (Tanaka et al., 1969) and consist of: Bran (259 gr), Brewer’s yeast (65 gr), Sugar (148 gr), Sodium Benzoate (2.6 gr), HCl-0.9 % (100 ml) and water (385 ml).

These devices consisted of a plastic dome with numerous small holes through which the females laid their eggs. The plastic covers a 9 cm diameter Petri dish containing a wet filter paper. One mature argan fruit was placed inside the petri dish so that its odor stimulated egg-laying. Eggs were carefully removed with a fine brush and by flushing the underside of the dome and the filter paper with water. After the eggs had sunk in the water they are incubated for 48h and then pipetted and seeded on 1 kg of artificial diet for larval development (27–28 °C, 55% R.H.) in trays and are allowed to develop for 5-6 days (Alonso et al., 2005). Four trays (replicates) were used for each generation. When larvae begin to crawl up the side of the tray, the pupation medium is introduced underneath the larval tray and both are covered in netting to minimize escapees. Pupation trays are refreshed daily to allow synchronization of adult eclosion. 5 days post pupation the pupae are placed in cages containing food and water until eclosion.

At the 7th generation onward, when the colony seemed established, adults were maintained in 77 cm × 72 cm × 7 cm wooden cages with food and water. Oviposition devices are removed and all eggs are collected throw cages mesh. The cages were suspended on metallic trolleys.

Life history and demographic parameters:

Egg production

For generations 1 to 13, records were kept on the following: 1) Preovipositional period (d), 2) Harvest duration (d), 3) Total egg production (ml), 4) Eggs collected on oviposition device (%), 5) Total Eggs/female (estimated by assuming that 1 ml of eggs is equivalent of 22 100 based on previously attained data in our laboratories), 6) Eggs eclosion % (by counting a sample of 100 eggs placed on a moistened filter paper within a 90-mm Petri dish).

Pupal production

For generations 2 to 14, records were kept on the following: 1) Larval development time (d), 2) Pupal collection duration (d), 3) Total pupal production (ml), 4) Egg to pupae recovery (%) calculated based on the initial number of eggs introduced into each container of rearing medium and assuming that 100 produce pupae measure 1,5 ml and 5) Emergence (%) based on lots of 100 puparia from each replicate.
Laboratory-adapted strain mass rearing optimization

To evaluate laboratory adapted strain optimal egg production, the 20th generation was studied. Each cage was populated with different volumes of pupae; the expected male: female ratio was 1:1. Twelve cage adult population densities were analyzed ranging from 2000 to 24,000 adults per cage. Each evaluation was repeated six times for statistical replication. Egg collections started from day 6 post adult eclosion and proceeded until the colony decline.

To evaluate optimal pupal production, the 21st generation was studied. Different volumes of eggs (0.25-3.00 ml) were seeded per 1 kg of diet as described above. Six replicates for each egg volume tested were analyzed. Following pupation, the volume (ml) of pupae was measured.

Statistical analysis

Data were subjected to one-way analysis of variance (ANOVA) and, where appropriate, means were separated by Tukey’s range tests. P-values of ≤ 0.05 were considered statistically significant. Analysis was performed using the Minitab 16 software.

RESULTS

Life history and demographic parameters

Egg production

In order to reveal the plasticity of Mediterranean fruit fly, C. capitata to be adapted to laboratory controlled environment, wild strain was kept among different generations. Figure 1 shows the total volume of eggs (ml) obtained from cages populated with 16000 of wild pupae over different generations. Significant differences in total egg production were observed among the different generations (F=76.58; df=6; p<0.001) (Figure 1). It was clear that the total egg production increases significantly over generations. During the first three generations, the production fluctuated around 26 ml. After, it increased to slightly more than 50 ml for generations 5 to 7. The best yields were reached from generation 11 onward (more than 90 ml) with a peak value of 98±5.92 on generation 13.

Table 1 shows that the period from adults emergence and first egg production (Preovipositional period) did not differ over the generations (F=0.42; df=6; p=0.857). It oscillates 5.5 to 6 days. No statistically significant differences were observed among harvest durations for all generations (F=2.68; df=6; p=0.045). It’s comprised between 14 and 15.5 days. Regarding the eclosion percentages of the eggs, it was different among generations (F=5.28; df=6; p<0.001). The lowest value was observed for first generation (63.67 ± 2.91). However, this percentage was higher for the following generations (more that 70%).

Our data suggest that the percentage of eggs collected on the ovipositor device dropped significantly from a given generation to the next one (F=1218.89; df=6; p<0.001). In fact, the first generation was not adapted at all to lay eggs throw cages mesh. 100 % of eggs were collected on the oviposition device. However significant changes in oviposition option were observed in the following generation. Hence, the percentage of eggs collected on oviposition device dropped to 63.98 ± 2.05 and 43.69 ± 1.23 c for generations 3 and 5 respectively. From generation 7 onward, all eggs were produced throw cages mesh.

Another parameter that shows the adaptation of medfly to laboratory conditions is the female daily fecundity. Our data suggest that the differences among the produced Egg/female/day over different generations were highly significant (F=80.84; df=6; p<0.000). It’s clear that females of first generations produced less eggs/female/day (about 5 for generations 1 and 3). This parameter considerably increased from generation 5 onward. The best value of 17.44 ± 0.82 was recorded for generation 13.

Pupal production

Figure 2 shows the total volume of pupae (ml) obtained from 1 ml of eggs seeded into 1 kg larval diet over different generations. Significant differences in total pupal production were observed among the different generations (F=27.45; df=6; p<0.001) (Figure 2). The total pupal production increases over generations. In fact, 1 ml of second generation eggs produced 127.78±6.81 ml. This
volume reached 158.74±1.17 ml for generation 8. From Generation 10 onward, the pupal production exceeded the volume of 160 ml. Hence a peak of 175.74±3.37 was recorded for generation 13.

In table 2, larval development time shortened with about 2 days from generation 2 to generation 14 (F=7.25; dl=6; p<0.001). It moved from 13±0.00 days to 10±75 respectively. The pupal collection time remained unchanged for all generations (F=0.14; dl=6; p=0.988). Concerning the total pupal weight, it reflects the total pupal production and followed the same trend of increasing over generations (F=27.45; dl=6; p<0.001). Pupal recovery did significantly improve as it passed from 38.55±2.05 % days for generation 2 to 53.01±0.61 for generation 14 (F=27.45; dl=6; p<0.001). The quality of pupae, reflected by the emergence rate, was also significantly enhanced (F=36.80; dl=6; p<0.001). The first generations had an emergence rate inferior to 80 %. From generation 8 onward this rate exceeded the limit of 80 % reached 87.25±1.70 % for generation 14.

**Laboratory-adapted strain mass rearing optimization**

Figure 3 shows the total pupal production for different seeded eggs densities tested. Significant differences in total pupal production were observed among the different densities (F=121.70; dl=11; p<0.001). The highest pupal rearing efficacy was achieved with a volume of eggs comprised between 1.25 and 1.5 ml seed in 1 kg of larval diet. Egg volumes lower or higher than the optimum recorded produced fewer pupae. The coefficient of determination (R²) between the total pupal production and pupal eggs volume, was found to be 86% (y = -4.59x^2 + 56.44x + 5.55; R²=0.8617) indicating that the curve provides a good fit to the data.

Regarding egg production optimization, figure 4 shows that the total egg production differ significantly among the different pupae densities (F= 49.65; dl=11; p<0.001). The highest egg production was achieved with 16000 pupae per cage. Pupae densities lower or higher than the optimum recorded produced fewer eggs. The coefficient of determination (R²) between the total egg production and pupal density per cage, was found to be 96% (y = -1.6587x^2 + 26.063 x - 12.61; R²=0.9627) indicating that the curve provides a good fit to the data.

**DISCUSSION**

Our study highlights the potential adaptability of the Mediterranean fruit fly to laboratory conditions. In fact, there were improvements in the parameters such as egg production, pupal production, eggs eclosion rate, pupae

**Table 1: Egg production parameters over different generations of wild strain populated on cages with 16000 pupae (Mean ± SE; n=4). Data followed by the same letter, in the same column, do not differ significantly according to ANOVA and Turkey’s HSD test (P>0.05)**

<table>
<thead>
<tr>
<th>Generations</th>
<th>Preovipositional period (d)</th>
<th>Harvest duration (d)</th>
<th>Eggs collected on oviposition device (%)</th>
<th>Egg/female/day</th>
<th>Eggs eclosion %</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1</td>
<td>6.00 ± 0.00 a</td>
<td>14.33 ± 0.33 a</td>
<td>100.00 ± 0.00 a</td>
<td>5.02 ± 0.39 d</td>
<td>63.67 ± 2.91 b</td>
</tr>
<tr>
<td>G3</td>
<td>5.75 ± 0.25 a</td>
<td>14.50 ± 0.29 a</td>
<td>63.98 ± 2.05 b</td>
<td>5.50 ± 0.45 d</td>
<td>70.50 ± 2.96 ab</td>
</tr>
<tr>
<td>G5</td>
<td>5.50 ± 0.29 a</td>
<td>15.50 ± 0.29 a</td>
<td>43.69 ± 1.23 c</td>
<td>8.19 ± 0.39 c</td>
<td>72.25 ± 1.49 ab</td>
</tr>
<tr>
<td>G7</td>
<td>5.75 ± 0.25 a</td>
<td>15.25 ± 0.25 a</td>
<td>0.00 ± 0.00 d</td>
<td>9.45 ± 0.43 c</td>
<td>75.50 ± 2.10 a</td>
</tr>
<tr>
<td>G9</td>
<td>5.75 ± 0.25 a</td>
<td>15.25 ± 0.25 a</td>
<td>0.00 ± 0.00 d</td>
<td>14.57 ± 0.42 b</td>
<td>75.50 ± 1.94 a</td>
</tr>
<tr>
<td>G11</td>
<td>5.50 ± 0.29 a</td>
<td>15.25 ± 0.25 a</td>
<td>0.00 ± 0.00 d</td>
<td>16.48 ± 0.79 ab</td>
<td>74.50 ± 0.87 a</td>
</tr>
<tr>
<td>G13</td>
<td>5.75 ± 0.25 a</td>
<td>15.50 ± 0.29 a</td>
<td>0.00 ± 0.00 d</td>
<td>17.44 ± 0.82 a</td>
<td>78.50 ± 0.65 a</td>
</tr>
</tbody>
</table>

**Figure 2: Comparison of pupal production (ml) for 1 ml of eggs seeded into 1 kg larval diet over different generation of wild strain. Mean volume of pupae produced is shown for each generation (mean ± SE; n = 4). Mean values labelled with the same letter are not significantly different according to ANOVA and Tukey’s HSD test (P > 0.05)**
emergence rate and different stages development time. In addition of these parameters, the main trait showing the adaptability of the strain is its ability to lay all eggs through the cages mesh instead of oviposition device. In fact, from generation 7 onward all eggs are collected from the mesh. Hence, for an insect population to adapt to a new diet, the most important behavioral modifications requires the females to oviposit on the new food (Bravo and Zucoloto, 1998). Leppla et al. (1983) reported that the Mediterranean fruit fly (medfly) \textit{C. capitata} required more than 5 generations to strain adaptation. Methods for insect rearing on artificial diet and the availability of laboratory strains have enabled a wide range of studies that could otherwise be infeasible (Grayson et al., 2015). The lack of success during the early stages of colonization suggests that the wild insects have to be adapted to laboratory conditions and that intense selection occurs in the F1 and following generations. However, it was difficult to determine if all insects have contributed to the process of adaptation during the first generations (Boller, 1972). For example, Selection on strains of the gypsy moth, \textit{Lymantria dispar} (L.), maintained on laboratory, has enhanced a variety of traits, resulting in faster development, shorter diapause, and greater fecundity (Grayson et al., 2015). Also, the laboratory strain of Caribbean fruit flies \textit{Anastrepha suspensa} (Loew) (Diptera:Tephritidae), began mating at an earlier age and mated more rapidly is small screen cages in the laboratory conditions (Mazomenos et al., 1977)

Our results demonstrate that the preoviposition period (about 6 days) and oviposition period (about 15 days) remained unchangeable over generations. The same observation was reported bay (Diamantidis et al., 2011) for Kenyan flies. Assuming that the 11th generation onward could be considered as laboratory adapted strains, these results are different from (Foote and Carey, 1987) and (Vargas and Carey, 1989) that reported that females in the laboratory population produced eggs at a significantly younger age than those in the wild populations.

For egg production, the total egg production increases significantly over generations. This finding is confirmed by different authors. Vargas and Carey (1989) found that egg production by the laboratory population was

### Table 2: Pupal production parameters for 1 ml of eggs seeded into 1 kg larval diet over different generation of wild strain. Mean volume of pupae produced is shown for each generation (mean ± SE; n = 4). Mean values labelled with the same letter are not significantly different according to ANOVA and Tukey’s HSD test (P > 0.05).

<table>
<thead>
<tr>
<th>Generations</th>
<th>Larval development time (d)</th>
<th>Pupal collection duration (d)</th>
<th>Total pupal weight (g)</th>
<th>Egg to pupae recovery (%)</th>
<th>Emergence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>G2</td>
<td>13.00 ± 0.00 a</td>
<td>3.33 ± 0.33 a</td>
<td>85.19 ± 4.54 d</td>
<td>38.55 ± 2.05 d</td>
<td>62.00 ± 1.15 d</td>
</tr>
<tr>
<td>G4</td>
<td>12.75 ± 0.25 ab</td>
<td>3.50 ± 0.29 a</td>
<td>94.49 ± 2.91 cd</td>
<td>42.76 ± 1.32 cd</td>
<td>72.00 ± 0.82  c</td>
</tr>
<tr>
<td>G6</td>
<td>12.25 ± 0.25 ab</td>
<td>3.25 ± 0.25 a</td>
<td>99.45 ± 1.26 bc</td>
<td>45.00 ± 0.57 bc</td>
<td>80.00 ± 1.41 b</td>
</tr>
<tr>
<td>G8</td>
<td>12.50 ± 0.29 ab</td>
<td>3.50 ± 0.29 a</td>
<td>105.83 ± 0.78 b</td>
<td>47.88 ± 0.35 b</td>
<td>86.50 ± 2.47 ab</td>
</tr>
<tr>
<td>G10</td>
<td>12.00 ± 0.00 abc</td>
<td>3.25 ± 0.25 a</td>
<td>108.52 ± 1.28 ab</td>
<td>49.10 ± 0.58 ab</td>
<td>84.98 ± 0.71 ab</td>
</tr>
<tr>
<td>G12</td>
<td>11.50 ± 0.29 bc</td>
<td>3.50 ± 0.29 a</td>
<td>116.31 ± 1.27 a</td>
<td>52.63 ± 0.58 a</td>
<td>87.25 ± 1.25 a</td>
</tr>
<tr>
<td>G14</td>
<td>10.75 ± 0.48 c</td>
<td>3.50 ± 0.50 a</td>
<td>117.16 ± 2.25 a</td>
<td>53.01 ± 0.61 a</td>
<td>87.25 ± 1.70 a</td>
</tr>
</tbody>
</table>

Figure 3: Comparison of pupal production (ml) for different egg volumes seeded into 1 kg larval diet over different generation of wild strain. Mean volume of pupae produced is shown for each generation (mean ± SE; n = 6). Mean values labelled with the same letter are not significantly different according to ANOVA and Tukey’s HSD test (P > 0.05)
significantly greater than by all wild populations. The laboratory-adapted strain produced about 5 times more eggs which is the same relation reported by Bravo and Zucoloto (1998).

The number of egg laid per female increased from 5.02 ± 0.39 for generation 2 to 17.44 ± 0.82 for generation 13. Neto et al. (2012) reported that in similar large-scale experiments, 18,000 VIENNA-8 TSL (Temperature Sensitive Lethal) adults per cage, yielded 7.54 ± 1.70 eggs per female per day, for a period of 10 days. Blay and Yuval (1999) reported the greatest number of eggs laid by a mated female per day to be 94 eggs.

With respect to the hatch rate of eggs collected from the mass-rearing cages, it was slightly improved over generation. During the 8 generations tested, Rempoulakis et al. (2016) found this parameter to remain stable.

There was significant improvement of quantitative and qualitative parameters such as pupal production, pupal emergence rate and larval development period during domestication process of the Mediterranean fruit fly in laboratory conditions, This is due to that fact that at least a certain percentage of immature accepted the new food and ingest it in adequate amounts (Bravo and Zucoloto, 1998). Our findings are confirmed by Vargas and Carey (1989) that reported that the duration of the larval stage differed significantly between laboratory and wild lines of flies. In fact, C. capitata is known for its plasticity that permits their adaptation to new habitats, reflected in its present world-wide geographical distribution (Souza et al., 1988).

The improvement of mass rearing performances should not lead to strains completely different from the initial one. Rodriguero et al., (2002) reported that copulatory success might be affected between wild and laboratory adapted strain, as a side effect due to selection of wild males over lab males due to differential sexual activity or other causes. Indeed, Certain level of sexual incompatibility was observed between strains under both laboratory and field conditions of the Mexican fruit fly, Anastrepha ludens (Loew) (Meza-Hernandez and Diaz-Fleischer, 2006). To counter any unwanted changes, there is a need to gain knowledge of the basic biology of the organism in question, including the basic population genetics and the effects of temperature, diet, and density (Jensen et al., 2017). It is therefore of fundamental importance to follow the adaptation process genetically and try and link the strain’s mating efficiency and ‘wild’ character with genetic markers (Zygouridis et al., 2014)

CONCLUSION

In conclusion, our study shows that wild Mediterranean fruit fly, C. capitata collected from infested argan fruits were well established in laboratory. Different population traits, such as egg production, pupal production, eggs eclosion rate, pupae emergence rate and larval development period, were improvement over the fourteen studied generations. Once adapted the laboratory conditions, the optimization of the mass rearing parameters is required as it influences insects quality and production cost. Our findings provide important insights of the optimal insects’ densities to use for eggs and larval production. Obviously, further studies are required to characterize occurring changes over different generations.

ACKNOWLEDGEMENTS

We thank Prof. Josep Anton Jaques Miret for his valuable advises to speed up the strain adaptation process.

Figure 4: Comparative analysis on the total volume of eggs (ml) obtained from cages populated with varying amounts of wild strain pupae (mean ± SD; n = 6). Bars superscripted with different letters are significantly different according to ANOVA and Tukey’s HSD test (P<0.05)
REFERENCES


