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#### Population dynamics of Cacopsylla melanoneura (Hemiptera: Psyllidae) in Northeast Italy and its role in the Apple proliferation epidemiology in apple orchards

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26	Population dynamics of Cacopsylla melan	<i>noneura</i> (Hemiptera: Psyllidae) in Northeast
27	Italy and its role in the Apple proliferation	on epidemiology in apple orchards
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#### 35 Abstract

36 In the present study, incidence of '*Candidatus* Phytoplasma mali' in an experimental apple 37 orchard in Northeast Italy, in addition to abundance and phytoplasma infectivity of Cacopsylla 38 melanoneura (Hemiptera: Psyllidae) were determined and the role of this psyllid as a vector of 39 'Ca. P. mali' in this region were reviewed. Insect samples collected in the orchard by the beating 40 method indicated high abundance of C. melanoneura, (up to 7.92 specimens/branch); however 41 the psyllid C. picta was not observed. Molecular analyses revealed presence of 'Ca. P. mali' in 42 6.25% of overwintered psyllids. This infection rate is quite high in comparison to other localities 43 where C. melanoneura is known as the main vector of the phytoplasma. This finding supports 44 the assumption that C. melanoneura also is paramount in the epidemiology of the apple 45 proliferation disease also in Northeast Italy. Moreover, we correlated immigration dynamics to 46 the temperatures registered in the apple orchard, and defined an immigration index to predict the 47 progressive arrival of the overwintered adults from winter sites. Psyllids start to reach the apple 48 orchards when either the average of the maximum temperature of the seven days is above 9.5°C 49 or the immigration index has a positive value. This index will be a useful tool for the growers to 50 prevent apple proliferation phytoplasma spread with well-timed insecticide treatments targeted 51 against C. melanoneura. However, further research is needed to validate or adjust the index to 52 other apple growing regions, which may affect more efficacious management of this disease and 53 psyllid vector.

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56 **KEY WORDS:** *psyllid vector, 'Candidatus Phytoplasma mali', immigration dynamics,* 

57 *temperature, apple orchards, overwintered adults,* 

#### 58 Introduction

Apple proliferation is a serious disease which has caused significant economic losses over the past ten years in all European apple-growing regions (Kunze 1989). Fruit weight is often reduced by 30-60%, the fruit color is unsatisfactory and the taste is poor, with the result that as much as 80% of the fruits is unmarketable.

63 The disease is induced by 'Candidatus Phytoplasma mali', a phytoplasma that colonizes the 64 phloem system of apple plants and is transmitted by insect vectors during their trophic activity. 65 The pathogen is acquired passively during feeding in the phloem of infected plants, moves 66 through the intestine and pass intracellularly through the epithelial cells to enter the hemocoel. 67 Then the phytoplasma circulates in the hemolymph, replicate and finally penetrates specific cells of the salivary glands. With the next feeding, the insect can transmit the phytoplasma to a healthy 68 69 plant. Affected plants lack vigor and fruits are markedly reduced in size with poor flavor and low 70 sugar and acidity content (Osler and Loi 1986). Since the first incidence occurred in the Trentino 71 region (northeast Italy) in the mid-1990s, many studies were carried out concerning the spread of 72 the phytoplasma and the insect vectors. High percentages of symptomatic plants were first 73 observed in several orchards of Trentino (Vindimian and Delaiti 1996, Vindimian et al. 2000) 74 affecting several varieties, mainly Golden Delicious, Florina and Renetta del Canada. Studies on 75 insect vectors revealed the important role of the Hemiptera, genus Cacopsylla in the transmission 76 of this phytoplasma. In Trentino two species, Cacopsylla melanoneura (Förster) and C. picta 77 (Förster), are regularly present in the orchards. Contrary to Northwestern Italy where C. 78 melanoneura is the primary vector of 'Ca. P. mali' (Tedeschi et al. 2002, 2003; Tedeschi and 79 Alma 2004), in Trentino low phytoplasma infection rate in this species and very low 80 transmission efficiency by this psyllid were recorded (Mattedi et al. 2007, 2008). But in a recent

81 report elevated phytoplasma infection rates in C. melanoneura from Trentino region were 82 correlated with the infection level in the plants (Malagnini et al. 2010). On the other hand, C. 83 *picta* showed higher natural infection rate and transmission efficiency even at low density 84 (Frisinghelli et al. 2000, Forno et al. 2002, Mattedi et al. 2007, 2008) in accordance also with 85 German results (Jarausch et al. 2003, 2004, 2008, Mayer et al. 2009). Both these insects have a 86 quite complex life cycle. The overwintered adults reach the apple orchards in winter, in the case 87 of C. melanoneura (Tedeschi et al. 2002; Mattedi et al. 2007) and at the end of March in the case 88 of C. picta (Mattedi et al. 2007). On this host plant they mate, lay eggs and develop. In late 89 spring the newly emerged adults rapidly move to shelter plants, mainly conifers, for aestivation 90 and overwintering.

91 Preliminary evidence suggests that overwintered adults are already infected when they re-92 migrate into apple orchards (Jarausch et al. 2004; Mattedi et al. 2008), so well-timed treatments 93 are very important to control the first individuals that reach the orchards. For this reason a 94 phenology model will be a useful support for vector management decisions. Many forecasting 95 models have been produced for other psyllid species, using means of driving variables, mainly 96 temperature, and based on developmental thresholds (DT) and degree-days (DD) (Beránková and 97 Kocourek 1994, Kapatos and Stratopoulou 1999, Kumral et al. 2008, Morgan and Solomon 98 1993; Schaub et al. 2005). These models refer to some phases of the life cycle and predict when 99 a developmental stage (generally the most damaging stage) will appear. None of these models 100 concerns the immigration period for psyllids that move between different host-plant species or 101 exploit non-host plant species as overwintering sites.

102 In the present study we analyzed in detail the incidence of '*Ca* P. mali' in an experimental

103 orchards in Northeast Italy in addition to the abundance and phytoplasma infection of *C*.

104 *melanoneura*, with the aim to revise the role of this psyllid as a vector of '*Ca* P. mali' in this 105 region. Moreover, we propose an index based on the maximum temperatures registered in the 106 apple orchard, to predict the arrival of the overwintered adults. This index will be a useful 107 management tool for the growers to prevent phytoplasma spread thanks to well-timed insecticide 108 treatments.

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#### Materials and methods

#### 111 Field samplings

112 The study was carried out over five growing seasons (2006-2010) in a conventionally treated 3 113 hectare apple orchard of the IASMA Research and Innovation Center, Fondazione E.Mach 114 (Borgo Valsugana, 419 m a.s.l., Trento, Italy). Observations on the incidence of Apple 115 proliferation disease and the population dynamics of C. melanoneura were performed in 4 116 untreated plots (area 1500 m<sup>2</sup>), which were randomly selected in the orchard. The main apple 117 variety was Golden delicious, with 15-25 years old trees, 4.0±0.5 m high, and spaced 0,5-1.6 m 118 within a row and 4.0-4.6 m between rows. The number of symptomatic trees was recorded at the end of September and a total number of twenty-one samples was randomly collected from 119 120 symptomatic plants to be analyzed with molecular assays. During each season, insect monitoring 121 started before the immigration in the orchard of the first overwintering adults (January/February) 122 and it lasted till the emergence of the springtime generation (May/June). Adults were collected 123 every 7 days, by means of beating method: for each replication plot, 25 branches (50±10 cm in 124 length) were considered and every branch was shaken 2 times above a beating tray (diameter 7 125 cm, 60 x 40 cm of cloth). The collected adults were counted and identified in the laboratory by 126 both morphological, examining female and male terminalia (Ossiannilsson, 1992) and molecular

tools (Tedeschi and Nardi 2010), then analyzed for phytoplasma presence. An in-depth study on
the frequency of '*Ca*. Phytoplasma mali'-positive psyllids was carried out in 2008 and a total
number of 194 batches, each of 5 specimens, was analyzed. The number of eggs and young
stages was assessed by examining under the microscope 30 apical shoots (20-25 cm long) from
each replication, randomly selected in the central rows of each plot. These controls were
performed every 7 days, from the beginning of the oviposition until the emergence of the
springtime generation.

134

#### 135 Phytoplasma detection and species identification

136 Plant DNA was isolated from 100 mg (wet weight) of phloem tissue from symptomatic plants, 137 previously ground with liquid nitrogen in a sterile mortar, using the QIAGEN's DNeasy® Plant 138 Mini Kit (Qiagen, Hilden, Germany). DNA was eluted in 100 µl of elution buffer and kept at -139 20°C until used. Total DNA was extracted from batches of five adult psyllids following a 140 protocol adapted from Marzachi et al. (1998) and previously applied to psyllids (Tedeschi et al. 141 2002). The final product was resuspended in 50 µl of TE. 142 Insect and plant DNAs were amplified firstly with the phytoplasma universal primer pair P1/P7 143 (Schneider et al. 1995) and then in nested PCR with the primers fO1/rO1 (Lorenz et al. 1995) 144 specific for the AP-group phytoplasmas, after a 1:40 dilution. Reaction conditions were as in the 145 original papers. Contamination by amplicons was avoided by using separate rooms and material 146 as well as decontamination procedures (UV exposure and bleaching of materials and surfaces). 147 Moreover, for each amplification, a negative control containing Milli-Q water was included. Amplification products were analyzed by 1% agarose gel electrophoresis, stained with 148 ethidium bromide, and visualized on a U.V. transilluminator. Specific 'Ca. Phytoplasma mali' 149

150 profiles were obtained by RFLP analysis with endonuclease SspI, digesting seven microliters of

the amplification product with 3U of SspI for 4.5 h at 37°C.

152 The proportion of infected insects was estimated by its maximum-likelihood estimator, ^p,

153 calculated according with Swallow (1985):  $^{p} = 1 - H^{1/k}$ , where H is the observed fraction of

154 healthy groups and k is the number of insects per group, five in this case.

155 The psyllid identification by morphological analyses was confirmed by molecular analyses using

156 the primer pairs MEL fw/MEL rev developed for *C. melanoneura* (Tedeschi and Nardi 2010).

157 Thirty-two randomly selected individuals were used to confirm morphological observations.

158 Reaction conditions were as in the original papers.

159

#### 160 **Population dynamics and index of immigration**

161 The population trend of *C. melanoneura* in the apple orchards was based on data collected during

162 the years 2006-2010 for adults and 2006-2008 for eggs and juveniles. In order to identify which

163 environmental parameters in the orchards were possibly involved in *C. melanoneura* 

164 colonization we developed an original model according to the following procedure. Starting,

165 from 2006 the maximum, and median daily and hourly temperatures were recorded at 2 meters

above the ground from a meteorological station inside the apple orchard. For daily temperatures

167 we also calculated the average of the seven days  $(T_{7n})$  preceding any sampling date  $(a_n)$ . Seven

168 days was chosen since field samplings were conducted with such a periodicity. Temperature

169 trends were associated with the insect dynamics, from the date of first detection  $(a_0)$  to the peak

170 of captures  $(\mathbf{a}_{max})$ . In the first instance, the absolute highest temperature among the 7 days

171 preceding  $a_0$  ( $T_{0max}$ ) was taken and then the minimum across the years among the  $T_{0max}$ . This

172 minimum was defined as hypothethical threshold temperature (**Tth**) of psylla orchard

173 immigration. The Tth, calculated both for median and maximal temperatures was checked

174 whether it had been previously passed without eliciting the psyllid immigration and we selected

175 only those that were not passed. The same procedure was done for the  $T_{7n}$  where a relative

176 hypothetic threshold  $(T_7 th)$  was defined.

177 The average of degree days were counted with temperature above the threshold for any  $T_{7n}(dd_n)$ .

178 The index of immigration was defined as:

179

180  $Ii = [(T_{7n} - T_7th) + dd_n]$ 

181 The immigrated population (**Ip**) as:

 $182 (p_n - p_{n-1})$ 

183 where  $(p_n - p_{n-1})$  is the difference of sampled adults (expressed as % referred to the population 184 peak) between two consecutive samplings.

185 A simple regression analysis was performed between Ii and Ip, from the  $a_0$  to the  $a_{max}$  of each

186 year (2006-2010). If the regression was significant then the previsions were verified concerning

187 the detection of any a<sub>0</sub> by calculating **Ii** related to any sampling from 2006 to 2010 *a posteriori*.

188 The first positive value of Ii was assessed whether it was associated to the first psylla detection

189 in the field. The same procedure was followed for historical data taken from the same orchard in

190 the period 2002-2004. Eventually, all data were associated to the apple phenology, in particular

- 191 bud breaking and first flowering.
- 192
- 193

#### **Results**

194 Apple proliferation incidence

195 The symptoms observed in the orchards were predominantly witches' brooms, reddening of the

196 leaves, enlarged stipulae, abnormally long flower stalks, flowering out of season and small fruits.

197 Basing on the percentage of symptomatic trees, the incidence of apple proliferation disease

ranged from 60 to 100%, according to the plot, with a mean annual increase of 5%.

199

#### 200 Psyllid identification and phytoplasma detection

201 Following the morphological observations, almost all the psyllids collected in the experimental

202 plots were identified as C. melanoneura. C. mali was sporadically observed; whereas, C. picta

203 was never found. Molecular analyses confirmed the identifications of *C. melanoneura*.

204

205 The PCR analyses performed with primer pair fO1/rO1 gave AP-group specific amplicons of the 206 expected size (1050 bp) with both plant and insect DNA. No amplification products were 207 obtained from the negative controls. All 21 samples of symptomatic plants were positive to AP-208 group phytoplasmas and the RFLP with SspI restriction enzyme confirmed the presence of 'Ca. 209 Phytoplasma mali'. The results of PCR detection in the insects are shown in table 1. One 210 hundred and seventy-four samples of overwintered adults, 5 samples of nymphs and 15 samples 211 of newly emerged adults were tested. Forty-eight out of 174 batches of overwintered psyllids 212 were positive for 'Ca. Phytoplasma mali', with an estimation of 6.25% of infected C. 213 melanoneura. No amplification was obtained from nymphs and newly emerged adults. 214

### 215 **Population dynamics and index of immigration**

216 The population trend of *C. melanoneura* overwintered adults was extremely variable during the

217 years . The immigrant adults were recorded starting from the beginning of February in 2008,

218	while in 2006 they were delayed to mid-March. The peak of overwintered adults was reported
219	between the 10th and the 13th week, depending of the year, but it was mostly concentrated in the
220	second half of March. Eggs of C. melanoneura were observed on apple branches starting from
221	the beginning of March in 2008 and the last ten days of the month in 2006 and 2007. The
222	oviposition lasted until mid-April in 2007 and the beginning of May in 2006 and 2008. Young
223	stages were recorded from the beginning of April in 2007 and 2008 and the end of the same
224	month in 2006. Nymphs remained in the orchards until the beginning of May in 2007 and the end
225	of May in 2006 and 2008 (Fig. 1).
226	Combining the population dynamics of <i>C. melanoneura</i> with the apple phenology (Table 2), a
227	good correspondence between overwintered adult peak – first egg detection and apple bud
228	breaking was found, with the peak of egg always preceding the first flowering.
229	Values and indexes referred to the psylla migration analysis are reported in table 3. The <b>Tth</b>
230	calculated from maximum (12.8°C) and median (3.3°C) temperatures did not explain the first
231	appearance of the psylla in the orchard, and no correlation was found with the insect migration
232	into the orchard. Similarly, the $T_7$ th taken from median temperatures (2.0°C) did not represent
233	an effective threshold value, because it has been passed several times in the years without
234	triggering the appearance of psylla in the orchard. Instead, the $T_7$ th calculated from maximum
235	values (9.5°C) was found to be a candidate for a possible migration threshold. The regression
236	analysis between <b>Ip</b> and <b>Ii</b> (Fig.2) was highly significant (F = 24.2; df = 1,26; $p < 0.001$ ).
237	Positive values of <b>Ii</b> corresponded with the first appearance of the psylla in the orchard in 2006
238	and 2008-10. Instead, in 2007 values slightly above zero (1.5 and 0.9) in the first five weeks did
239	not correspond to the first presence. However, when the Ii reached for a more relevant value

240 (6.5), individuals were eventually found. Positive **Ii** fitted in verifying all the  $\mathbf{a}_{0}$  also for 241 historical data of 2002-2004.

- 242
- 243

#### Discussion

244 The present research provides important and new information on the epidemiology of apple 245 proliferation in northeast Italy and develops an index of immigration that enables well-timed 246 control measures against the insect vectors. The role of C. melanoneura as vector of 'Ca. 247 Phytoplasma mali' has been revised in this area confirming the results obtained by Malagnini et 248 al. (2010) on psyllid infectivity. Previous reports concerning Trentino region pointed out the 249 predominance of C. picta in Val d'Adige and C. melanoneura in Val di Sole and Val di Non (up 250 to 2 overwintered adults/branch) (Mattedi et al. 2007, 2008). Transmission trials and 251 phytoplasma quantification in insect bodies revealed a low transmission efficiency for C. 252 melanoneura while an important increase in phytoplasma concentration was observed only in C. 253 *picta*, indicating an efficient phytoplasma multiplication in this species (Pedrazzoli et al. 2007). 254 For this reason it is now popularly held belief that C. picta is the most important vector of 'Ca. 255 Phytoplasma mali' in northeast Italy, as well as in Germany (Jarausch et al. 2003, 2004, 2008; 256 Mayer et al. 2009), while in northwestern Italy, C. melanoneura represents the only psyllid 257 vector of this phytoplasma (due also to the absence of *C. picta*). 258 Evidence supports the role of C. melanoneura as a vector in Valsugana, a valley in the south east 259 of the Trentino region with a long tradition in apple production. This area is characterized by an high incidence of the apple proliferation disease and the only important psyllid species which 260 261 was found in the repeated collections trough a five year period was C. melanoneura; C. picta was 262 never recorded. The molecular analyses revealed the presence of 'Ca. Phytoplasma mali' (as the

Swallow's estimated proportion p<sup>^</sup>) in 6.25% of overwintered psyllids, confirming roughly the results obtained by Malagnini et al. (2010) with specimens coming from the same area. This discrepancy may be due to the different number of psyllids analyzed. During the peak presence there were up to 7.92 overwintered adults/branch in 2007, and from the beginning until the end of March in all the three years the number of recorded *C. melanoneura* was almost always higher than 2 psyllids/branch. Thus, in this valley, *C. melanoneura* should be considered the only significant vector associated with the rapid spread of this disease.

The infection rate of the overwintered adults tested high in comparison to other localities where *C. melanoneura* is already known as the main vector of '*Ca.* phytoplasma mali' (an estimation of 6.25% vs 3.6% of '*Ca.* Phytoplasma mali'-positive insects) (Tedeschi et al. 2003). This fact strengthens the evidence that *C. melanoneura* has an important role in the epidemiology of the apple proliferation disease also in northeast Italy, where its control is already regulated by insecticide treatments (Baldessari et al. 2007, 2009).

276 Once ascertained that C. melanoneura represents an important risk for apple growers, a new 277 approach to the management of this vector needs to be developed. Data collected during the 278 years concerning the population dynamics of this species, the apple phenology and the weather 279 parameters allowed us to define an index to predict the re-migration of overwintered adults into 280 apple orchards. The aim was to provide to the growers an easy to use tool which would predict 281 the time of arrival of C. meanoneura.in order to refine the pest management decisions. The 282 efficiency of an insecticide treatment depends on the product and on the timing of application in 283 order to affect the most sensitive or harmful stage on the crop. In the case of C. melanoneura, 284 overwintered adults showed, in comparison with all the other stages, the highest population density, the highest percentage of 'Ca. Phytoplasma mali'-positive specimens, and the longest 285

286 time spent in apple orchards. All these characteristics suggest that this is the crucial role of this 287 stage in vectoring AP-phytoplasma (Tedeschi et al. 2002, 2003). For this reason the treatments 288 should be focused on the overwintered adults as soon as they start to colonize apple orchards. 289 As a consequence, we established a index of immigration related to the immigrant psylla adults. 290 Similar studies were built in the past to forecast the occurrence of a particular life stage of a pest 291 with the aim to improve its control (Beránková and Kocourek 1994, Kapatos and Stratopoulou 292 1999, Kumral et al. 2008, Morgan and Solomon 1993, Schaub et al. 2005), but never to predict 293 the adult immigration in the orchards. Here we propose an empirical model which cannot be 294 supported by laboratory trials due to the impossibility of reproducing the entire biological cycle 295 of C. melanoneura in controlled conditions. This correlation is based on the temperatures 296 recorded in the apple orchards, a variable that can be easily monitored by the growers or at least 297 by the phytosanitary services.

It is known that temperature influences apple phenology and that the migration could also be influenced by certain phenological stages of the trees. Immigration into orchards take place during budding, which may be detectable by the overwintering psylla, thereby attracting them to the stimuli associated to budding.

However, the distances occurring between psyllid host plants and shelter plants can be
considerable (Čermák and Lauterer 2008, Thebaud et al. 2009), thus it is improbable and
temperature is most likely thethe critical factor. There was no correlation between immigration
dynamics and apple phenology demonstrated, however oviposition occurs at bud burst while egg
peak and hatchings are always before the first flowering. This confirmed a good degree of
synchrony between *C. melanoneura* and host-plant growth, being linked with temperatures as

stated for psyllids in general by Hodkinson (2009). These data are likewise useful for thegrowers for possible further treatments during the season.

310 In the present study, the hypothetic threshold calculated as an average of the maximum 311 temperatures of the seven days  $(T_7 th)$  proved to be a good tool to forecast immigration, by 312 calculating a proper Index of immigration (Ii). Psyllids started to reach the apple orchards when 313 the  $T_7$ th was 9.5°C. Also including in the model single episodes, meant as hours above this 314 threshold, we could individuate with extremely precision the time of first occurrence and the 315 following immigration trend for a period of 9 years. It is feasible that any Ii over 0 can elicit 316 psyllid migration, with direct correlation between **Ii** value and number of migrant specimens, and 317 our failure in detecting individuals in correspondence of Ii < 2 depended likely on the extremely 318 low number occurring in the field in those circumstances.

319 On the other hand, there are still some relevant limits that we need to point out. The calculated 320 threshold fits well for apple orchards located in the Valsugana valley, but not necessarily for 321 other locations, where the correlation needs to be validated by adjusting the threshold value, 322 either according to historical collection data or by programming periodical field collections. 323 Other geographical factors associated with the winter sites location (e.g. the regional orography, 324 the main air streams and distance from apple orchards) may differently affect the psylla 325 migration process and influence its presence/absence, both in terms of time and quantity, in a 326 given apple orchard.

A straight correlation between maximum temperature and adult mobility was found in addition to a, temperature threshold that, independently on the period in the winter, favors the psylla to abandon overwintering sites. However, more research is required to set up a proper forecasting model, which could be applied in different apple-growing regions.

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# 423 Table 1 - Estimated proportion (p<sup>^</sup>) of '*Candidatus* Phytoplasma mali'-infected *Cacopsylla*

# *melanoneura* (Förster) collected by beat tray samplings in 2008 in the experimental plots

	Date	AP+/tot	$\mathbf{p}^{\wedge}$		AP+/tot	$p^{\wedge}$
	1 February	0/3	0			
	8 February	5/10	12.94			
	15 February	6/8	24.2			
	22 February	1/8	2.63			
overwintered	28 February	5/30	3.58		48/174	6.25
overwintered	7 March	11/30	8.73			
	13 March	8/30	6.01			
	21 March	5/30	3.58			
	28 March	6/20	6.88			
	4 April	1/5	4.36	)		
nymphs	15 May	0/15	0	}	0/15	0
offspring	16 April	0/1	0	7	0/5	0
onspring	15 May	0/4	0	ſ		

Table 2 - Number of weeks when an event of psylla or apple phenology was recorded for
each year of investigation (2006-2010). For the historical data (2002-2004) only the first
detection and the peak of captures of psylla were available.

	2002	2003	2004	2006	2007	2008	2009	2010
First Psylla	5	8	6	11	7	4	8	9
Psylla Peak	10	10	11	13	11	10	12	12
First Egg	-	-	-	13	11	10	12	12
Egg Peak	-	-	-	14	13	14	14	13
Bud break	-	-	-	13	10	11	12	12
Flowering	-	-	-	16	14	14	15	15

434	Table 3 - Absolute (T) and average of 7 days (T7) median (med) and maximal (max)
435	temperature (°C) associated with the first psylla detection (suffix $a_0$ ) and with periods
436	before the first detection (suffix a $_{-n}$ ) in the years 2002-2004 and 2006-2010 . Ii is the
437	Immigration index. In bold are indicated the minimal $a_0$ and maximal a $_{-n}$ values across the
438	years for each parameter.

	2002	2003	2004	2006	2007	2008	2009	2010
$Ta_{0med}$	-	-	-	4.9	7.5	12.9	3.3	5.4
Ta-nmed	-	-	-	4.4	8.5	4.2	4.5	4.4
$Ta_{0max}$	-	-	-	12.8	16.4	20.3	12.5	13.0
Ta <sub>-nmax</sub>	-	-	-	11.8	21.8	11.4	9.6	9.8
$T7a_{0 med}$	-	-	-	3.5	5.1	5.3	2,0	4.5
T7a-n med	-	-	-	2.3	3.7	3.1	3.7	3.0
$T7a_{0 max}$	9.8	10.2	10.2	10.1	12.1	11.2	9.5	10.4
T7a-nmax	5.8	6.6	7.7	8.8	9.4	7.0	8.0	7.7

- 442 Table 4. Calculated Index of immigration (Ii) and Immigrated population (Ip) from the
- 443 first week of January until the population peak, in the periods 2002-2004 and 2006-2010.
- 444 Samplings (s) were conducted weekly. In gray cells, in bold, are highlighted values of first
- 445 positive Ii and first Psylla detection in the orchard.
- 446
- 447

	2002		2002 2003		20	2004 200		06 2007			2008		2009		2010	
S	Ii	Ip	Ii	Ip	Ii	Ір	Ii	Ip	Ii	Ip	Ii	Ip	Ii	Ip	Ii	Ip
1	-8.0	0	-7.0	0	-7.5	0	-7.0	0	-3,2	0	-7.9	0	-9.5	0	-6.7	0
2	-6.4	0	-9.0	0	-5.2	0	-8.3	0	-0,4	0	-4.8	0	-6.3	0	-6.0	0
3	-6.1	0	-7.0	0	-6.0	0	-8.2	0	1.5	0	-2.1	0	-5.6	0	-6.3	0
4	-3.6	0	-2.9	0	-8.2	0	-7.0	0	-3.9	0	6.1	0.03	-3.8	0	-7.5	0
5	2.2	0.04	-4.9	0	-1.2	0	-1.7	0	0.9	0	0.0	0.05	-0.9	0	-6.6	0
6	4.3	0.09	-5.0	0	2.3	0.07	-5.1	0	-1.2	0	1.4	-0.02	-2.6	0	-5.3	0
7	-1.3	-0.09	-1.7	0	2.3	0.11	-2.8	0	6.5	0.01	-0.2	0.00	-2.6	0	-3.3	0
8	4.0	0.27	3.4	0.05	-5.7	-0.04	-3.4	0	7.8	0.01	9.3	0.50	1.5	0.04	-1.5	0
9	5.1	0.08	6.9	0.50	-0.3	-0.07	-2.5	0	15.3	0.50	12.2	0.26	2.3	0.15	2.8	0.01
10	14.6	0.61	11.3	0.45	3.2	0.07	-0.1	0	12.2	0.13	4.1	0.19	6.9	0.22	-1.6	0.02
11					6.1	0.85	3.2	0.41	9.2	0.35			11.4	0.60	6.9	0.17
12							7.8	0.50							15.2	0.81
13							10.8	0.10								

## 450 Figure Captions

- 451 Fig. 1. Presence of eggs and nymphal stages of *Cacopsylla melanoneura* (Förster) in the orchard
- 452 in 2006-2008.
- 453 Fig. 2. Simple Regression analysis (F = 24.2; df = 1,26; p < 0.001) between Immigration Index
- 454 (Ii) and Immigrated population (Ip) calculated from the average of maximal temperature of the 7
- 455 days preceding any adult sampling of the period 2002-2010.
- 456
- 457
- 458

Eggs	2006 2007 2008															
Nymphal stages	2006 2007 2008															
Week		9	10	11	12	13	14	15	16	17	18	19	20	21	22	23

464 Fig. 1



469 Fig. 2