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Additional Information

1 **Deployment of mating disruption dispensers before and after first seasonal male flights for**
2 **the control of *Aonidiella aurantii* in citrus**

3

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10

11 **Abstract** The rejection of citrus fruit caused by infestations of the California red scale (CRS), *Aonidiella aurantii*
12 (Maskell) (Hemiptera: Diaspididae), raises concerns about its management. This fact has led to the introduction of
13 new integrated control methods in citrus orchards, including the implementation of techniques based on pheromones.
14 Previous works described efficient mating disruption pheromone dispensers to control *A. aurantii* in the
15 Mediterranean region. The main aims of the present study were to adjust the timing of dispenser applications and
16 study the importance of controlling the early first generation of *A. aurantii* by testing two different application dates:
17 before and after the first CRS male flight. The efficacy of the different mating disruption strategies was tested during
18 2010 in an experimental orchard and these results were confirmed during 2011 in a commercial citrus farm. Results
19 showed that every mating disruption strategy achieved significantly lower male captures in monitoring pheromone
20 traps compared with untreated plots, as well as mean fruit infestation reductions of about 80%. The control of the
21 first CRS generation is not essential for achieving a good efficacy as demonstrated in two locations with different
22 pest pressure. The late application of MD dispensers before the second CRS male flight has proven to be effective,
23 suggesting a new advantageous way to apply mating disruption.

24

25 **Keywords** California Red Scale, Hemiptera: Diaspididae, pheromone, semiochemical, mesoporous dispenser,
26 integrated pest management

27

28 **Key Message**

29 Mating disruption (MD) of *Aonidiella aurantii* proved successful in citrus orchards. The importance of controlling
30 the first generation of *A. aurantii* has been investigated by checking the efficacy of MD applied before and after the
31 first generation. We concluded that the control of the first generation is not essential and the application of MD
32 before the second male flight has proven to effectively reduce fruit infestation. This late deployment allows a
33 reduction in the required quantity of pheromone, thus increasing the economic viability of MD in citrus crops.

34

35 **Author Contribution Statement**

36 VN and JP conceived and designed research. SV, CA and VN conducted experiments. SV analyzed data. SV and VN
37 wrote the manuscript. All authors read and approved the manuscript.

38

39 **Introduction**

40 Infestations of California red scale (CRS), *Aonidiella aurantii* (Maskell) (Hemiptera: Diaspididae), pose a serious
41 problem for citrus growers, as CRS may lead to a reduction in tree vigor and the downgrading or commercial
42 rejection of fruits. The economic importance of this armored scale is due to its presence on the surface of the fruits,
43 as a cosmetic damage, and the cost of the strategies needed to control it, even more intense in fresh citrus market.

44 Since CRS control has been affected by the development of resistances to insecticides (Yust et al. 1943; Collins et al.
45 1994; Grafton-Cardwell and Vehrs 1995; Levitin and Cohen 1998), the development and introduction of new
46 integrated and biological control programs was essential for citrus crops. The use of mineral and vegetable oils
47 (University of California 1991; Grafton-Cardwell and Reagan 1995; Rongai et al. 2008) and insect growth regulators
48 (Yarom et al. 1988; Grout and Richards 1991a; Grafton-Cardwell et al. 2006; Eliahu et al. 2007; Rill et al. 2007)
49 appeared as good alternatives to conventional pesticides. These products have provided good control results but they
50 can be harmful to beneficial arthropods (Grafton-Cardwell and Gu 2003; Grafton-Cardwell et al. 2006; Desneux et
51 al. 2007; Vanaclocha et al. 2013). The main enemies of CRS are species of the *Aphytis* parasitoids (Hymenoptera:
52 Aphelinidae) (DeBach 1959; DeBach and Argyriou 1967). Specifically, *Aphytis melinus* (DeBach) is the most
53 successful agent but the control of CRS by augmentative releases of this parasitoid is still under study in Spain
54 (Sorribas et al. 2008; Pekas et al. 2010; Sorribas et al. 2012; Tena et al. 2013). Although in some regions the
55 augmentative biocontrol of CRS through *A. melinus* has achieved good results (Avidov et al. 1970; McLaren and
56 Buchanan 1973; Furness et al. 1983; Moreno and Luck 1992; Bedford 1996), the current cosmetic thresholds in
57 citrus fresh fruit market makes the *A. melinus* success quite improbable in the short term.

58 Integrated pest management programs include the implementation of control methods based on pheromones. Tashiro
59 and Chambers (1967) demonstrated the production of a sex pheromone in CRS, whose chemical structures were
60 reported by Roelofs et al. (1977), as 3-methyl-6-isopropenyl-9-decen-1-yl acetate and (Z)-3-methyl-6-isopropenyl-
61 3,9-decadien-1-yl acetate. Since then, synthetic sex pheromone traps have been widely employed as a detection and
62 monitoring tool for CRS populations (Kennett and Hoffmann 1985; Moreno and Kennett 1985; Grout et al. 1989;
63 Grout and Richards 1991b). The efficacy of mating disruption (MD) technique against CRS was not clearly
64 demonstrated in the first experiments using rubber pheromone dispensers (Barzakay et al. 1986; Hefetz et al. 1988).
65 However, by studying different pheromone doses, Vacas et al. (2009) described a new mesoporous dispenser capable

66 of interfering with normal *A. aurantii* chemical communication. The efficacy of these mesoporous dispensers was
67 further verified, when CRS male catches and fruit infestation were significantly reduced by applying doses of about
68 40 g pheromone/ ha for six months (Vacas et al. 2010). Moreover, the pheromone environment in MD-treated
69 orchards resulted harmless for the performance of *A. melinus* in field and laboratory trials (Vacas et al. 2011;
70 Vanaclocha et al. 2012). By means of all these studies, it was found that CRS mating disruption achieved control at
71 least equal to conventional oil sprays, providing growers with an advantageous control tool. It is commonly known
72 that oil sprays need careful planning and application for a satisfactory control level. On the other hand, the
73 deployment of MD dispensers does not need replacement or qualified hand-labor, and only a minimum flight
74 monitoring is needed to determine the moment of application; all these are important advantages to take into account.
75 However, research had shown the need for additional trials to adjust the timing of dispenser application to cover all
76 the CRS generational cycles. In Spain CRS shows between three and four complete generations with four male
77 flights; the first taking place from mid-April to mid-May, the second from mid-June to mid-July, the third from end-
78 August to September and a late fourth flight during October-beginning of November. Following this dynamic, if all
79 the generations must be affected, mating disruption should be maintained for eight months what would mean a large
80 amount of pheromone, making this control method unaffordable. However, several authors have stated that the first
81 flight of *A. aurantii* males is not correlated with fruit infestation and abundance of the following flights (Moreno and
82 Kennett 1985; Hernández-Penadés et al. 2002; Campos-Rivela et al. 2012). Thus, the importance of controlling the
83 early first generation of *A. aurantii* has been investigated and the efficacy of MD applied before and after the
84 development of the CRS first generation has been examined both in small plots and in a commercial large size
85 orchard.

86

87 **Materials and Methods**

88 Mesoporous dispenser and device

89 The pheromone dispensers applied in the mating disruption (MD) treatments are based on the technology of
90 inorganic molecular sieves (Corma et al. 1999, 2000). Pheromone is impregnated in a natural clay mineral matrix
91 called sepiolite. Its structure, with a high specific surface area, confers to the dispenser good properties for the
92 adsorption and release of organic molecules. Besides the pheromone, formulations include different additives to give

93 consistency and protect the dispenser against humidity. The impregnated material is then compressed in a cylindrical
94 mold by means of a hydraulic press. The manufacturing process has been licensed to Ecología y Protección Agrícola
95 S.L. (Valencia, Spain) who manufactured the dispensers for these trials. Currently, these MD dispensers are
96 commercialized since 2013 by Syngenta Agro (Madrid, Spain) under the name Dardo[®].
97 Dispensers contained 70 mg (a.i.) of the CRS sex pheromone as the diastereomeric mixture (3S,6R and 3S,6S) of the
98 3-methyl-6-isopropenyl-9-decen-1-yl acetate with 75% chemical purity; the remaining 25% belongs to the by-
99 product 3-methyl-6-isopropylidene-9-decen-1-yl acetate, without pheromonal activity. Dispensers were attached to
100 tree branches inside polypropylene baskets 50 mm wide and 90 mm long with a hanger at the top (Ecología y
101 Protección Agrícola SL, Valencia, Spain). Pheromone is released through the 6 × 5 mm grid walls of the basket.

102

103 Experimental design - Trial 2010

104 The field trial was conducted in a 3 ha mandarin (*Citrus reticulata* × *sinensis*; var. Ortanique) experimental orchard
105 located in Denia (Alicante, Spain; UTM: X243500 Y4303900) under Mediterranean climate conditions (mean
106 temperature = 19.9°C, mean relative humidity = 71.2%). Trees were 20 years-old and spaced 6 m by 4 m. The trial
107 was designed with 11 plots: nine plots of 0.3 ha alternately arranged to test three different procedures for the
108 application of mating disruption and two plots of 0.1 ha as reference untreated plots. Pheromone treated plots were at
109 least 50 m apart, whereas untreated plots were separated 60 m from any MD plot. Pheromone dispensers were
110 applied on 29 March 2010 in three plots before the appearance of the first CRS flight (MD-I treatment). Another
111 three plots had dispenser application on 28 May 2010, before the CRS second flight (MD-II treatment). Timings of
112 dispensers' deployment were determined according to general population dynamics in the study area and degree-day
113 accumulation ($DD = [(T_{max} + T_{min})/2] - T_{critical}$; being $T_{critical} = 11.7^{\circ}C$) (Kennet and Hoffmann 1985). Finally, the
114 combination of MD-II application with an oil treatment against nymphs of the first generation was tested in another
115 three plots (MD-II + Oil), where dispensers were also placed on 28 May 2010 and oil treatments were applied on 28
116 May in only these three plots. Pheromone dispensers in every plot were placed at a density of one per tree (420
117 dispensers ha⁻¹) and were not replaced during the experiment.

118

119 Experimental design - Trial 2011

120 A new field trial was conducted in a commercial citrus farm, with larger plots, to confirm the result obtained in the
121 previous trial 2010. MD timings were tested in a 10 ha orange (*Citrus sinensis* Osbeck, var. Navelina) orchard
122 located in Nerva (Huelva, Spain; UTM: X717288 Y4177107) under semi-continental climate conditions (mean
123 temperature = 21.6°C, mean relative humidity = 56.6%). Trees were 25 years-old and spaced 7 m by 3.5 m. In this
124 case, plots with the same MD timing were contiguous as mating disruption is more efficient when applied in large
125 areas. Thus, pheromone dispensers were applied on 7 April 2011 in a whole 1 ha as MD-I treatment, while for MD-II
126 treatment, 1.5 ha had dispenser application on 6 June 2011. Timings were determined according to population
127 dynamics and degree-day accumulation. MD pheromone dispensers in both strategies were placed at a density of 410
128 dispensers ha⁻¹ and were not replaced during the experiment. A third 0.3 ha plot was left without treatments as
129 untreated plot, which was 100 m apart from the pheromone treated areas.

130

131 Oil sprays

132 Given that fruit infestation was over 20% in the previous season, oil treatments in the corresponding plots were timed
133 for the presence of crawlers, which were monitored according to the sampling method suggested by the Valencian
134 regional IPM programme (DOCV 2008). A total of 25 infested branches (2–3 years old) were randomly sampled
135 each week from the date of the first flight and taken to the laboratory. Leaves and twigs were removed from the
136 branches and cut into 10 cm long pieces. Using a binocular scope, a total of 100 live scales were identified as first,
137 second and third instars, adult females, and adult females with crawlers. The oil treatment was applied when first and
138 second instars represented 70% of live scales and more than 90% of adult females had crawlers. Paraffinic oil (15 g l⁻¹
139 ¹) (Argenfrut RV; GulfOil Argentina SA, Argentina) applications were made with an airblast sprayer calibrated to
140 deliver 3500 l ha⁻¹ at 150 psi with the tractor driven at 1.55 km h⁻¹.

141

142 Evaluation of treatment efficacy

143 The efficacy of the different strategies was evaluated according to the CRS male flight inhibition and the fruit
144 infestation assessment. Both parameters were studied in the center of each plot to avoid possible edge effects due to
145 pheromone drift between contiguous treated areas, as buffer areas are considered to be 15 m from the plot borders
146 (Vacas et al. 2009). One commercial white sticky pheromone trap (Pherocon® V Trap; Trécé Inc., Adair, OK) was
147 placed in the center of each plot in Trial 2010 to compare male catches between the different control strategies. In the

148 case of Trial 2011, captures were also evaluated in triplicate, as in the center of three subplots within each above
149 described area. All the monitoring traps were revised every 7 or 15 days, from April to November, and the recorded
150 captures in each trap were divided by the corresponding number of days (7 or 15) to obtain the number of males
151 captured per trap and day (MTD). Pherocon® monitoring lures (Trécé Inc., Adair, OK), loaded with 250 µg sex
152 pheromone, were replaced every 42 days.

153 To measure the inhibition of male catches that occurred in pheromone-treated plots, the mating disruption index
154 (MDI) for each strategy was calculated as an indicator of the treatment efficacy using the following formula: $MDI =$
155 $(1-(x/y)) \times 100$, where x is the number of males captured in MD plots and y is the number of males captured in
156 untreated plots.

157 Fruit were evaluated for scale infestation of Ortanique mandarins on 21 September 2010 and 29 September 2011 in
158 Navelina oranges. Forty fruit per tree (10 fruit per direction: north, east, south and west) were evaluated on trees
159 located on the center of each plot (250-300 fruits/ha) in Trial 2010 and in the center of three subplots within each
160 area in Trial 2011 (250-300 fruits/ha). A fruit was considered to be scale-infested when it had more than three scales
161 on its surface, as suggested by the treatment threshold published in the Valencian regional IPM guidelines (DOCV
162 2008). The percentage of fruit with more than 10 scales was also recorded to perform a sensitivity analysis.

163

164 Field data analysis

165 Generalized linear model (GLM) techniques assuming negative-binomial error variance were employed to compare
166 the number of CRS males captured in the different treated plots. Models were constructed with MTD data as the
167 dependent variable and treatment, time (week of the study period) and their interaction as the explanatory variables.
168 Given that the MD-II application of dispensers was not carried out until 28 May in Trial 2010 and 6 June in Trial
169 2011, we constructed different models for data from the first male flight and data from the rest of flights (second and
170 third). In this way, we evaluate the initial disruption level in MD-I plots during the first flight when MD-II was not
171 yet installed (from April to beginning of June), and also the significance of the male captures inhibition during the
172 second and third flights (from mid-June to October) for both trials.

173 The significance of the explanatory variables was assessed by backward elimination from the model. When
174 significant effects were found, the *glht* function in the multcomp package (Hothorn et al. 2008) was used to perform
175 Tukey HSD tests for post-hoc pairwise comparisons.

176 Likewise, we used GLM techniques assuming negative-binomial error variance to assess scale-infestation differences
177 between the different treatments. For both trials, models were constructed with the percentages of scale-infested fruit
178 in the trees inspected at the end of the trial as the dependent variable and treatment as the explanatory variable.

179 All statistical analyses were conducted with R (R Development Core Team 2012).

180

181 Pheromone release profile

182 The pheromone release profile of the mesoporous dispensers was studied during the trials. Additional dispensers
183 were aged under field conditions in a nearby area, 500 m away from trial orchard, in order to extract their residual
184 pheromone content at different aging times (from 0 to 250 days of field exposure). Three dispensers were taken per
185 aging time, to be extracted at 40°C for 2 h, with magnetic agitation, in 15 ml of dichloromethane/methanol (2/3, v/v)
186 as solvent. After 1 h of extraction, 0.5 ml of the internal standard 1-dodecanol (20 mg ml⁻¹) were added to the
187 extracts. The extracts were centrifuged (3000 rpm for 8 min) and then filtered with syringe filters before
188 chromatographic analysis. Pheromone was then quantified by gas chromatography with flame ionization detector
189 (GC/FID; Clarus® 500 gas chromatograph; PerkinElmer Inc., Wellesley, MA). All injections were made onto a ZB-
190 5ms (30 × 0.25 mm × 0.25 µm) column (Phenomenex Inc., Torrance, CA), held at 160°C for 5 min, and then
191 programmed at 2°C min⁻¹ up to 180°C, where it was held for 1 min, and then programmed at 45°C min⁻¹ up to 250°C.
192 The carrier gas was helium at 1.2 ml min⁻¹. The pheromone amount was estimated by means of a calibration curve
193 which was previously built by preparing standard solutions with the following concentrations: 10.0, 5.0, 2.0, 1.0 and
194 0.4 mg ml⁻¹ of pheromone and 1 mg ml⁻¹ of the internal standard. Calibration curve was described by the equation y
195 = $a + bx$, where y is the FID peak area ratio of the pheromone and the internal standard ($area_{ph}/area_{IS}$) and x is the
196 known amount of pheromone.

197 Simple regression was used to determine the evolution of the residual pheromone load (mg) according to ageing time
198 for the dispenser employed. The quantified residual pheromone contents were employed in a polynomial regression
199 as the dependent variable, to study the significance of the linear and quadratic effect of time (days and days²) on
200 pheromone emission and check whether the pheromone emission was constant during the time under study. In this
201 case, the residual pheromone load decrease at a constant level and the mean emission rate is given by the slope of the
202 fitted linear model. Statgraphics Centurion XVI v16.1 software (StatPoint Technologies Inc., Warrenton, VA) was
203 used for these analyses.

204

205 **Results**

206 Trial 2010

207 Population dynamics of *A. aurantii* in the area of Denia can be observed by the data obtained with traps located in
208 the untreated plots (Fig. 1). The first flight took place during May with a maximum of 6.93 CRS males per trap and
209 day (MTD). The second flight began at the end of June with the maximum number of males captured in mid-July.
210 Male captures of *A. aurantii* increased from the first week of August. They reached a maximum on 31 August and
211 began to decrease slowly up to the beginning of November, when only 0.4 MTD were registered in the untreated
212 plots.

213 Male catches in plots treated with pheromone remained low throughout the entire season, and only slight peaks were
214 registered according with the three described male flights (Fig. 1). The effect of time factor on male catches during
215 the first flight was statistically significant ($F_{1,16} = 5.30$, $P = 0.035$) according to the natural population dynamics. Yet
216 more crucial, the treatment applied significantly affected male captures ($F_{3,16} = 6.67$, $P = 0.004$) as follows. When
217 first flight was taking place (April to beginning of June), MD was already installed on MD-I plots and captures were
218 94.2% inhibited relative to the untreated plots ($P < 0.001$). Meanwhile, MD was not yet established in MD-II
219 strategies but mean initial population levels were lower compared to the untreated plots although not significantly
220 different (MD-II: $P = 0.226$; MD-II + Oil: $P = 0.310$) (Tukey test; adjusted P values with single step method).

221 Considering, catches from the most abundant flights (mid-June to October), time factor resulted statistically
222 significant ($F_{7,75} = 27.89$, $P < 0.001$), as well as the effect of treatment ($F_{3,75} = 84.75$, $P < 0.001$). Compared to the
223 untreated, male catches were significantly lower in the MD-I, MD-II and MD-II + Oil plots ($P < 0.001$). Thus,
224 communication disruption occurred during this period with the three tested mating disruption strategies, resulting in
225 average male flight inhibitions of 81.5% with the MD-I timing, 87.7% in plots with MD-II and 88.9% with the
226 combined strategy MD-II + Oil (without significant differences among them, $P > 0.2$).

227 Regarding fruit damage assessment, 31% of fruit were scale-infested with more than three scales on the surface when
228 no treatment was applied in the orchard (untreated plot in Fig. 2). Results of every MD treatment differed
229 significantly from the absence of treatments for both infestation levels recorded (more than three scales: $F_{3,76} = 20.98$,
230 $P < 0.001$; more than 10 scales: $F_{3,76} = 22.97$, $P < 0.001$). All of the mating disruption deployments achieved a
231 reduction in scale-infested fruit compared to the untreated plot: 81% damage reduction for MD-I ($P = 0.006$); 95%

232 for MD-II; and 96% by the combination MD-II + Oil ($P < 0.001$). Although CRS fruit infestation was significantly
233 reduced by MD-I application, it differed significantly from the MD-II timing ($P = 0.003$) and MD-II + Oil strategy (P
234 < 0.001) (Fig. 2). Fruit infestation observed in MD-I plots did not exceed 6%, whereas less than 1.5% of fruit was
235 found to be scale-infested in the MD-II plots.

236

237 Trial 2011

238 The male population level was significantly higher in Nerva regarding to level of captures obtained in Denia ($F_{1,7} =$
239 9.19 , $P = 0.02$). In the area of Nerva, the first male flight peaked during April and the first weeks of May (Fig. 3)
240 with a maximum of 16.57 MTD in the untreated plot. The first catches belonging to the second flight were obtained
241 on 22 June, while the third flight peaked on 31 August.

242 Male catches in plots treated with pheromone remained low throughout the entire season with statistical differences
243 regarding to the untreated plot (Fig. 3). Time had a significant effect on catches during the entire period of study
244 (first flight: $F_{9,18} = 21.78$, $P < 0.001$; rest of flights: $F_{18,36} = 9.05$, $P < 0.001$), according to natural population
245 dynamics. The effect of treatment factor was significant (first flight: $F_{2,18} = 64.66$, $P < 0.001$; rest of flights: $F_{2,36} =$
246 248.55 , $P < 0.001$). Considering catches from the first flight, MD-I achieved a significant mean male flight reduction
247 ($P < 0.001$) of 97.3% relative to the control. Given that MD-II was not yet installed when the first flight took place,
248 mean male captures during April and May in MD-II plots were significantly higher compared to MD-I ($P < 0.001$).
249 When multiple comparison was performed with data from the most abundant flights (July-October), CRS male
250 captures were 96.2% and 99.1% reduced, relative to control, with MD-I and MD-II respectively (with significant
251 differences between MD strategies, $P < 0.001$).

252 Damage assessment revealed that pheromone treatments had a significant effect on CRS fruit infestation (more than
253 three scales: $F_{2,67} = 27.16$, $P < 0.001$; more than 10 scales: $F_{2,67} = 16.33$, $P < 0.001$). Damage was significantly lower
254 in both pheromone treated plots, regarding to the untreated which had 55% fruit with more than three scales ($P <$
255 0.001). The percentage of fruit with more than three scales was 71.8% reduced with MD-I treatment, while this
256 reduction achieved 82.7% with the MD-II application, although they were not significantly different ($P = 0.145$)
257 (Fig. 4).

258

259 Pheromone release profile

260 Pheromone release profile of mesoporous dispensers is depicted in Fig. 5. The complete model was fitted to an
261 exponential model (solid line in Fig. 5, equation 1) resulting in $R^2 = 0.95$.

$$262 \quad y = 71.322 \times e^{-0.008x} \quad (\text{equation 1})$$

263 However, statistical analysis showed that the curvature of this model was due to data from the last three months of
264 the dispenser life-span. Polynomial regression of data from 0 to 154 days of aging (end March to August), evaluated
265 the significance of the quadratic (days²) and linear (days) effects of time and confirmed the absence of curvature
266 (quadratic effect not significant: $P = 0.31$; linear effect: $P < 0.001$). Thus, the release profile was fitted to the line
267 given by equation 2 (discontinuous line in Fig. 5), resulting $R^2 = 0.99$. This means that emission of mesoporous
268 dispensers is assumed to be constant from 0 to 154 days (until August), and the mean release rate given by the slope
269 of the linear model is $334 \mu\text{g day}^{-1}$. From this date on, emission level decreased below $100 \mu\text{g day}^{-1}$ during the last
270 months of field exposure.

$$271 \quad y = 70.241 - 0.334x \quad (\text{equation 2})$$

272

273 **Discussion**

274 The efficacy of the mating disruption technique against CRS infestations was previously demonstrated by Vacas and
275 coworkers (Vacas et al. 2009; Vacas et al. 2010; Vacas et al. 2011); nevertheless, timing of dispensers' deployment
276 needed to be adjusted for an optimal practical application. In the present work, the efficacy of the late application of
277 MD dispensers, before the second CRS male flight (MD-II), has been demonstrated in two trials carried out in 2010
278 and 2011, in two different locations of Spain, with different climates, male population levels and citrus varieties.
279 CRS is able to develop from three to five generations per year, mainly influenced by temperature (Kennett and
280 Hoffmann 1985; Grout et al. 1989). Under the climatic conditions of Spanish citrus areas, CRS shows three complete
281 generations and a possible fourth generation in some areas and during warmer autumns. Generally, the first male
282 flight takes place between mid-April and mid-May, and is usually not too abundant. In the present work, the
283 importance of controlling the first generation of *A. aurantii* was investigated in 2010 by the application of mating
284 disruption at two different times of deployment: before (MD-I) and after (MD-II) the occurrence of the first male
285 flight. According to CRS male flight monitoring in trial 2010, the first flight was significantly inhibited in MD-I
286 plots. Once MD was also installed in MD-II and MD-II + Oil plots, catches were maintained at low levels throughout
287 the study and both MD timings achieved more than 80% reduction of CRS male catches. This disorientation effect

288 was confirmed in trial 2011 in larger commercial plots with higher male population levels, where both MD strategies
289 achieved mean flight inhibition of about 97%.

290 In the trial carried out in 2010, the scale-infestation assessment revealed that the MD-II results were significantly
291 better than the MD-I strategy, despite reducing damage by around 80%. This could be explained by the release
292 profile and life-span of the pheromone dispensers. According to the results, flight inhibition in MD-I plots was not
293 significantly different from MD-II plots during the third flight but MD-II achieved significantly greater reduction of
294 fruit infestation. Extraction and quantification of the pheromone remaining in the aged dispensers showed that
295 release rate was constant for five months and equal to $334 \mu\text{g day}^{-1}$, and thereafter it decreased. This reduction in
296 release rate is not due to climate factors but to dispenser formulation itself. The release profile of this kind of
297 dispensers is highly temperature independent (Domínguez-Ruiz et al. 2008) and release rate is lower as closer the
298 pheromone load to the residual content that remains retained in the dispenser. If the mesoporous dispensers are
299 applied in March (MD-I), this five month period with proper pheromone emission would protect the crop only until
300 the end of August, without covering the entire third flight of CRS males. Mean pheromone emission during
301 September in MD-I plots was $130 \mu\text{g day}^{-1}$, clearly under the optimum release level of $250 \mu\text{g day}^{-1}$ suggested by
302 Vacas et al. (2009). We think that this lower emission rate could still have a disorientation effect of males towards
303 traps, but could be insufficient to disrupt the short-range attraction and mating of males. It has been described for
304 moth pests that the amount of pheromone needed to disrupt male orientation to traps is lower than the amount needed
305 to disrupt mating (Ioratti et al. 2011). In this way, when dispensers are applied on May (MD-II), pheromone emission
306 is maintained at suitable levels to protect the crop against the entire third flight.

307 The present work confirms that the late deployment of dispensers is at least as efficient as the deployment before the
308 first flight. Therefore, it has been observed with different infestation levels that the control of the CRS first
309 generation is not crucial and damage can be controlled by establishing MD before the second flight with mesoporous
310 dispensers releasing $334 \mu\text{g day}^{-1}$ constantly for at least five months. This could be related with the fact highlighted
311 by several authors who stated that the first flight is not correlated with the abundance of the following flights
312 (Moreno and Kennett 1985; Hernández-Penadés et al. 2002; Campos-Rivela et al. 2012). Thus, the first flight is not a
313 good predictor of infestation later in the season, probably because survival and activity of the first generation is
314 highly affected by more unstable weather conditions. However, mating disruption of the first emerging moths is
315 crucial for the development of the subsequent generations throughout the season in Lepidoptera species. Several

316 authors demonstrated that early pheromone applications prevent mid-season increases in Lepidoptera populations
317 (Staten et al. 1987; Kehat et al. 1995; Lykouressis et al. 2005); these populations being responsible for high yield
318 losses. By contrast, we have observed that the control of the first CRS generation is not so essential for achieving a
319 good efficacy. Moreover, this first generation does not usually colonize the fruit and the second annual crawler
320 generation, which takes place in the summer, is generally considered to be mainly responsible for the infestation of
321 fruit (Rodrigo et al. 2004).

322 It has been demonstrated that the late application of dispensers allows a reduction in the required quantity of
323 pheromone, thus increasing the economic viability of CRS mating disruption. Currently, available dispensers are not
324 able to release pheromone at a suitable level during all the CRS male flights. The development of pheromone
325 dispensers with higher load and longer lifespan would not be cost-effective, as the current cost of pheromone
326 synthesis is a limiting factor for MD implementation. At the moment, pheromone represents about 90% of the value
327 of the dispenser (Ecología y Protección Agrícola, pers. comm.) and this technique is already in the upper limit of the
328 costs affordable by the growers (300 € ha⁻¹ for the MD treatment vs. 200-250 € ha⁻¹ for two oil sprays). Therefore,
329 with the dispensers available in the market, the recommended strategy will be the application of mating disruption
330 before the second CRS male flight.

331

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339 **References**

340 Avidov Z, Balshin M, Gerson U (1970) Studies on *Aphytis coheni*, a parasite of the California red scale, *Aonidiella*
341 *aurantii* in Israel. BioControl. 15:191-207

342 Barzakay I, Hefetz A, Sternlicht M, Peleg BA, Gokkes M, Singer G, Geffen D, Kronenberg S (1986) Further field
343 trials on management of the California red scale, *Aonidiella aurantii*, by mating disruption with its sex-
344 pheromone. *Phytoparasitica* 14:160-161

345 Bedford ECG (1996) Problems which we face in bringing red scale, *Aonidiella aurantii* (Maskell), under biological
346 control in citrus in South Africa. *Proc Int Soc Citriculture* 1:485-492

347 Campos-Rivela JM, Martínez-Ferrer MT, Fibla-Queral JM (2012) Population dynamics and seasonal trend of
348 California red scale (*Aonidiella aurantii* Maskell) in citrus in Northern Spain. *Span J Agric Res* 10:198-208

349 Collins PJ, Lambkin TM, Bodnaruk P (1994) Suspected resistance to methidation in *Aonidiella aurantii* (Maskell)
350 (Homoptera: Diaspididae) from Queensland. *J Aust Entomol Soc* 33:325-326

351 Corma A, Muñoz-Pallares J, Primo-Yufera E (1999) Production of semiochemical emitters having a controlled
352 emission speed which are based on inorganic molecular sieves. World Patent WO9944420

353 Corma A, Muñoz-Pallares J, Primo-Yufera E (2000) Emitter of semiochemical substances supported on a sepiolite,
354 preparation process and applications. World Patent WO0002448

355 DeBach P (1959) New species and strains of *Aphytis* (Hymenoptera: Eulophidae) parasitic on the California red
356 scale, *Aonidiella aurantii* (Mask.), in the Orient. *Ann Entomol Soc Am* 52:354-362

357 DeBach P, Argyriou L (1967) The colonization and success in Greece of some imported *Aphytis* spp. (Hymenoptera:
358 Aphelinidae) parasitic on citrus scale insects (Homoptera: Diaspididae). *BioControl* 12:325-342

359 Desneux N, Decourtye A, Delpuech JM (2007) The sublethal effects of pesticides on beneficial arthropods. *Ann Rev*
360 *Entomol* 52:81-106

361 (DOCV) Diari Oficial de la Comunitat Valenciana (2008) DOCV no. 5901, 26. Resolution 27 October 2008 of
362 Consellería de Agricultura, Pesca y Alimentación; November 2008.
363 http://www.docv.gva.es/datos/2008/11/26/pdf/2008_13692.pdf

364 Domínguez-Ruiz J, Sanchis J, Navarro-Llopis V, Primo J (2008) A new long-life trimedlure dispenser for
365 Mediterranean fruit fly. *J Econ Entomol* 101:1325-1330

366 Eliahu M, Blumberg D, Horowitz AR, Ishaaya I (2007) Effect of pyriproxyfen on developing stages and
367 embryogenesis of California red scale (CRS), *Aonidiella aurantii*. *Pest Manag Sci* 63:743-746

368 Furness G, Buchanan G, George R, Richardson N (1983) A history of the biological and integrated control of red
369 scale, *Aonidiella aurantii* on citrus in the lower Murray Valley of Australia. *BioControl* 28:99-212

370 Grafton-Cardwell EE, Lee JE, Stewart JR, Olsen KD (2006) Role of two insect growth regulators in integrated pest
371 management of citrus scales. *J Econ Entomol* 99:733-744

372 Grafton-Cardwell EE, Gu P (2003) Conserving vedalia beetle, *Rodolia cardinalis* (Mulsant) (Coleoptera :
373 Coccinellidae), in citrus: a continuing challenge as new insecticides gain registration. *J Econ Entomol* 96:1388-
374 1398

375 Grafton-Cardwell EE, Reagan CA (1995) Selective use of insecticides for control of armored scale (Homoptera:
376 Diaspididae) in San-Joaquin Valley California citrus. *J Econ Entomol* 88:1717-1725

377 Grafton-Cardwell EE, Vehrs SLC (1995) Monitoring for organophosphate-resistant and carbamate-resistant armored
378 scale (Homoptera: Diaspididae) in San-Joaquin Valley citrus. *J Econ Entomol* 88:495-504

379 Grout TG, Richards GI (1991a) Effect of buprofezin applications at different phenological times on California red
380 scale (Homoptera: Diaspididae). *J Econ Entomol* 84:1802-1805

381 Grout TG, Richards GI (1991b) Value of pheromone traps for predicting infestations of red scale, *Aonidiella aurantii*
382 (Maskell) (Homoptera: Diaspididae), limited by natural enemy activity and insecticides used to control citrus
383 thrips, *Scirtothrips aurantii* Faure (Thysanoptera: Thripidae). *J Appl Entomol* 111:20-27

384 Grout TG, Du Toit WJ, Hofmeyr JH, Richards GI (1989) California red scale (Homoptera: Diaspididae) phenology
385 on citrus in South Africa. *J Econ Entomol* 82:793-798

386 Hefetz A, Kronengerg S, Peleg BA, Bar-zakay I (1988) Mating disruption of the California red scale *Aonidiella*
387 *aurantii* (Homoptera: Diaspididae). Proc 6th Int Citrus Congress, Tel Aviv (Israel), pp 1121-1127

388 Hernández-Penadés P, Rodríguez-Reina JM, García-Marí F (2002) Umbrales de tratamiento para cóccidos
389 diaspídidos en cítricos. *Bol San Veg Plagas* 28:469-478

390 Hothorn T, Bretz F, Westfall P (2008) Simultaneous Inference in General Parametric Models. *Biometrical J* 50:346-
391 363

392 Ioratti C, Anfora G, Tasin M, De Cristofaro A, Witzgall P, Lucchi A (2011) Chemical ecology and management of
393 *Lobesia botrana* (Lepidoptera: Tortricidae). *J Econ Entomol* 104:1125-1137

394 Kehat M, Anshelevich L, Harel M, Dunkelblum E (1995) Control of the codling moth (*Cydia pomonella*) in apple
395 and pear orchards in Israel by mating disruption. *Phytoparasitica* 23:285-296

396 Kennett CE, Hoffmann RW (1985) Seasonal development of the California red scale (Homoptera: Diaspididae) in
397 San Joaquin Valley citrus based on degree-day accumulation. *J Econ Entomol* 78:73-79

398 Levitin E, Cohen E (1998) The involvement of acetylcholinesterase in resistance of the California red scale shape
399 *Aonidiella aurantii* to organophosphorus pesticides. Entomol Exp Appl 88:115-121

400 Lykouressis D, Perdakis D, Samartzis D, Fantinou A, Toutouzas S (2005) Management of the pink bollworm
401 *Pectinophora gossypiella* (Saunders) (Lepidoptera: Gelechiidae) by mating disruption in cotton fields. Crop Prot
402 24:177-183

403 McLaren IW, Buchanan GA (1973) Parasitism by *Aphytis chrysomphali* Mercet and *A. melinus* Debach of
404 Californian red scale, *Aonidiella aurantii* (Maskell), in relation to seasonal availability of suitable stages of the
405 scale. Austr J Zool 21:111-117

406 Moreno DS, Luck RF (1992) Augmentative releases of *Aphytis melinus* (Hymenoptera: Aphelinidae) to suppress
407 California red scale (Homoptera: Diaspididae) in southern California lemon orchards. J Econ Entomol 85:1112-
408 1119

409 Moreno DS, Kennett CE (1985) Predictive year-end California red scale (Homoptera: Diaspididae) orange fruit
410 infestations based on catches of males in the San-Joaquin Valley. J Econ Entomol 78:1-9

411 Pekas A, Aguilar A, Tena A, García-Marí F (2010) Influence of host size on parasitism by *Aphytis chrysomphali* and
412 *A. melinus* (Hymenoptera: Aphelinidae) in Mediterranean populations of California red scale *Aonidiella aurantii*
413 (Hemiptera: Diaspididae). Biol Control 55:132-140

414 Rill S, Grafton-Cardwell EE, Morse JG (2007) Effects of pyriproxyfen on California red scale (Hemiptera:
415 Diaspididae) development and reproduction. J Econ Entomol 100:1435-1443

416 Rodrigo E, Troncho P, García-Marí F (1996) Parasitoids (Hymenoptera: Aphelinidae) of three scale insects
417 (Homoptera: Diaspididae) in a citrus grove in Valencia, Spain. Entomophaga 41:77-94

418 Roelofs WL, Gieselmann MJ, Cardé AM, Tashiro H, Moreno DS, Henrick CA, Anderson RJ (1977) Sex-pheromone
419 of California red scale, *Aonidiella aurantii*. Nature 26:698-699

420 Rongai D, Cerato C, Lazzeri L, Palmieri S, Patalano G (2008) Vegetable oil formulation as biopesticide to control
421 California red scale (*Aonidiella aurantii* Maskell). J Pest Sci 81: 179-185.

422 Sorribas JJ, Rodríguez R, Rodrigo E, García-Marí F (2008) Niveles de parasitismo y especies de parasitoides del
423 piojo rojo de california *Aonidiella aurantii* (Hemiptera: Diaspididae) en cítricos de la Comunidad Valenciana.
424 Bol San Veg Plagas 34:201-210

425 Sorribas J, van Baaren J, Garcia-Marí F (2012) Effects of climate on the introduction, distribution and biotic
426 potential of parasitoids: Applications to biological control of California red scale. *Biol Control* 62:103-112

427 Staten RT, Flint HM, Weddle RC, Quintero E, Zarate RE, Finell CM, Hernandez M, Yamamoto A (1987) Pink
428 bollworm (Lepidoptera: Gelechiidae): Large-scale field trials with a high-rate gossypure formulation. *J Econ*
429 *Entomol* 80:1267-1271

430 Tashiro H, Chambers DL (1967) Reproduction in the California Red Scale, *Aonidiella aurantii* (Homoptera:
431 Diaspididae). I. Discovery and extraction of a female sex pheromone. *Ann Entomol Soc Am* 60:1166-1170

432 Tena A, Llácer E, Urbaneja A (2013) Biological control of a non-honeydew producer mediated by a distinct
433 hierarchy of honeydew quality. *Biol Control* 67:117-122

434 University of California (1991) Integrated pest management for citrus. University of California, Berkeley, California

435 Vacas S, Alfaro C, Navarro-Llopis V, Primo J (2009) The first account of the mating disruption technique for the
436 control of California red scale *Aonidiella aurantii* Maskell (Homoptera: Diaspididae) using new biodegradable
437 dispensers. *Bull Entomol Res* 99:415-423

438 Vacas S, Alfaro C, Navarro-Llopis V, Primo J (2010) Mating disruption of California red scale, *Aonidiella aurantii*
439 Maskell (Homoptera: Diaspididae), using biodegradable mesoporous pheromone dispensers. *Pest Manag Sci*
440 66:745-751

441 Vacas S, Vanaclocha P, Alfaro C, Primo J, Verdú MJ, Urbaneja A, Navarro-Llopis V (2011) Mating disruption for
442 the control of *Aonidiella aurantii* Maskell (Homoptera: Diaspididae) may contribute to increased effectiveness of
443 natural enemies. *Pest Manag Sci* 68:142-148

444 Vanaclocha P, Vacas S, Alfaro C, Primo J, Verdú MJ, Navarro-Llopis V, Urbaneja A (2012) Life history parameters
445 and scale-cover surface area of *Aonidiella aurantii* are altered in a mating disruption environment: Implications
446 for biological control. *Pest Manag Sci* 68:1092-1097

447 Vanaclocha P, Vidal-Quist C, Oheix S, Montón H, Planes L, Catalán J, Tena A, Verdú MJ, Urbaneja A (2013) Acute
448 toxicity in laboratory tests of fresh and aged residues of pesticides used in citrus on the parasitoid *Aphytis*
449 *melinus*. *J Pest Sci* 86: 329-336

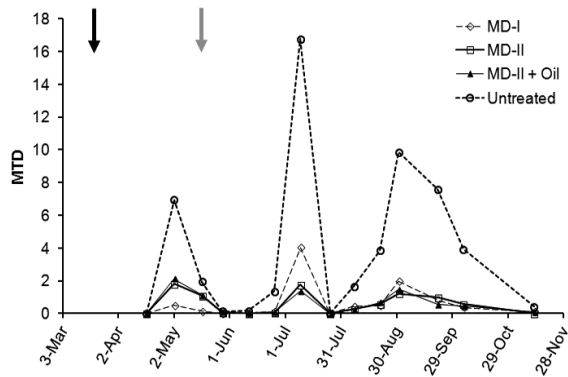
450 Yarom I, Blumberg D, Ishaaya I (1988) Effects of buprofezin on California red scale (Homoptera: Diaspididae) and
451 Mediterranean black scale (Homoptera: Coccidae). *J Econ Entomol* 81:1581-1585

452 Yust HR, Nelson HD, Busbey RL (1943) Comparative susceptibility of two strains of California red scale to HCN,
453 with special reference to the inheritance of resistance. J Econ Entomol 36:744-749

454

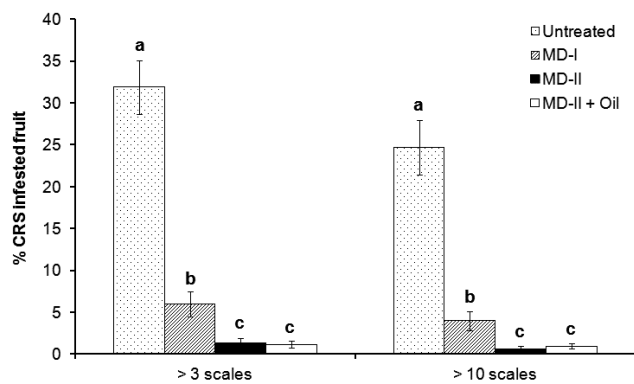
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456 **Fig. 1** Population dynamics of *Aonidiella aurantii* in trial 2010 (Alicante, Spain), shown as males per trap per day
 457 (MTD) captured on the different plots: dispenser application before the first CRS male flight (MD-I), application
 458 before the second flight (MD-II), the combination of MD-II application with an oil treatment, and the untreated plots.
 459 Black arrow indicates dispenser application in MD-I strategy (29 March 2010). The grey arrow points out pheromone
 460 application in MD-II strategy (28 May 2010) and oil application in the May+Oil plots.



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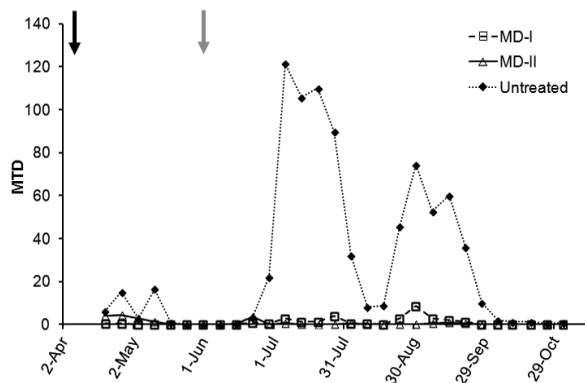
464 **Fig. 2** Mean percentage of scale-infested fruits observed in the different plots of trial 2010: untreated, MD-I, MD-II
 465 and the combination of MD-II with oil spray. Bars labeled with the same letter do not differ significantly (Tukey
 466 HSD tests, $P > 0.05$).



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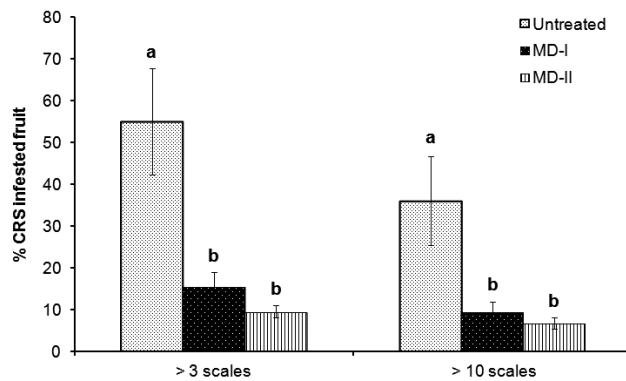
470 **Fig. 3** Population dynamics of *Aonidiella aurantii* in trial 2011 (Huelva, Spain), shown as males per trap per day
 471 (MTD) captured on the different plots: dispenser application before the first CRS male flight (MD-I), application

472 before the second flight (MD-II) and the untreated plot. Black arrow indicates dispenser application in MD-I strategy
 473 (7 April 2011) and the grey arrow points out pheromone application in MD-II strategy (6 June 2011).



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477 **Fig. 4** Mean percentage of scale-infested fruits observed in the different plots of trial 2011: MD-I, MD-II and the
 478 untreated plot. Bars labeled with the same letter do not differ significantly (Tukey HSD tests, $P > 0.05$).



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482 **Fig. 5** Evolution of the remaining load of pheromone (mg) on the mesoporous dispensers versus time (days in
 483 orchard). The complete release profile was fitted to an exponential model (equation 1, $R^2 = 0.95$), although
 484 pheromone release rate was constant until 154 days of field exposure and fitted a linear model (equation 2, $R^2 =$
 485 0.99). The x-axis represents the dates corresponding to ageing time.

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