Effects of delayed mating on the reproductive biology of the vine mealybug, Planococcus ficus (Hemiptera: Pseudococcidae)

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1	Effects of delayed mating on the reproductive biology of the vine mealybug, <i>Planococcus</i>
2	ficus (Hemiptera: Pseudococcidae)
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13 The effect of increasing mating delay on the reproductive performance and population growth 14 rates of the vine mealybug, *Planococcus ficus* (Signoret) (Hemiptera: Pseudococcidae), was 15 investigated under laboratory conditions. Virgin females were mated at 1, 3, 5, 7, 14, 21 and 16 28 days after emergence and reproductive and life table parameters were estimated. The pre-17 oviposition period (number of days between mating and the onset of oviposition) significantly 18 decreased in females mated within 7 days, whereas females mated at older ages showed 19 equivalent pre-oviposition periods (< 4 days). The length of the oviposition period did not 20 vary with increasing age at mating. Female longevity significantly increased in females mated 21 at 21 and 28 days, as a consequence of a longer pre-reproductive period. Fecundity and sex 22 ratio were not affected by the female age at mating, whereas fertility was higher in mealybugs 23 mated at older ages. Additional field observations highlighted that young and old virgin 24 females were equally able to attract males, as both mated on the same day as the field release. 25 Mating delay also affected the life table parameters of P. ficus, as the intrinsic and finite rates 26 of increase did not differ in mealybugs mated within 7 days and significantly decreased in 27 females mated at older ages. The mean generation time and the population doubling time 28 were overall similar in females mated at 1-7 days, and increased significantly in females that 29 experienced longer mating delays. In terms of the mating disruption control of P. ficus, our 30 findings indicate that this method would be effective if mating is delayed > 7 days, as shorter 31 delays in mating did not reduce the population growth rates.

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33 Keywords: mating delay; fecundity; fertility; life history; population growth parameters;
34 mating disruption.

36 Introduction

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The vine mealybug, *Planococcus ficus* (Signoret) (Hemiptera: Pseudococcidae), is found throughout Central and South America, California, the Mediterranean basin, South Africa and the Middle East. It completes a variable number of generations per year, depending on the climatic conditions. In the Mediterranean basin, the annual generations of *P. ficus* range from three to four in Italy (Duso, 1989; Lentini *et al.*, 2008) to six in Tunisia (Mansour *et al.*, 2009). In contrast, four to six generations per year have been reported in California (Millar *et al.*, 2002) and five to six generations in South Africa (Walton & Pringle, 2004).

This pest causes increasing yield losses in wine and table grape vineyards (Daane *et al.*, 2012). The conventional control strategy against *P. ficus* consists of multiple applications of insecticides in spring-summer, in order to reduce the population density and avoid fruit damage. However, the chemical control of *P. ficus* is often of limited effectiveness, as mealybugs reside above all underneath leaves, under bark and in roots, being thus protected from contact insecticides (Walton *et al.*, 2004).

The identification and synthesis of the *P. ficus* sex pheromone (Hinkens *et al.*, 2001) have facilitated the development of monitoring (Millar *et al.*, 2002; Walton *et al.*, 2004) and mating disruption control strategies. The pheromone-mediated control of *P. ficus* has been successfully applied in California and the Mediterranean basin using a microencapsulated formulation and reservoir dispensers (Walton *et al.*, 2006; Cocco *et al.*, 2014; Sharon *et al.*, 2016).

57 High sex pheromone concentrations influence insect mating behaviour in several ways, 58 including the reduced responsiveness of males to pheromone stimuli due to adaptation or 59 habituation, the delayed or diminished response of males due to the continuous presence of 60 pheromones, false trails by synthetic pheromone sources that divert males from calling 61 females, and modifications in the ratios of components of the natural sex pheromone blend 62 which lead to a sensory imbalance effect (Bartell, 1982; Barclay & Judd, 1995; Cardè & 63 Minks, 1995). Irrespective of the modes of action, the effectiveness of mating disruption can 64 be evaluated by assessing damage reduction or variations in the reproductive and 65 demographic parameters of natural pest populations or sentinel females artificially deployed 66 in the field (Baker & Heath, 2005). Studies on insect demography show that the rate of 67 population increase is affected not only by fecundity and fertility but also by the age at first 68 mating (Carey, 1993). Therefore, any factor causing a delay in mating has a significant 69 influence on population dynamics.

70 The effects of delayed mating on the reproductive output have been studied in several 71 lepidopteran species, generally highlighting a higher longevity and pre-oviposition period and 72 a lower fecundity and fertility with increasing female age at mating (Proshold, 1996; 73 Fadamiro & Baker, 1999; Jones & Aihara-Sasaki, 2001; Torres-Vila et al., 2002; Jiao et al., 74 2006). However, the delay of mating differentially affects the biological parameters of the 75 investigated species, thereby altering their fitness (Mori & Evenden, 2013). The decrease of 76 the target pest population density can also be due to a delay of mating encounters rather than 77 the suppression of matings, as observed in the European corn borer, Ostrinia nubilalis 78 (Hübner) (Lepidoptera: Crambidae), subjected to mating disruption (Fadamiro et al., 1999).

Delayed mating has also been reported to affect the reproductive output of females in other orders of insects. In the oriental beetle, *Anomala orientalis* (Waterhouse) (Coleoptera: Scarabaeidae), the fecundity gradually decreased with increasing female age at mating (Wenninger & Averill, 2006). Similarly, the solenopsis mealybug, *Phenacoccus solenopsis* Tinsley (Hemiptera: Pseudococcidae), mated 30 days after emergence, exhibited a decreased fecundity and pre-oviposition period compared to females mated at the age of two days (Huang *et al.*, 2013). 86 The influence of delayed mating on the reproductive biology of *P. ficus* has not previously 87 been explored. In the congeneric citrus mealybug, Planococcus citri (Risso) (Hemiptera: Pseudococcidae), delayed mating and high temperature modified the sex ratio and fecundity 88 89 (Nelson-Reese, 1961; Ross et al., 2010). To our knowledge, no specific studies have been carried out on the factors influencing the P. ficus sex ratio. On the other hand, the 90 91 reproductive biology of *P. ficus* has been studied in detail by Waterworth et al. (2011), who 92 demonstrated that it cannot reproduce parthenogenetically, even though virgin females are 93 capable of laying unviable eggs. The same authors also reported a very long lifespan of 94 unmated females (94 days), whereas males lived approximately 4.5 days and mated multiple 95 times per day.

A deeper understanding of the reproductive biology of *P. ficus* is required in order to evaluate the influence of mating disruption on its populations. Laboratory and field experiments were thus carried out to investigate the effects of increasing delays in mating on reproductive and demographic parameters of *P. ficus*.

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101 Materials and methods

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103 Insect rearing and source of adults

The stock colony of *P. ficus* originated from ovipositing females collected in infested vineyards in north-western Sardinia (Italy). It was maintained on sprouted potato tubers (cv. Spunta) in constant darkness at 22-25 °C and 60-70% RH. Species-specific identification was initially confirmed by molecular analysis (Demontis *et al.*, 2007) and is routinely tested approximately every six months to rule out the potential contamination of *P. citri*, which is morphologically very similar to *P. ficus*.

110 Males and females used in the experiments were obtained from eggs of the same age. Thus, 111 approximately 80 mated females at the onset of oviposition were isolated in plastic cylinders 112 (8 cm tall, 6 cm in diameter) sealed with a double layer of paper napkins and a rubber band. 113 At daily intervals, the oviposited eggs were gently moved with a soft sable-hair brush to 114 sprouted potatoes placed inside Tupperware containers $(13.5 \times 13.5 \times 6.5 \text{ cm})$ covered with 115 ventilated lids. Containers were marked with the date of egg collection and stored in a growth 116 chamber kept at 25 °C, 50% RH, 12L:12D photoperiod. Since the preimaginal development 117 time was approximately 30 days (see Results), mealybugs were observed daily after 20 days 118 under a dissecting microscope equipped with an ocular micrometer in order to separate 119 immature males and females. Mature third-instar nymphs, approximately 1.2 mm long, were 120 isolated on a sprouted potato (weight range 25-60 g) inside a plastic cylinder (8 cm tall, 6 cm 121 in diameter), whereas male prepupae were moved to paper towel strips and placed in plastic 122 vials (5.5 cm tall, 1.2 cm in diameter). Vials and cylinders were kept in the growth chamber 123 and checked daily until male and female emergence, the dates of which were recorded. Adult 124 mealybugs from each stock colony were randomly assigned to the different treatments.

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126 Delayed mating experiment

127 In order to evaluate the influence of delayed mating on the reproductive output of P. ficus, the 128 following reproductive parameters were investigated: i) length of the pre-oviposition period, 129 from mating to the onset of oviposition; ii) length of the pre-reproductive period, from adult 130 emergence to the onset of oviposition, which also corresponds to the sum of the female age at 131 mating and the pre-oviposition period; iii) length of the oviposition period; and iv) total 132 longevity of females. In addition, the following were also recorded: daily fecundity, i.e. the 133 number of eggs oviposited by each female; fertility, calculated as the percentage of viable 134 eggs over the total oviposited eggs; sex ratio of offspring, calculated as the percentage of female offspring; and survival. Experiments were carried out in the above-described growth
chamber at 25 °C, which is the optimal developmental temperature of *P. ficus* (Walton &
Pringle, 2005).

In order to allow mating, virgin females at ages 1, 3, 5, 7, 14, 21 and 28 days were individually paired at the beginning of the photophase with two 3-day-old virgin males inside the plastic cylinders and were observed until mating occurred. At least 25 females were tested for each age at mating. After mating, females were checked daily until the onset of oviposition, after which eggs were collected daily with a sable-hair brush until female death and were gently placed inside plastic cylinders (4.0 tall, 3.5 cm in diameter). Dates of mating, beginning and end of oviposition, and death of each female were recorded.

The offspring sex ratio was preliminarily determined in 10 females mated 1, 7 and 28 days after emergence following the method described by Ross et al. (2010) for *P. citri* eggs. Since there was no significant difference in the sex ratio among the tested treatments (see Results), the sex ratio of other treatments was not evaluated. Therefore, the mean value (59% female offspring) was used in the calculation of the life table parameters.

Fertility was assessed in 10 females mated 1, 7, 14, 21 and 28 days after emergence by collecting the oviposited eggs with a sable-hair brush every 2 days. Eggs were placed in cardboards inside sealed plastic cylinders (4.5 tall, 3.5 cm in diameter) and stored for two months under natural laboratory conditions. Fertility was determined as the percentage of egg eclosion, calculated as [first-instar nymphs / (first-instar nymphs + unhatched eggs)] ×100.

The ability of *P. ficus* females of different ages to attract and mate with wild males was investigated in field observations. Thus, 20 females at ages 1, 7 and 28 days were reared in the laboratory as described previously, and released in a naturally-infested vineyard. Females were placed individually on a sprouted potato inside a plastic cylinder (8 cm tall, 6 cm in diameter), which had 4 holes on the upper part of the wall (1.5 cm in diameter) to allow male 160 access. The positions of the three treatments were alternated in the central five rows of a 0.5-161 ha plot in a commercial vineyard (18 years old, cv. Carignano) (for further details about the experimental vineyard, see Muscas et al., 2017) and spaced approximately 3 m along and 162 163 within rows. Cylinders were sheltered from direct sunlight and rain by laminated cardboard, 164 and hung from trellis wires inside the canopy. In order to prevent ants and wild females from 165 entering the cylinders, the support wires were coated with insect trapping glue. Each female 166 was checked daily in order to estimate the length of the pre-oviposition period, given by the 167 time interval from the field release to the beginning of oviposition. Mating was assumed to 168 occur on the same day as the release. Females accidentally injured during the daily checks 169 were excluded from the analysis.

170

171 Data analysis

172 Reproductive parameters (fecundity, fertility, sex ratio, pre-oviposition period, pre-173 reproductive period, and oviposition period) and female longevity were compared among 174 treatments by a generalized linear mixed model, setting normal and binominal distributions 175 for numerical and percentage data, respectively. Significantly different means were separated 176 by Tukey's multiple comparison test (P < 0.05) (PROC GLMM, SAS Institute, 2008).

177 Life tables for females mated at different ages were built from age-specific fecundity (m_x) and 178 survival (l_x) rates, which were calculated from reproductive and survivorship data. The 179 following population growth parameters were estimated (Carey, 1993; Maia *et al.*, 2000):

- 180 net reproductive rate, $R_0 = \sum (l_x m_x)$;
- 181 intrinsic rate of increase (r_m), obtained by the iterative solution of the equation $\sum e^{-r_m x} l_x m_x = 1$;
- 182 finite rate of increase, $\lambda = e^{r_m}$;
- 183 mean generation time, $T_G = lnR_0 / r_m$;
- 184 population doubling time, $DT = ln2 / r_m$.

The mean values of the life table parameters were estimated with the jackknife method using a Microsoft Excel 2007 spreadsheet (Microsoft, Redmond, WA, USA), which generates pseudo-values of the investigated parameter through a subsampling replication technique (Meyer *et al.*, 1986). Jackknife pseudo-values were compared using analyses of variance (Bari *et al.*, 2015), followed by Tukey's tests to separate means (P < 0.05) (PROC GLMM, SAS Institute, 2008).

191

192 **Results**

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194 The preimaginal development time did not vary significantly among female groups ($F_{6,160}$ = 195 1.88, P = 0.189), ranging from 29.36 \pm 0.41 to 30.71 \pm 0.35 days (mean \pm SE, data not 196 shown). The duration of the pre-oviposition period decreased significantly in females mated 197 within 7 days, whereas 14-, 21- and 28-day-old mated females exhibited a non-significantly different pre-oviposition period (< 4 days) ($F_{6,160} = 41.95$, P < 0.001) (Table 1). The pre-198 199 reproductive period, also on account of being the female age at mating plus the pre-200 oviposition period, showed a significantly increasing pattern with increased mating delay 201 $(F_{6,160} = 614.78, P < 0.001)$ (Table 1). The length of the pre-reproductive period increased by 202 about two days from 1- to 7-day-old females and then increased proportionally by about 14 203 days in females mated from 14 to 28 days. The oviposition period ranged from 13.04 to 16.33 204 days revealing a slight, yet not significant, decrease in females mated 14, 21 and 28 days after 205 emergence ($F_{6,160} = 1.42$, P = 0.212). Females mated at 21 and 28 days lived significantly longer than those mated at younger ages ($F_{6,160} = 29.83$, P < 0.001), with the total lifetime 206 207 being modified above all by differences in the pre-reproductive period among treatments 208 (Table 1).

Females mated within 7 days showed similar daily oviposition patterns, as peaks were reached 5-7 days after the beginning of oviposition and ranged from 27.54 to 33.78 eggs/day (Fig. 1). In contrast, females mated at older ages reached higher peaks (range 50.08-63.68 eggs/day) in a shorter time (4-5 days). Only 1-day-old females laid fewer than 300 eggs, whereas the mean fecundity ranged from 302.5 to 355.6 eggs in females mated at older ages. However, no significant difference was revealed among groups ($F_{6,160} = 1.41$, P = 0.212).

215 The fertility of the investigated groups (females mated at 1, 7, 14, 21 and 28 days) was 216 87.51% and 85.86% in females mated at 1 and 7 days of age, respectively, whereas it was above 90% in the other groups (range 90.74-94.00%) (Fig. 2). However, a significant 217 218 difference appeared only between females mated at 1 and 7 days compared with those mated at 28 days ($F_{4,45} = 4.60$, P = 0.010), suggesting that older females had higher reproductive 219 performance than young mated females. The percentage of female offspring did not vary 220 221 significantly among the investigated ages at mating, showing a predominance of female offspring (range = 57.3-62.6%) (F_{2.27} = 0.90, P = 0.419). 222

223 The effects of the delayed mating on reproductive traits of P. ficus also affected the life table 224 parameters of cohorts originating from females mated at different ages. The net reproductive rate was significantly higher in females mated at 5 and 21 days post-emergence, whereas 225 those mated at 1 day showed the lowest rate ($F_{6,160} = 879.58$; P < 0.001) (Table 2). The 226 227 intrinsic and finite rates of increase did not differ significantly among 1- to 7-day-old groups, 228 whereas it decreased significantly in the other treatments (r_m : F_{6,160} = 600.37; P < 0.001; λ : $F_{6,160} = 776.01$; P < 0.001). The mean generation time ranged from 43.50 to 46.93 days in 229 230 females mated at 1-7 days, showing a significant increase in females mated at older ages (14-28 days) (F_{6,160} = 660.28; P < 0.001). The doubling time exhibited a pattern similar to T_G, 231 232 with values of the same magnitude in females mated at 1-7 days and significantly higher values in females mated 14-28 days after emergence ($F_{6,160} = 30.12$; P < 0.001). 233

Females released in the field 1 day after emergence showed a pre-oviposition period of 11.67 days, which was significantly longer than that of 7- and 28-day-old females (3.60 and 3.84 days, respectively) ($F_{2,54} = 112.24$; P < 0.001) (Fig. 3). The lengths of pre-oviposition periods confirmed the laboratory findings, showing that females of different ages equally attracted males and mated on the same day as the field release.

239

240 **Discussion**

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The present study illustrates the reproductive traits of P. ficus females mated at increasing 242 243 ages and indicates the influence of delayed mating on the population growth potential of this 244 pest. The preimaginal development time of mealybugs reared on sprouted potatoes was about 30.5 days at 25 °C, similarly to the findings of Walton & Pringle (2005) in mealybugs reared 245 246 on fresh grapevine leaves. The significantly longer development time observed by Varikou et 247 al. (2010) in *P. ficus* immatures developed on grape leaf discs (43.3 days at 25 °C) is likely 248 due to the different rearing substrate. Because the development times of P. ficus on sprouted 249 tubers and fresh grape leaves were similar, it is likely that phloem-sap feeders, such as 250 mealybugs, have a feeding preference for fresh substrate over excised plant organs. In 251 addition, the grape cultivar and possible genetic differences among the mealybug populations 252 has been shown to affect the preimaginal development time (Varikou et al., 2010).

Planococcus ficus females can mate shortly after emergence (Waterworth *et al.*, 2011), although a variable time interval before the beginning of oviposition is required for ovarian maturation. Time intervals between mating and the beginning of oviposition (pre-oviposition period) decreased in females mated within 7 days and were almost equivalent in older females. This trend has also been observed in *Ph. solenopsis*: females mated at 30 days showed a significantly shorter pre-oviposition period than that of females mated 2 days after 259 emergence (Huang et al., 2013), thus suggesting that newly-emerged females have a longer 260 pre-oviposition period than older females. This phenomenon can be explained by the egg 261 production and oviposition within the female reproductive tract (Bloch Qazi et al., 2003). 262 Oocytes are firstly generated within the ovaries (oogenesis) and then released in the oviducts (ovulation), through which eggs move down into the common oviduct, where they are 263 264 fertilized and finally laid. Virgin females retain oocytes in the ovaries and ovulation starts 265 after mating. In P. citri, meiotic ovarioles become visible 6 days after female emergence 266 (Nelson-Rees, 1961), which means that meiosis initiates after at least 6 days even if females mate on emergence. Subsequent ovulation, fertilization and oviposition may require a fairly 267 268 constant period of time (about 3-4 days). Therefore, in our opinion, P. ficus virgin females 269 mated at age \geq 7 days have almost completed the obgenesis of a portion of eggs, and exhibit a 270 nearly constant and slightly longer pre-oviposition period than 3 days.

271 The progressive decrease of the pre-oviposition period in females mated within 7 days 272 affected the length of the pre-reproductive period. In fact, in our study, the pre-reproductive 273 period increased slightly in females mated within 7 days, whereas, in contrast, it increased 274 proportionally with the mating delay in females mated at 14, 21 and 28 days. A prereproductive period similar to P. ficus females mated at 1 day has been found in both the 275 276 passionvine mealybug, Planococcus minor (Maskell) (Hemiptera: Pseudococcidae), and P. 277 citri. These species showed a pre-reproductive period of 10.2 and 12.3 days when reared at 25 278 °C on sprouted potato and grapevine, respectively (Francis et al., 2012; da Silva et al., 2014). 279 However, no information is available on the pre-reproductive period in mealybugs mated at 280 older ages.

The reproductive performance of *P. ficus* females mated at increasing ages differed considerably from that of other insects. A delayed mating of 4-6 days in some lepidopteran species has led to a significant reduction of fecundity and fertility (Vickers, 1997; Jones & 284 Aihara-Sasaki, 2001; Walker & Allen, 2011), whereas P. ficus has not been found to be 285 negatively affected by the age at mating. In fact, in our experiment, P. ficus fecundity was 286 267.4 eggs in 1-day-old females, ranging instead from 302.5 to 355.6 eggs in females mated 287 at older ages. No studies have been carried out on P. ficus egg production. However, it is 288 reasonable to assume that P. ficus oogenesis is similar to P. citri, for which Nelson-Rees 289 (1961) observed the highest fecundity in 10-20-day-old females and the decay of ovarioles 290 and egg resorption in 30-day-old virgin females. The same author demonstrated that P. citri is 291 a capital breeder, as starved newly-emerged females had a reduced body size and 292 undeveloped ovarioles.

293 The different performance of mealybugs relative to other systematic groups could be due to 294 their different evolution. In fact, many lepidopteran species enter in diapause during the 295 winter months and have a short adult lifespan and reproductive period. For instance, the 296 autumn gum moth, Mnesampela privata (Guenée) (Lepidoptera: Geometridae), exhibits 297 continuous ovulation after adult emergence and both fecundity and fertility are linearly related 298 to longevity (Walker & Allen, 2010). Unmated M. privata females live about 10 days; 299 therefore, a mating delay of 7 days reduces the time available for oviposition by 70% (Walker 300 & Allen, 2011). In contrast, P. ficus does not enter into diapause but slows down its life cycle, 301 overwintering mainly as mated adult females and preserving the reproductive potential for 302 several months (Lentini et al., 2008). In fact, overwintering females are usually larger and 303 have higher fecundity than summer females (A. Lentini, personal observation). The minimum 304 development threshold temperature of insects is lower than the reproductive threshold 305 (Gilbert & Raworth, 1996). Therefore, mealybug adult females at low temperatures can feed 306 and increase their 'energy reserve' in order to produce a high number of eggs under 307 favourable climatic conditions.

308 Contrary to findings in Lepidoptera, the fertility of *P. ficus* females did not decrease with age, 309 in fact females mated at 28 days showed the highest fertility values. Similarly, the fertility of 310 *A. orientalis* was not affected by the age at mating, although about 50% of females mated at 311 older ages did not lay fertile eggs (Wenninger & Averill, 2006).

312 Age of mating did not affect the offspring sex ratio, which was about 59% female regardless 313 of the mating delay. The mean sex ratio observed is in accordance with our previous 314 observations carried out in over 11,000 P. ficus eggs (female percentage = 60.3%) (Cocco et 315 al., 2015). To date, no studies have reported variations in the P. ficus sex ratio due to the age 316 of females at mating, whereas research carried out on P. citri provided contrasting results. 317 Ross et al. (2011) observed nearly 50% of female eggs in young mated females and about 318 60% in females mated after 30 days. On the other hand, Nelson-Rees (1961) found a 319 prevalence of female offspring among females mated within 20 days and a majority of male 320 offspring among 30-day-old females. Sex ratio in P. citri and, by extension, in P. ficus is 321 likely influenced not only by female age but also by environmental factors such as food 322 quality, feeding frequency, and population density (Ross et al., 2010, 2011). In addition, 323 climatic conditions may play a role in biasing the sex ratio, as Walton & Pringle (2005) observed a lower proportion of females at 18 °C and 30 °C under laboratory conditions. This 324 325 could be an adaptive response to adverse environmental conditions in order to ensure higher 326 genetic variability, as reported in some mite species (Margolies & Wrensch, 1996; Rencken & 327 Pringle, 1998).

The findings of our field experiment were in accordance with our laboratory results. In fact, mating delay did not impair the calling behaviour of *P. ficus* females and their mating ability, as females of different ages mated on the same day as the field release. Similarly, the female age at mating in the European grapevine moth, *Lobesia botrana* Den. & Schiff. (Lepidoptera: 332 Tortricidae) did not alter the calling activity of females and their receptivity (Torres-Vila *et*333 *al.*, 2002).

334 Female mating at different ages affected all the investigated demographic parameters. The 335 intrinsic and the finite rates of increase showed a decreasing pattern in females mated at 14-28 336 days, whereas the population doubling time increased when mating was delayed for more than 337 14 days. These patterns indicate a lower growth potential of *P. ficus* populations at increasing 338 ages at mating. Increasing mating delay also led to a lengthening of the mean time required to 339 complete a generation. The net reproductive rate did not show a clear pattern and it is not 340 clear why females mated at 5 and 21 days exhibited the highest values. However, overall the 341 demographic parameters showed a consistent pattern with the finding that a mating delay > 7342 days reduces the population growth potential.

343 Biological and reproductive parameters of P. ficus are affected by several factors, including 344 the host species (Ahmed & Abd-Rabou, 2010), grapevine cultivar (Morandi Filho et al., 2008; 345 Gonzalez Luna & La Rossa, 2016), climatic conditions (Cocco et al., 2017), nitrogen 346 fertilization (Cocco et al., 2015) and vineyard floor management (Muscas et al., 2017). The 347 main aim of this study was to highlight differences in the reproductive output of P. ficus 348 females mated at increasing ages. Therefore, life table values may vary under different 349 conditions, but nonetheless they provide an insight into the influence of the female age at 350 mating on the reproductive output and indicate the distinct response of P. ficus females to 351 delayed mating. The demographic parameters observed by Gonzalez Luna & La Rossa (2016) 352 on P. ficus reared in two-year-old grapevines were comparable with those observed during 353 our experiment in females mated at 1-3 days, which are the natural mating conditions in the 354 field.

Changes in the reproductive output of *P. ficus* due to delayed mating could affect pheromonemediated pest control strategies, such as mating disruption. Mating disruption against *P. ficus*

357 would be effective if mating is suppressed or delayed for more than 7 days. In fact, delayed 358 mating did not reduce mealybug fecundity but increased the development time of P. ficus 359 populations, thereby reducing the number of generations per year. The effectiveness of mating 360 disruption is influenced by the persistence of the synthetic pheromone in the vineyard. Our 361 ongoing studies investigating the influence of mating disruption on the reproductive traits of 362 P. ficus in the field have indicated a significant reduction of ovipositing females as well as the 363 lengthening of the pre-oviposition period in the pheromone-treated plots throughout the 364 season (A. Cocco, unpublished results). The mating disruption control technique applied to 365 lepidopteran pests is potentially more effective, because a simple mating delay of a few days, 366 even without mating suppression, greatly affects female fecundity, thus reducing the 367 population growth potential (Vickers, 1997; Jones & Aihara-Sasaki, 2001; Walker & Allen, 368 2011). For instance in the koa seedworm, Cryptophlebia illepida (Butler) (Lepidoptera: 369 Tortricidae), a mating delay of 6 days reduced the population growth rate by about 20% 370 (Jones & Aihara-Sasaki, 2001), whereas in P. ficus a similar growth rate reduction was 371 obtained with a mating delay \geq 3 weeks. In vineyards protected by mating disruption, a higher 372 percentage of unmated P. ficus females, a lower percentage of ovipositing females and a lower fecundity were observed, although the contribution of mating delay on the reduction of 373 374 the pest population density was not estimated (Walton et al., 2006; Cocco et al., 2014). In 375 order to better understand the influence of the reproductive traits on the effectiveness of 376 mating disruption to control *P. ficus*, further field studies are ongoing to quantify the length of 377 the mating delay caused by the control method adopted.

378

379 Acknowledgements

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539 Tables

541	Table 1.	Effect	of delayed	mating o	on ovip	osition	behaviour	and	lifetime	(mean	\pm SE)	of
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Female age at mating (d)	Pre-oviposition period (d)	Pre-reproductive period (d)	Oviposition period (d)	Total lifetime (d)
1	$9.18\pm0.39a$	$10.14\pm0.40a$	$14.92 \pm 1.21 a$	$56.96 \pm 1.43a$
3	$7.06\pm0.42b$	$10.56\pm0.42ab$	$15.44 \pm 1.82a$	$56.42 \pm 1.90a$
5	$6.27\pm0.52 bc$	$11.78\pm0.52 bc$	$16.33 \pm 1.40a$	$60.27\pm2.08a$
7	$5.04 \pm 0.42 cd$	$12.34\pm0.43c$	$14.80 \pm 1.24a$	$57.42 \pm 1.27a$
14	$3.52\pm0.19e$	$17.92\pm0.20d$	$13.20\pm0.49a$	$61.38\pm0.77a$
21	$3.38\pm0.11e$	$24.82\pm0.12e$	$13.20\pm0.70a$	$68.60 \pm 1.00 b$
28	$3.66 \pm 0.12 ed$	$32.08\pm0.11f$	$13.04\pm0.50a$	$76.16\pm0.77c$
	$F_{6,160} = 41.95$ P < 0.001	$F_{6,160} = 614.78$ P < 0.001	$F_{6,160} = 1.42$ P = 0.212	$F_{6,160} = 29.83$ P < 0.001

Planococcus ficus females.

544 Means within columns followed by the same letter are not significantly different by Tukey's

545 test (P < 0.05).

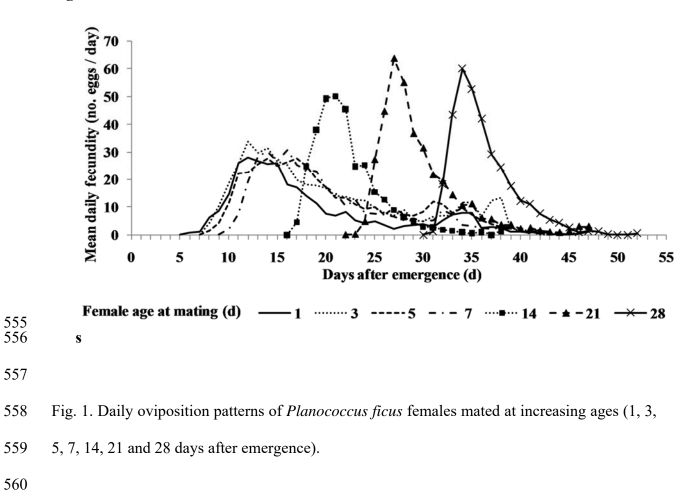
547 Table 2. Life table parameters (mean \pm SE) of *Planococcus ficus* in relation to female age at

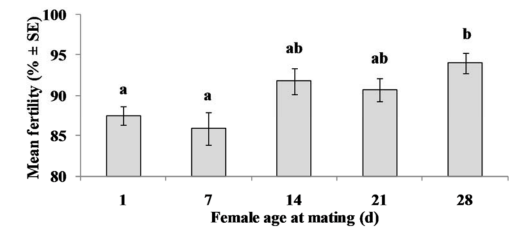
- 548 mating.
- 549

Female age at mating (d)	Net reproductive rate (R_0) (\bigcirc / \bigcirc)	Intrinsic rate of increase (r_m) $(\bigcirc / \bigcirc / d)$	Finite rate of increase (λ) (\bigcirc / \bigcirc / d)	Mean generation time (T _G) (d)	Population doubling time (DT) (d)
1	$140.23\pm0.42a$	$0.114\pm0.001a$	$1.120\pm0.002a$	$43.50\pm0.33a$	$6.10\pm0.06a$
3	$175.44 \pm 1.00 \text{cd}$	$0.115\pm0.002a$	$1.121\pm0.002a$	$44.94\pm0.55b$	$6.03\pm0.11a$
5	$190.45\pm0.66e$	$0.112\pm0.002a$	$1.119\pm0.001a$	$46.93\pm0.74\text{c}$	$6.20\pm0.10\text{a}$
7	$170.77\pm0.58b$	$0.113\pm0.001a$	$1.120\pm0.001\text{a}$	$45.50\pm0.79 \text{bc}$	$6.14\pm0.08a$
14	$172.88\pm0.48bc$	$0.105\pm0.002b$	$1.110\pm0.002b$	$49.19\pm0.31\text{d}$	$6.62\pm0.07b$
21	$193.63\pm0.61f$	$0.091 \pm 0.001 \texttt{c}$	$1.095\pm0.002c$	$57.85 \pm 0.68 e$	$7.62\pm0.04c$
28	$177.11 \pm 0.50 d$	$0.080 \pm 0.002 d$	$1.084 \pm 0.002 d$	$64.43\pm0.52f$	$8.63 \pm 0.06 \text{d}$
	$F_{6,160} = 879.58$ P < 0.001	$F_{6,160} = 600.37$ P < 0.001	$F_{6,160} = 776.01$ P < 0.001	$F_{6,160} = 660.28$ P < 0.001	$F_{6,160} = 30.12$ P < 0.001

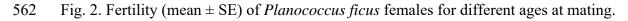
551 Means within columns followed by the same letter are not significantly different by Tukey's

552 test (P < 0.05).









563 Different letters above columns indicate significant differences (Tukey's test, P < 0.05). Note

that the *y*-axis starts at 80%.

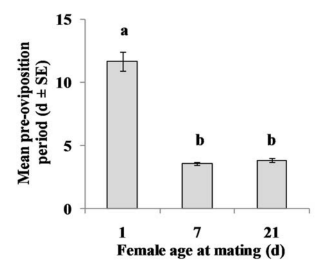


Fig. 3. Length of the pre-oviposition period of *Planococcus ficus* females released in the fieldat different ages.