



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Tracking the dispersion of Scaphoideus titanus Ball (Hemiptera: Cicadellidae) from wild to cultivated grapevine: use of a novel mark-capture technique

This is the author's manuscript

Original Citation:

Availability:

This version is available http://hdl.handle.net/2318/151386 since 2016-10-12T15:01:13Z

Terms of use:

Open Access

Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)



UNIVERSITÀ DEGLI STUDI DI TORINO

This is an author version of the contribution published on:

LESSIO F., TOTA F., ALMA A. Tracking the dispersion of Scaphoideus titanus Ball (Hemiptera: Cicadellidae) from wild to cultivated grapevine: use of a novel mark-capture technique BULLETIN OF ENTOMOLOGICAL RESEARCH (2014) 104 DOI: 1017/S0007485314000030

The definitive version is available at: http://journals.cambridge.org/action/displayAbstract?fromPage=online&aid;= 9286098&full

Bulletin of Entomological Research

Tracking the dispersion of Scaphoideus titanus Ball (Hemiptera: Cicadellidae) from wild to cultivated grapevine: use of a novel mark-capture technique --Manuscript Draft--

Manuscript Number: BER-D-13-00136R1 Full Title: Tracking the dispersion of Scaphoideus titanus Ball (Hemiptera: Cicadellidae) from wild to cultivated grapevine: use of a novel mark-capture technique Article Type: Full research paper Corresponding Author: Alberto Alma University of Turin Grugliasco, TO ITALY Corresponding Author Secondary Information: Corresponding Author's Institution: University of Turin Corresponding Author's Secondary Institution: First Author: Federico Lessio First Author Secondary Information: Order of Authors: Federico Lessio Federica Tota Alberto Alma Order of Authors Secondary Information: Abstract: The dispersion of Scaphoideus titanus Ball adults was studied applying a water solution of cow milk (marker: casein) or chicken egg whites (marker: albumin) onto the canopy of wild grapevine at a distance from vineyards ranging from 5 to 330 m. Yellow sticky traps were placed on the canopy of grapes, and captured insects were analyzed via an indirect ELISA for markers' identification. Data were subject to exponential regression as a function of distance from wild grapevine, and to spatial interpolation (Inverse Distance Weighted and Kernel interpolation with barriers) using ArcGIS Desktop 10.1 software. The influence of rainfall and time elapsed after marking on markers' effectiveness, and the different dispersion of males and females were studied with regression analyses. Of a total of 5417 insects analyzed, 43% were positive to egg; whereas 18% of 536 tested resulted marked with milk. No influence of rainfall or time elapsed was observed for egg, whereas milk was affected by the time elapsed. Males and females showed no difference in dispersal. Marked adults decreased exponentially along with distance from wild grapevine and up to 80% of them were captured within 30 m. However, there was evidence of long-range dispersal up to 330 m. The interpolation maps showed a clear clustering of marked S. titanus close to the treated wild grapevine, and the pathways to the vineyards did not always seem to go along straight lines but mainly along ecological corridors. S. titanus adults are therefore capable of dispersing from wild to cultivated grapevine, and this may affect pest management strategies.

Tracking the movement dispersion of Scaphoideus titanus Ball (Hemiptera:
 Cicadellidae) from wild to cultivated grapevine: use of a novel mark capture technique
 4

5 Federico Lessio, Federica Tota, Alberto Alma

6

7

Abstract

8 The movement dispersion of Scaphoideus titanus Ball adults from wild to cultivated 9 grapevine was studied with a novel mark capture technique applying. aA water solution of cow milk (marker: casein) or chicken egg whites (marker: albumin) was applied directly onto 10 11 the canopy of wild grapevine more or less in close proximity (5 350 m) to at a distance from 12 vineyards ranging from 5 to 330 m.; Y+ellow sticky traps were placed on the canopy of 13 grapes, and captured S. titanus adultsinsects were analyzed via an indirect ELISA for 14 markers' identification. Data were subject to exponential regression as a function of distance 15 from wild grapevine, and to spatial interpolation analyses (Inverse Distance Weighted and Kernel interpolation with barriers) were performed using ArcGIS Desktop 10.1 software.; Tthe 16 17 influence of rainfall and time elapsed after marking on markers' effectiveness, and the 18 different dispersal patternsdispersion of males and females were also studied with regression 19 analyses. Of a total of 5417 insects analyzed for egg, 43% were positive to egg; whereas 18% 20 of 536 tested were milk-resulted marked with milkpositive. No influence of rainfall or time 21 since the marker's applicationelapsed was observed for egg-marked specimens, whereas milk-22 marked were-was affected by the time elapsed. Males and females showed no difference in 23 dispersal. Marked adults decreased exponentially along with distance from wild grapevine 24 and up to 80% of them were captured within 30 m.; Hhowever, there was evidence of long-25 range dispersal up to 350-330 m. The interpolation maps showed a clear clustering of marked

Formatted: Font: Not Italic Formatted: Font: Not Italic S. titanus close to the treated wild grapevine, and the pathways to the vineyards did not always seem to go along straight lines but mainly along ecological corridors. S. titanus adults are therefore capable of moving-dispersing from wild to cultivated grapevine, and these new findingsthis must be considered when deciding onmay affect pest management strategies.

30

31 Key words: leafhopper vector, dispersal, immunomarking, ELISA, spatial interpolation

- 32
- 33

Introduction

34 The nearctic leafhopper Scaphoideus titanus Ball (Hemiptera: Cicadellidae) was introduced 35 into Europe in the late 1950s (Bonfils & Schvester, 1960) and is now widespread in many 36 European countries from Portugal to Bulgaria (COST Action FA0807). This species is a 37 grapevine specialist, and develops on both wild and cultivated grapevine (Vitis spp.). It is 38 univoltine and overwinters in the egg stage, which is laid under the bark of wood 2-yrs of age 39 or more (Vidano, 1964); eggs start to hatch in the middle of May and nymphs (which include 40 five instars) are present until the end of July, whereas adults usually appear at the beginning 41 of July and are observed up to the middle of October (Vidano, 1964). S. titanus is an 42 important pest, as it is the main vector of grapevine's Flavescence dorée (FD), a disease 43 caused by 16SrV phytoplasmas (subgroups C and D) (Malembic-Maher et al., 2011). Nymphs from the 3rd instar on acquire phytoplasmas by feeding on infected plants (acquisition access 44 45 period, AAP), and following a latency access period (LAP) of 4-5 weeks they become adults and able to transmit FD to healthy plants (IAP) (Bressan et al., 2005). Since FD is a cause of 46 47 great economic losses, insecticidal sprays against S. titanus are mandatory in Italy: active ingredients include neonichotinoids, organophosphates, etofenprox, and natural pyrethrum, 48 49 the latter in organic farming (Lessio et al., 2011a). However, there are still many ecosystems 50 suitable to S. titanus' survival such as untreated vineyards, organic farming vineyards, cast-51 away vineyards, and woods or uncultivated areas colonized by wild grapevine (mainly from

Formatted: Superscript

overgrown rootstocks: *Vitis rupestris, V. riparia × berlandieri*, etc.).: <u>T</u>the easiest way to
assess the threat of these areas to viticulture by serving as reservoirs for this leafhopper vector
is to apply mark-release-recapture (MRR) or mark-capture (MC) techniques.

Marking methods used in entomology include fluorescent dusts (Garcia-Salazar & Landis, 55 1997; Takken et al., 1998; Skovgard, 2002), radioisotopes (Hagler & Jackson, 2001), and 56 57 immunomarking (Hagler & Jackson, 2001; Jones et al., 2006; Hagler & Jones, 2010). In 58 mark-release-recapture (MRR) experiments, insects (obtained under laboratory conditions or 59 captured in the field) are marked, released at a certain point in the field, and then recaptured, 60 usually by means of traps. However, there are many drawbacks in applying MRR methods, both generally and especially concerning S. titanus. First of all, it isn't possible to mark and 61 62 release a quantity of insects as large as the effective population in the field, Mmoreover, the 63 number of marked individuals recaptured is generally small, up to 8-10% (Zhou et al., 2003; Lessio et al., 2008). In addition, the marker may affect the insects' flight behaviour to some 64 65 extent, and it is sometimes difficult to obtain a large quantity of insects, especially with 66 species like S. titanus that have just one generation per year and an obligatory diapause and 67 therefore cannot are difficult be to reared continuously under lab conditions. The possibility of applying a marker directly on the host plants overcomes these problems, and it is possible 68 since the development of ELISA mark detection techniques. The first immunomarking 69 70 method available was based on vertebrate proteins, such as chicken or rabbit immunoglobulin 71 G (IgG) (Hagler, 1997; Blackmer et al., 2004, 2006), but it hasn't been much used because it 72 is too expensive. The development of low-cost markers, such as food proteins like cow milk, 73 soy milk, or chicken egg whites, widened the possibility of using mark-capture techniques in 74 entomology on large-scale experiments (Jones et al., 2006). A recent study compared the 75 performances of so-called first (IgGs) and second (food proteins) generation markers, and 76 found that egg whites have a longer persistence than IgGs, whereas no difference was 77 observed in the insects' mortality (Slosky et al., 2012). For these reasons (the need to mark

78	field-born insect populations, low cost and high reliability of the markers), we decided to
79	apply this novel large-scale mark-capture technique to track the movements of S. titanus
80	adults from wild to cultivated grapevine in Northwestern Italy. As markers, we used cow milk
81	and chicken egg whites (see materials and methods for details).
82	
83	Materials and methods
84	Large scale field marking and sampling of S. titanus
85	Field studies were conducted during 2010- <u>and</u> 2011 in the district of Portacomaro (AT),
86	Piedmont, Italy-(44.97029-44.94596 °N, 8.24774-8.26120 °E). We set up four experimental
87	sites, called A, B, C and D; each site consisted of one or two vineyards (A-1 and A-2 for site
88	A, etc.) more or less in close proximity which disted from 5 to 330 m from to woods colonized
89	by wild grapevine (WGV). All the vineyards were subject to insecticidal sprays: vineyard B
90	received two sprays with Etofenprox on the 26 June and 25 July, whereas all others were
91	sprayed with Thiamethoxam and Chlorpirifos-methyl on the first and second date,
92	<u>respectively.</u> I in the middle of June, before the first spray, we assessed the presence of S .
93	titanus nymphs by visual inspection according to a sequential sampling plan with a fixed-
94	precision level of 75%, based on Green's equation (Lessio & Alma, 2006) (Table 1).
95	As markers we used albumin (pasteurized chicken egg whites: Eurovo SRL, S. Maria in
96	Fabiano Lugo, RA, Italy, approximate cost 5.00 €/lt.), and casein (sterilized Ultra High
97	Temperature, UHT cow whole fat milk: by Centrale del latte di Torino, Italy, approximate
98	cost 0.50 ϵ /lt.), henceforth referred to as egg and milk, which have a greater reliability
99	compared to soy milk (Jones et al., 2006). The markers were used as tap water solutions at a
100	ratio (volume/volume) of 10 and 20% for egg and milk, respectively; Nowe didn't use any
101	water softener and/or wetting agent <u>was used</u> , as they don't significantly improve insect
102	marking in the field (Boina et al., 2009). The markers were applied every 10–20 days from 8 th
103	July to 10 th September (Table 1) using a hand jet sprayer with a 15 l tank, at an approxa- rate

of 40<u>00</u> l/100 mha², directly onto WGV. When two separate WGV stands were present in the
same site, we applied a different marker on each of them; otherwise, we applied only egg,
which is more detectable than milk (Jones *et al.*, 2006). The daily amount of rainfall (mm)
was recorded from a meteorological station nearbyset at the same distance (2 km) from each
of the experimental sites.

109 Yellow sticky traps (cm 20 \times 30) were placed in the vineyards at a distance of $15-20 \pm 2$ m 110 from each other on the vine row, and $5-6 \pm 0.5$ m between rows, depending on plot size (for 111 larger plots, we increased the distances in order to cover evenly the whole plot size), and 112 directly on stands of WGV, at a distance of $15-20 \pm 2$ m from each other (Table 1; Figs. 3-6) 113 to capture marked S. titanus adults; each trap was geo-referenced with a Garmin® GPS 114 receiver and the distance between traps was confirmed by measuring with a graduated tape. 115 Eight to 19 days after each marker's application, captured adults were carefully removed from 116 the traps directly in the field using a wooden toothpick (using a new one every time to prevent 117 cross-contamination), placed into sterilized 1.5 ml microcentrifuge tubes (one insect/tube), 118 and stored at -20° C before analyses. The traps were placed at the beginning of July and 119 replaced after each insect removal up to the middle of October, which represents the window 120 of S. titanus adults' presence in North-western Italy (Lessio & Alma, 2004b).

- 121
- 122

Laboratory analyses

An indirect ELISA was performed to detect protein markers acquired by the leafhoppers; when egg and milk were used in the same sampling site, insects were analyzed so as to detect both markers at once. Commercially available antibodies for chicken egg albumin (RAE, (rabbit anti egg) (C6534, Sigma-Aldrich, St. Louis, MO, USA) and bovine casein (SAC, Sheep anti casein) (antibodies-online GmbH, Aachen, Germany) were used. The secondary antibodies used for the chicken egg albumin and bovine casein assays were peroxidase conjugated donkey anti-rabbit IgG (H + L) (DAR) (31458; Pierce Biotechnology, Rockford, IL, USA) and peroxidase conjugated rabbit anti-sheep IgG (H + L) (RAS) (31480; Pierce
Biotechnology, Rockford, IL, USA), respectively.

132 Reagents included: TBS-EDTA (Tris Buffered Saline, pH 8.0 plus 0.3 g/l sodium 133 ethylenediamine tetra acetate) (Sigma-Aldrich, St. Louis, MO, USA); PBS-BS (Phosphate 134 Buffered Saline + 20% Bovine Serum) (Sigma-Aldrich, St. Louis, MO, USA); PBSS-BS 20 (Phosphate Buffered Saline + 20% Bovine Serum + 1300 ppm Silweet L-77) (Silwet, 135 136 Chemtura Manufacturing, Manchester, UK)); PBSS-BS 30 (Phosphate Buffered Saline + 30% 137 Bovine Serum + 1300 ppm Silweet L-77); PBST (Phosphate Buffered Saline + 0.09% Triton 138 X-100) (Triton-X-100; Sigma-Aldrich, St. Louis, MO, USA), PBS-SDS (Phosphate Buffered 139 Saline + 2.3 g/l Sodium dodecyl sulfate), sulphuric acid (H_2SO_4) 2N; and immuno-pure ultra 140 TMB substrate (Pierce Biotechnology, Rockford, IL, USA).

141 For the chicken egg assay, the primary antibody was diluted 1:4000 (2 μ l in 8.0 ml) in PBSS-

142 BS20, while the secondary antibody was diluted 1:6000 (1.4 μl in 8.4 ml) in PBSS-BS20.

143 For the casein assay, the primary antibody was diluted 1:500 (16 μ l in 8.0 ml) in PBSS-BS30, 144 while the secondary antibody was diluted 1:1500 (5.4 µl in 8.1 ml) in PBSS-BS20. The 145 following protocol, slightly modified after Jones et al. (2006), was applied: 1 ml TBS-EDTA 146 was added to the 1.5 ml microcentrifuge tube with the insect, vortexed for 2-4 seconds and 147 left in stand-by mode for 3 minutes. From each tube, three 80 µl aliquots (replicates) were 148 retrieved and placed in individual wells of a 96-well microplate (Nunc Polysorp, Nalge Nunc, 149 Naperville, IL, USA) (to minimize contamination during washings, the 6 wells closest to the 150 negative and blank controls were left empty); the micro-plate was then covered with 151 aluminium foil and incubated at 37°C for 2 hrs. (at the end of this step, the leafhoppers were 152 sexed by observing the external genitalia with a stereomicroscope and then discarded). The 153 plate was then emptied and washed 5 times with 300 µl PBST using a LT-3000 micro-plate 154 washer (Labtech International Ltd, Uckfield, UK);-). Tthen 300 µl PBSS-BS (for egg) or 300 155 µl PBS-BS (for milk) were added, and the plate was incubated at 37°C for 1 hr. Afterwards, it 156 was washed 2 times with 300 µl PBST, and 80 µl of the first antibody (RAE for egg, SAC for 157 milk) were added and the plate was incubated at 37°C for 30 min. The plate was then 158 emptied, washed 5 times with 300 µl PBST, 80 µl of the second antibody (DAR for egg, RAS 159 for milk) was added, and the plate was incubated at 37°C for 2 hrs. After incubation, the plate 160 was washed 3 times with 300 µl PBS-SDS and 3 times with 300 µl PBST. Then 80 µl TMB 161 were added and the plate was incubated at room temperature $(25^{\circ}C)$ in the dark on a shaker 162 for 10 min. The reaction was then stopped by adding 80 μ l of 2N H₂SO₄ and the plate was 163 scanned with a LT-4000 micro-plate reader (Labtech International Ltd, Uckfield, UK) at 164 wavelengths of λ =450 nm and 492 nm (reference standard).

As positive standards, we used adults of Euscelidius variegatus (Kirschbaum) (Hemiptera: 165 166 Cicadellidae) reared on oat (Avena sativa L.) under laboratory conditions. Potted plants of 167 either oat or broad bean (Vicia faba L.) were sprayed with the markers using a hand vaporizer, 168 and then placed into insect-proof cages (cm $20 \times 20 \times 40$) made of mesh and Plexiglas in a 169 climatic chamber (T=23 ± 2 °C, RH=60%, L:D=16:8 h). In each cage (placed in the climatic 170 chamber) we put some-90 E. variegatus adults; 7 days later, the leafhoppers were removed, 171 killed by freezing, and preserved at -20° C before analyses; some untreated leafhoppers were 172 used as negative controls, and extraction buffer alone was the blank control.

173 Each sample (=insect) was associated with 3 values of optical density (ODS) for each 174 wavelength. The mean ODS at 450 was subtracted from the mean at 492: $ODS_{(450-)}$ 175 492)=ODS450-ODS492; and the same equation was applied to the optical densities of the 176 negative control: ODN₍₄₅₀₋₄₉₂₎=ODN₄₅₀-ODN₄₉₂; and blank: ODB₍₄₅₀₋₄₉₂₎=ODB₄₅₀-ODB₄₉₂. 177 Finally, we obtained the corrected (blanked) optical density for each sample as: 178 $ODCS=(ODS_{450-492})-(ODB_{450-492})$, and of the negative control as $ODCN=(ODN_{450-492})-(ODB_{450-492})-$ 179 (ODB₄₅₀₋₄₉₂). A sample was considered marked when the ODCS was greater than the mean 180 ODCN added plus 4 times its standard deviation (SD): ODCS>ODCN+4SD, providing 181 additional protection against false positives (Jones et al., 2006).

Data analyses

184	The movement-dispersion of S. titanus adults from WGV to the vineyards was studied by
185	fitting an exponential model: $N(r) = a \exp(-br)$, where N is the percentage of marked
186	individuals caught at the minimum distance r from the treated area (5 \pm 1.5 m step), weighted
187	by the number of traps displayed at the same distance r (being P_i the number of positive
188	specimens captured on the total number of traps t_i placed at the t_i^{th} minimum distance r from
189	treated WGV, we have the grand total $\underline{T=\sum P_i(t_i)}$; and subsequently, we calculated $\underline{N=P_i/T}$ as
190	the percentage of marked individuals per trap at the i^{th} distance r); a is a scaling parameter
191	that estimates the number of S. <i>titanus</i> collected at $r = 0$; and b is the spatial scale parameter
192	that models the rate of variation in insects captured. The choice of an exponential model was
193	made to verify if marked S. titanus would decrease at increasing distances from the source
194	(treated WGV) following an exponential decay pattern. For the same reason, for each
195	regression, we calculated the median dispersal index $r_{0.5}$ (that is, the distance where 50% of
196	the marked individuals are found) using the negative half-life equation: $r_{0.5} = \ln(2)/b$
197	(Northfield <i>et al.</i> , 2009).
198	In order to assess differences in dispersal between genders, regression equations were
199	obtained separately for females and males and the homogeneity of the regression test was
200	evaluated (Sokal & Rohlf, 1995). The influence of rainfall occurred and time elapsed between
201	since the marker's application and insect sampling (independent variables) on the percentage
202	of positive individuals captured on traps placed within the treated points (dependent variable)
203	was studied by applying a weighted least square (WLS) linear regression, using the total
204	number of insects captured as the weight variable (Sokal & Rohlf, 1995). All regression
205	analyses were carried out with the SPSS 20.0® statistical package (<u>http://www.spss.it</u>).
206	percentage <u>All percentage</u> data were previously arcsin square root transformed.

206	percentage <u>All percentage</u> data were previously arcsin square root the

Formatted: Font: Italic
Formatted: Font: (Tipo di carattere testo asiati, Subscript
Formatted: Font: Italic
Formatted: Font: Italic
Formatted: Font: (Tipo di carattere testo asiati, Subscript
Formatted: Font: Italic
Formatted: Font: (Tipo di carattere testo asiati, Italic, Superscript
Formatted: Font: Italic
Formatted: Font: Italic
Formatted: Font: Italic
Formatted: Font: (Tipo di carattere testo asiati, Subscript
Formatted: Font: Italic
Formatted: Font: (Tipo di carattere testo asiati, Subscript
Formatted: Font: Italic
Formatted: Font: (Tipo di carattere testo asiati, Italic, Subscript
Formatted: Font: Italic

207	To individuate the pathways of S. titanus adults from WGV to vineyards, spatial interpolation
208	of the marked insects captured was performed applying Inverse Distance Weighting (IDW)
209	and Kernel interpolation with barrier (KB), both available in the ArcMap toolbox of ArcGIS
210	Desktop 10.1 (http://esri.com). The choice of these two models rather than others was made in
211	order to detect a movement pattern of S. titanus based solely on line of sight distances
212	between sampling points (IDW), to another one that might be influenced by the presence of
213	breaklines (KB). The IDW is a deterministic method, based on the Euclidean distance
214	between sampling points (Bartier & Keller, 1996). It is easy and rapid to use, and is
215	appropriate for aggregated data, as it highlights the hot spots (Tillman et al., 2009). The
216	generic IDW equation is: $z_{x,y} = \sum z_i w_i / \sum w_i$, where $z_{x,y}$ is the value to be estimated, z_i is the
217	control value for the i^{th} sample point, and $w_i = (d_{x,y,i})^{-\beta}$ is the weight that states the
218	contribution of each z_i in determining $z_{x,y}$, where <i>d</i> is the distance between sampling points $z_{x,y}$
219	and z_i , and β is defined by the user (the larger the value of β , the smaller the reciprocal
220	influence of the sampling points; in this research we chose $\beta=2$, which is the most widely
221	used). Kernel interpolation is used to determine the "utilization distribution" (UD) of a
222	resource by an animal (Sheather & Jones, 1991; Benhamou & Cornélis, 2010). The kernel
223	<u>Kernel</u> density estimate f_h^h of an univariate density f based on a random sample $X_l,, X_n$ of
224	size <i>n</i> is: $f_h^{(x)} = n^{-1} \sum h^{-1} K [h^{-1} (x-X_i)]$, where <i>K</i> is the kernel function and <i>h</i> is the
225	bandwidth, a smoothing parameter (Sheather & Jones, 1991). Kernel interpolation with
226	barriers (KB) is a variant that uses a non Euclidean distance rather than a line of sight
227	approach, so that the shortest distance between two points within the defined search
228	neighbourhood is used to connect them; in this case, we used as Kernel function the
229	exponential equation, which was used during the regression analysis (whereas no transfer
230	function is needed to apply the IDW method)as kernel function, whereas the bandwidth was
231	calculated as a default by ArcMap. Barriers were crops or natural vegetation stands between
232	treated WGV and vineyards; however, they were considered partially open, as some

Formatted: Font: Italic

233	movement within non-grapevine ecosystems may occasionally occur. The obtained	
234	interpolation maps were tested for accuracy via cross-validation: we calculated the mean	
235	prediction error: $ME = [\sum_{j=l,n} (x_i - x_i)/n]$, and the root mean square error: $RMSE = sqrt[\sum_{j=l,n} (x_i - x_i)/n]$	
236	$(x_i^{-} - x_i)^2/n$], where x_i^{-} is the predicted value, x_i the observed value, and n the sample size. Both	
237	<u>ME and <u>RMSE</u> are given in the same units of measure of the data: an ideal model should have</u>	<
238	a <u>ME equal 0, and a RSME as small as possible.</u> While <u>RMSE gives an estimate of the error as</u>	
239	a whole, ME mainly provides an estimate of the bias: that is, positive and negative ME values	
240	indicate that the model over or underestimates the data, respectively. (Rhodes et al., 2011).	
241		
242	Results	
243	In total, 1675 and 3901 S. titanus adults were captured in 2010 and 2011, respectively;	
244	flight peak occurred between the first ten days of August and the beginning of September. We	
245	analyzed 4881 insects by detecting egg alone (1664 in 2010 and 3217 in 2011), and screened	
246	536 for both egg and milk (all in 2011). The total net percentages Without considering	
247	differences in sites and position of traps, of egg-positive individuals were 32 and 55% in 2010	
248	and 2011, respectively (mean 43%). In 2010, the rate of egg-marked adults captured on WGV	
249	and in vineyards ranged from 36 to 44% and 9 to 68%, respectively (Fig. 1A);). Hhowever,	
250	the minimum value of 9% refers to vineyard C-2, placed at a minimum distance of 220 m	
251	from the treated WGV, where few insects were captured. In vineyard B (minimum distance	
252	from WGV: Dmin.=6 m), although many insects were captured, there were few marked	
253	specimens (<4025%) probably because of a high residential population of S. titanus; in fact,	
254	pest management in this site was different from (and probably less effective with respect to)	
255	the others-(Table 1). In 2011, we found 46–78% and 38–68% of egg-marked adults in WGV	
256	and vineyards, respectively (Fig. 1B). Milk was only used in site D in 2011 on one stand of	
257	WGV (Dmin.=110 m), whereas a second stand (Dmin.=120 m) was sprayed with egg: 97	
258	(18%) of the 536 tested leafhoppers were milk-positive, and 82 of them were captured on	

Formatted: Font: Italic
Formatted: Font: Italic

milk-sprayed WGV; 206 (38%) were egg-positive, and 131 were captured on egg-treated 259 260 WGV (Fig. 1B); finally, 58 (11%) of them were positive for both egg and milk at the same 261 time. The optical density values of positive specimens calculated on 5 plates chosen at 262 random (mean \pm s.e.) were 0.67 \pm 0.09 for egg, and 0.56 \pm 0.19 for milk; positive reference 263 standards (E. variegatus maintained on treated broadbean or oat) scored 2.26 ± 0.03 for milk 264 and 2.28 ± 0.06 for egg, whereas negative controls (untreated *E. variegatus*) were 0.01 ± 0.00 . Rainfall occurred eight times both in 2010 (min. 1.4 mm, max. 35 mm, total amount 125 265 266 mm), and 2011 (min. 0.4 mm, max. 31 mm, total amount 67 mm). No influence of either 267 rainfall or time between applications was observed on the rate of egg-marked S. titanus; on 268 the other hand, milk-marked specimens were negatively related to time (Table 2).

269 The sex ratio (M/F) was generally female biased, both for total (0.39-0.55) and marked 270 (0.35–0.99) individuals; site C in 2010 represents an exception; it was investigated only from 271 the first week of August on, and the sex ratio was 0.08 for both total and marked insects. Egg-272 marked specimens ranged from 33 to 66% for males, and 18-54% for females; whereas milk-273 marked males and females were 17% and 19% of the total captured, respectively. The 274 homogeneity of regression test between the distribution of marked males and females as a 275 function of distance of capture from the treated point was never significant within different 276 experimental sites and years (Table 3). Therefore, the exponential models were fitted to the 277 experimental data (and the subsequent median dispersal indexes calculated) without taking 278 gender into account.

Exponential regression analyses provided a good-significant fit of marked *S. titanus* adults as a function of the minimum distance from the treated point, although in site D we obtained low R^2 values; the subsequent median dispersal indexes ranged from 14 to 70 m within the different experimental plots (Table 4). The cumulative distribution functions show how the main captures (80%) occurred within 20–30 m from WGV (Fig. 2A, B:); however, there was also evidence of long-range dispersal up to $\frac{350-320}{20}$ m (Fig. 2C, D). In site A, captures

285 decreased asymptotically after 25-30 m, although a slight increase was observed between 65 286 and 70 m (Fig. 2A), whereas in site B (investigated only during 2010) they were almost 287 constant with increasing distance (Fig. 2B). In site C, in 2010 there was a clear point break 288 (increase) at a distance of 30 m, and thereafter captures didn't increase anymore; but this site 289 was only observed from the beginning of August in 2010. In the second vineyard (C-2), 290 further from the treated zone, only a single marked specimen was captured. In 2011, the trend 291 was smoother with a constant decrease in captures up to 60 m (maximum distance of the first 292 vineyard, C-1, from WGV); up to 10% of the total marked insects were found in the second 293 vineyard (C-2) (Fig. 2C). In site D, 70% of the egg-marked adults were captured on treated 294 WGV and a cumulative 30% in the vineyard, at a 120-160 m distance, without any clear 295 break point; on the other hand, only 60% of the milk-marked specimens were captured at the 296 treated point, and 40% were found in the vineyard at a distance of 100-220 m (Fig. 2D).

297 On the whole, both IDW and KB interpolation methods showed a clear clustering of marked 298 adults on the edges of the experimental vineyards. In many cases, when WGV was distributed 299 along two edges, the clustering was much more evident if the European grapevine's rows 300 were parallel rather than perpendicular to the edge, e.g. sites A (Fig. 3), and C, concerning the 301 first vineyard (C-1) close to WGV (Fig. 5). Site B, only studied in 2010, shows almost the 302 same pattern (Fig. 4); however, these results should be considered carefully because of the 303 small size of the vineyard. In site D, egg and milk-marked individuals showed almost the 304 same pattern independent of the interpolation method used (Fig. 6), suggesting how an 305 ecological corridor may exist between the two areas colonized by WGV. On the other hand, 306 in site C long distance dispersal from the WGV to vineyard C-2 had a different pattern 307 depending upon the interpolation method used: IDW produced a more uniform map, whereas 308 KB showed how the possible ecological corridors are displaced along the rows (Fig. 5). On 309 the whole, the cross-validation results showed lower ME and RMSE values for KB rather 310 than for the IDW (with the exception of sites B and D, concerning egg-marked specimens),

311 indicating a better interpolation power of the first model compared to the second-interpolation 312 method; the only exception was represented by egg-marked specimens in site D. The ME was 313 generally positive for KB (overestimation) and negative (underestimation) for IDW, however 314 KB always had a lower absolute value (the only exception was represented by egg-marked 315 specimens in site D) (Table 5). Insects marked with both egg and milk were too few in 316 number to perform cross-validation.

- 317
- 318

Discussion

319 The marking method proposed, used in large-scale application on S. titanus, was quite reliable 320 with egg, as up to 78% of the insects captured on the traps placed into the treated wild 321 grapevine (WGV) were marked; on the other hand, milk had a poorer performance (22%). 322 These data are in accord with Jones et al. (2006), who obtained roughly 70% and 23% of 323 marked Cydia pomonella L. in apple orchards treated with egg and milk, respectively; 324 whereas Boina et al. (2009) obtained higher rates of Diaphorina citri Kuwayama marked 325 with egg (88%) and milk (80%). In our research, one of the main problems was to properly 326 treat the WGV canopy, as it develops up to 6 m above ground level in certain places and is 327 sometimes very dense and difficult to reach. In order to study the movement of S. titanus 328 during the entire period of the adults' presence in the field, we applied the markers constantly 329 but sometimes with a longer window of time between application and the insects' removal 330 from traps; otherwise, it would become too time-consuming. We found a higher rate of 331 positive individuals in 2011, probably because of a smaller amount of rainfall; <u>H</u>however, 332 concerning egg, there was no influence of rainfall or time after the marker's application on the 333 rates of positive individuals; Oon the other hand, the time between application and removal 334 did affect the rate of milk-marked S. titanus. In other researches, the rate of marked 335 individuals decreased along with time after application and the amount of (simulated) rainfall 336 (Jones et al., 2006; Boina et al., 2009). Under laboratory conditions, a residue egg-treatment

Formatted: Font color: Auto

13

337 on true bugs remained 68-100% positive up to 10 days after marking, and 27-88% positive 338 from 11 to 20 days after marking (Hagler & Jones, 2010). In addition, direct egg treatment of 339 Hippodamia convergens Guérin-Méleville allowed detection of egg proteins on 100% of the 340 individuals up to 26 days after marking (Sloski et al., 2012). The problem with marking plants 341 is that insects must come into contact with the marker before it dries up or is washed off. In 342 addition, direct marking of S. titanus adults would not be reliable because of the difficulty in 343 obtaining a very large number of specimens, and we couldn't release this leafhopper in the 344 vineyards as it is subject to compulsory pest management. However, our data set (30-50% of 345 egg-marked specimens out of more than 5000 captured) seemed large enough to analyse and 346 interpret the movement patterns of this leafhopper vector.

347 S. titanus adults are therefore capable of both short and long range dispersal from wild 348 (WGV) to cultivated grapevine. This behaviour was previously theorized both in Italy (Pavan 349 et al., 2012), and in the US (Beanland et al., 2006) by comparing captures in traps placed at 350 different distances from potential S. titanus sources: the results of our mark-capture 351 experiments clearly demonstrate how these movements actually occur. The majority of 352 individuals seem to cover short distances: when WGV is close to the edge of the vineyards, 353 up to 80% of the marked individuals are captured within 30 m. However, long distance flight is also possible: S. titanus captures on the local scale are spatially related up to 200 m, 354 355 whereas at greater distances they seem to depend on local factors, mainly pest management 356 strategies (Lessio et al., 2011b). The results of this research confirm this aspect, as some 357 movement occurred up to more than 200 m. In vineyard B, although many insects were 358 captured, there were few marked specimens (<25%) probably because of a high residential 359 population of S. titanus; in fact, pest management in this site was different from (and probably 360 less effective with respect to) the others. Concerning site D, in the vineyard, the majority of 361 marked adults was captured in the North-West corner, suggesting how the infestation may 362 have mainly occurred from the second uncultivated area, treated with milk; however, this area

363 may also have recruited adults from other areas, as suggested by the double-marked 364 individuals, and milk-marked adults being captured in the egg-treated zone and vice versa. On 365 the whole, the Kernel with barriers (KB) interpolation method showed smaller errors (RMSE 366 and absolute ME values) compared to inverse distance weighting (IDW): the first model, 367 which derives partially from the exponential regression (used as a transfer function in the 368 Kernel interpolation process) is therefore more accurate than the latter (due to lower RMSE values), and its overestimation of observed data (ME>0) has a lower absolute value than the 369 underestimation given by IDW (ME<0)., These differences suggesting how the movement 370 371 patterns of S. titanus adults may not depend solely upon their distance from sources but also 372 upon ecological corridors or natural barriers. It seems therefore that this leafhopper is less 373 likely to perform direct long-distance flights, whereas it rather moves along more roundabout 374 pathways. S. titanus adults have a crepuscular flight activity, which makes them not rely on 375 the wind for dispersal (Lessio & Alma, 2004b), and this may be in accord with an active 376 wandering movement rather than a passive wind-borne transport. Moreover, marked adults 377 were generally clustered along the same row of cultivated grapevine rather than on different 378 rows; this is in accord with the fact that they move mainly along the same row, and captures 379 on the same row are more spatially related (Lessio et al., 2009b). Males and females showed 380 no differences in dispersal from wild to cultivated grapes. Usually, males of S. titanus start to 381 fly earlier than females, however, in the late part of the season the presence and flight activity 382 of females is increased, whereas males tend to decrease (Lessio et al., 2009a). This long-range 383 dispersion of females may have a consequence during the next year, resulting in a higher 384 population of S. titanus in vineyards because of egg-laying. 385

Because WGV may also host 16SrV phytoplasmas (Lessio *et al.*, 2007), incoming *S. titanus* adults may also be capable of transmitting FD to cultivated grapevine: in fact, symptomatic grapes are often clustered at the edges, consistent with *S. titanus* coming in from outside Formatted: Font: Italic

389 (Pavan et al., 2012). Within this frame, pest management strategies against S. titanus in NW 390 Italy should be revisited, as the main problem seems to be represented by adults entering the 391 vineyards in the late part of the season; at present, PM focuses on a first spray against nymphs 392 at the end of June, a second one against adults at the middle-end of July, and a further one 393 sometime after harvest (Lessio et al., 2011a). It is perhaps necessary to change this calendar, 394 using a more persistent active ingredient in the late part of the season to protect grapes from 395 inoculation; for instance, neonicotinoids are much more efficient than organophosphates in 396 preventing transmission (Saracco et al., 2008).

397 Other strategies should be directed toward avoidance: the first action to be applied should be 398 to erase WGV as a source of S. titanus; however, such an action must not be done when adults 399 (both males and females) are present, as it may cause an increase of their movement onto 400 European grapevine. The same problem occurs when dealing with Hyalesthes obsoletus 401 Signoret, the vector of Stolbur phytoplasmas causing Bois Noir (Weber & Maixner, 1998), 402 which lives on weeds and only occasionally feeds on grapes as an adult (Alma et al., 1987): if 403 weeds are erased, adults are compelled to move onto grapevine; for example, in Israel, where 404 H. obsoletus has two generations per year, the second generation is more likely to move to 405 grapes if its host plant is harvested or dries up because of summer heat (Orestein et al., 2003). 406 Another means of preventing leafhoppers from entering the vineyard may be the use of insect-407 proof fences (nets). These devices were successfully used in Israel against some Diptera 408 (Vernon & MacKenzie, 1998; Päts & Vernon, 1999; Bomford et al., 2000). A five metres 409 high screen barrier was successfully evaluated in Californian citrus orchards and nurseries 410 against Homalodisca vitripennis (=coagulata) (Say), a vector for Xilella fastidiosa causing 411 Pierce's disease (Blua et al., 2005). Such a protective device against S. titanus should be at 412 least 2.5 m, as high as the flight boundary layer of this leafhopper (Lessio & Alma, 2004a). 413 Moreover, the screen should be provided with an overhang to avoid insects double crossing it 414 by walking on it (Bomford et al., 2000). On the other hand, plantation of trees had

415 inconsistent effects in limiting invasion into vineyards by *Graphocephala atropunctata*416 (Signoret), another vector for *X. fastidiosa* (Daugherty *et al.*, 2012).

In conclusion, the presence of wild grapevines in vine growing areas must be addressed with an integrated pest management strategy that includes: area-wide sprays and use of suitable active ingredients to prevent such transmission as much as possible; avoidance of new vine plantations in regions with a high presence of WGV; destruction of WGV whenever possible, which would decrease the pathways available to this leafhopper; and the development of new tools such as physical barriers to avoid the entrance of *S. titanus* adults into vineyards from outside.

- 424
- 425

Acknowledgments

We are grateful to Edoardo Sala and Francesca Martina for the help given in field collections and laboratory analyses. Meteorological data were kindly provided by "Regione Piemonte Direzione Agricoltura, Settore Fitosanitario - Sezione Agrometeorologica". This work was realized within the frame of the "FLADO" research project, supported by "Regione Piemonte, Servizi di Sviluppo Agricolo".

431

432

References

Alma, A, Arnò, C., Arzone A. & Vidano, C. (1987) New biological reports on
Auchenorrhyncha in vineyards. pp. 509-516 in *proceedings of the sixth Auchenorrhyncha meeting, Turin, 7-11 September 1987* University of Turin, Italy.

Bartier, P.M. & Keller, C.P. (1996) Multivariate interpolation to incorporate thematic
surface data using inverse distance weighting (IDW). *Computers & Geosciences* 22 (7),
795-799.

439	Beanland, L., Noble, R. & Wolf, T.K. (2006) Spatial and temporal distribution of North
440	American grapevine yellows disease and of potential vectors of the causal
441	phytoplasmas in Virginia. Journal of economic-Economic Entomology 35(2), 332-344.
442	Benhamou, S. & Cornélis, D. (2010) Incorporating movement behavior and barriers to
443	improve kernel home range space use estimates. Journal of wildlife Wildlife
444	management Management 74(6), 1353-1360.
445	Blackmer, J.L., Hagler, J.R., Simmons, G.S. & Cañas, LA. (2004) Comparative dispersal
446	of Homalodisca coagulata and Homalodisca liturata (Homoptera: Cicadellidae).
447	Environmental Entomology 33, 88-99.
448	Blackmer, J.L., Hagler, J.R., Simmons, G.S. & Henneberry, T.J. (2006) Dispersal of
449	Homalodisca vitripennis (Homoptera: Cicadellidae) from a point release site in citrus.
450	Environmental Entomology 35, 1617-1625.
451	
452	Blua, M.J., Campbell, K., Morgan, D.J.W. & Redak, R.A. (2005) Impact of a screen
453	barrier on dispersion behaviour of Homalodisca coagulata (Hemiptera: Cicadellidae).
454	Journal of economic <u>Economic</u> Entomology 98 (5), 1664-1668.
455	Boina, D.R., Meyer, W.L., Onagbola, E.O. & Stelinski, L.L. (2009) Quantifying dispersal
456	of Diaphorina citri (Hemiptera: Psyllidae) by immunomarking and potential impact of
457	unmanaged groves on commercial citrus management. Environmental Entomology
458	38 (4), 1250-1258.
459	Bomford, M.K, Vernon, R.S. & Päts, P. (2000) Importance of collection overhangs on the
460	efficacy of exclusion fences for managing cabbage flies (Diptera: Anthomyidae).
461	Environmental Entomology 29(4), 795-799.
462	Bonfils J. & Schvester, D. (1960) Les Cicadelles (Homoptera Auchenorrhyncha) dans leur
463	rapports avec la vigne dans le Sud-Ouest de la France. Annales Epiphyties 11, 325-336.

Formatted: Font: Bold

464	Bressan A., Spiazzi S., Girolami V. & Boudon-Padieu, E. (2005) Acquisition efficiency of	Formatted: Font: Bold
165	Elevencence derée absterleans by Seanheideus titaus Dell from infected telerent er	Formatted: Font: Bold, English (U.K.)
405	Flavescence doree phytoplasma by Scapholaeus manus Ball from infected tolerant or	Formatted: English (U.K.)
466	susceptible grapevine cultivars or experimental host plants. Vitis 44, 143-146.	Formatted: Font: Italic
467	COST Action FA0807. Integrated management of phytoplasma epidemics in different crop	Formatted: Font: Bold
468	systems: phytoplasma diseases and vectors in Europe and surroundings.	
469	http://www.costphytoplasma.eu/WG2/Phytoplasma%20Vectors%20and%20Diseases%	
470	20in%20Europe%20and%20Surroundings.pdf (accessed 23 April, 2013).	
471	Daugherty, M.P., Gruber, B.R., Almeida, R.P.P., Anderson, M.M., Cooper, M.L.,	
472	Rasmussen, Y.D. & Weber, E.A. (2012) Testing the efficacy of barrier plantings for	
473	limiting sharpshooter spread. American Journal of Enology and Viticulture 63(1), 139-	
474	143.	
475	Garcia-Salazar, C. & Landis, D. (1997) Marking Trichogramma brassicae (Hymenoptera:	
476	Trichogrammatidae) with fluorescent marker dust and its effect on survival and flight	
477	behavior. Journal of Economic Entomology 90, 1546-1550.	
478	Hagler, J.R. (1997) Field retention of a novel mark-release-recapture method. Environmental	
479	Entomology 26 , 1079-1086.	
480	Hagler, J.R. & Jackson, C.G. (2001) Methods for marking insects: current techniques and	
481	future prospects. Annual Review of Entomology 46, 511-543.	
482	Hagler, J.R. & Jones, V.P. (2010) A protein-based approach to mark arthropods for mark-	
483	capture type research. Entomologia Experimentalis et Applicata 135, 177-192.	
484	Jones, V.P., Hagler, J.R., Brunner, J.F., Baker, C.C. & Wilburn, T.D. (2006) An	
485	inexpensive immunomarking technique for studying movement patterns of naturally	
486	occurring insect populations. Environmental Entomology 35(4), 827-836.	
487	Lessio, F. & Alma, A. (2004a) Dispersal patterns and chromatic response of Scaphoideus	
488	titanus Ball (Homoptera: Cicadellidae), vector of the phytoplasma agent of grapevine	
489	flavescence dorée. Agricultural and Forest Entomology 6, 121-127.	

19

- 490 Lessio, F. & Alma, A. (2004b) Seasonal and daily movement of *Scaphoideus titanus* Ball
 491 (Homoptera Cicadellidae). *Environmental Entomology* 33(6), 1689-1694.
- 492 Lessio, F. & Alma, A. (2006) Spatial distribution of nymphs of *Scaphoideus titanus* Ball
 493 (Homoptera Cicadellidae) in grapes, and evaluation of sequential sampling plans.
- 494 *Journal of economic Economic Entomology* **99**(2), 578-582.
- Lessio, F., Tedeschi, R. & Alma, A. (2007) Presence of *Scaphoideus titanus* on American
 grapevine in woodlands, and infection with "flavescence dorée" phytoplasmas. *Bulletin of Insectology* 60, 373-374.
- 498 Lessio, F, Chiusano, P. & Alma, A. (2008) Rilascio e cattura di *Scaphoideus titanus* Ball per
 499 lo studio della dispersione. *Petria* 18(2), 232-233.
- 500 Lessio, F., Tedeschi, R., Pajoro, M. & Alma A. (2009a) Seasonal progression of sex ratio
 501 and phytoplasma infection in *Scaphoideus titanus*, Ball (Hemiptera: Cicadellidae).
 502 Bulletin of Entomological Research 99, 377-383.
- 503
- Lessio F., Borgogno Mondino, E. & Alma, A. (2009b) Spatial correlation of *Scaphoideus titanus* Ball adults on European grapevine at a plot scale: a case study. pp. 166-167 in *Extended abstracts 16th meeting of ICVG, Dijon, 31 August-4 September 2009*. Dijon,
 INRA.
- Lessio, F., Albertin, I., Lombardo, D.M., Gotta, P., Alma, A. (2011a) Monitoring *Scaphoideus titanus* for IPM purposes: results of a pilot-project in Piedmont (NW
 Italy). *Bulletin of Insectology* 64 (Supplement), 269-270.
- Lessio F., Borgogno Mondino, E., Alma, A. (2011b) Spatial patterns of *Scaphoideus titanus*(Hemiptera: Cicadellidae): a geostatistical and neural network approach. *International Journal of Pest Management* 57(3), 205-216.
- Malembic-Maher, S., Salar, P., Filippin, L., Carle, P., Angelini E. & Foissac X. (2011)
 Genetic diversity of European phytoplasmas of the 16SrV taxonomic group and

Formatted: Font: Bold, English (U.K.)
Formatted: English (U.K.)
Formatted: Indent: Left: 0", Hanging: 0.4
Formatted: English (U.K.)
Formatted: Font: Italic, English (U.K.)
Formatted: English (U.K.)
Formatted: Font: Italic, English (U.K.)
Formatted: English (U.K.)
Formatted: Font: Bold, English (U.K.)
Formatted: English (U.K.)

- proposal of 'Candidatus Phytoplasma rubi' *International Journal of Systematic and Evolutionary Microbiology* 61, 2129–2134.
- 518 Northfield, T.D, Mizell III, R.F., Paini, D.R., Andersen, P.C., Brodbeck, B.V., Riddle,
- T.C. & Hunter, W.B. (2009) Dispersal, patch leaving, and distribution of *Homalodisca vitripennis* (Hemiptera: Cicadelldae). *Environmental Entomology* 38(1), 183-191.

521 Orenstein, S., Zahavi, T., Nestel, D., Sharon, R., Barkalifa, M. & Weintraub, P.G. (2003)

- 522 Spatial dispersion of potential leafhopper and planthopper (Homoptera) vectors of 523 phytoplasma in wine vineyards. *Annals of applied Applied Biology* **142**, 341-348.
- 524 Päts, P. & Vernon, R.S. (1999) Fences excluding cabbage maggot flies and tiger flies
 525 (Diptera: Anthomyidae) from large planting of radish. *Environmental Entomology*526 28(6), 1999.
- 527 Pavan, F., Mori, N., Bigot, G. & Zandigiacomo, P. (2012) Border effect in spatial
 528 distribution of Flavescence dorée affected grapevines and outside source of
 529 Scaphoideus titanus vectors. Bulletin of Insectology 65 (2), 281-290.
- **Rhodes, E.M., Liburd, O.E. & Grunwald, S.** (2011) Examining the spatial distribution of
 flower thrips in southern highbush blueberries by utilizing geostatistical methods. *Environmental Entomology* 40, 893-903.
- Saracco, P., Marzachi, C. & Bosco, D. (2008) Activity of some insecticides in preventing
 transmission of chrysanthemum yellows phytoplasma ("Candidatus Phytoplasma
 asteris") by the leafhopper *Macrosteles quadripunctulatus* Kirschbaum. *Crop Protection* 27(1), 130-136.
- 537 Sheather, S.J. & Jones, M.C. (1991) A reliable data-based bandwidth selection method for
 538 kernel density estimation. *Journal of the Royal Statistical Society* 53(3), 683-690.
- 539 Skovgärd, H. (2002) Dispersal of the filth fly parasitoid *Spalangia cameroni* (Hymenoptera:
 540 Pteromalidae) in a swine facility using fluorescent dust marking and sentinel pupal
- 541 bags. Environmental Entomology **31**, 425-431.

542	Slosky, L.M., Hoffmann, E.J. & Hagler, J.R. (2012) A comparative study of the retention	
543	and lethality of the first and second generation arthropod protein markers. Entomologia	
544	Experimentali <u>s</u> et Applicata 144 , 165-171.	
545	Sokal, R.R. & Rohlf, F.J. (1995) Assumption of analysis of variance pp. 392-450 in Sokal,	
546	R.R. & Rohlf, F.J. (Eds.) Biometry: the principles and practice of statistics in	
547	biological research. New York, Freeman & co.	
548	Takken, W., Charlwood, J.D., Billingsley, P.F. & Gort, G. (1998) Dispersal and survival	
549	of Anopheles funestus and A. gambiae s.l. (Diptera: Culicidae) during the rainy season	
550	in southeast Tanzania. Bulletin of Entomological Research 88, 561-566.	
551	Tillman, P.G, Northfield, T.D., Mizell, R.F. & Riddle, T.C. (2009) Spatiotemporal patterns	Formatted: Font: Bold
552	and dispersal of stink bugs (Heteroptera: Pentatomidae) in peanut-cotton farmscapes.	
553	Environmental Entomology 38, 1038-1052.	
554	Vernon, R.S. & MacKenzie, J.R. (1998) The effect of exclusion fences on the colonization	
555	of rutagabas by cabbage flies (Diptera: Anthomyidae). The Canadian Entomologist	
556	130 , 153-162.	
557	Vidano, C. (1964) Scoperta in Italia dello Scaphoideus littoralis Ball cicalina americana	
558	collegata alla "Flavescence dorée" della vite. L'Italia Agricola 88, 1031-1049.	
559	Weber, A. & Maixner, M. (1998) Survey of populations of the planthopper Hyalesthes	
560	obsoletus Sign. (Auchenorrhyncha, Cixiidae) for infection with the phytoplasma	
561	causing grapevine yellows in Germany. Journal of Applied Entomology 122, 375-381.	
562	Zhou, L., Hoy, C.W., Miller, S.A. & Nault, L.R. (2003) Marking methods and field	
563	experiments to estimate aster leafhopper (Macrosteles quadrilineatus) dispersal rates.	
564	Environmental Entomology 32 (5), 1177-1186.	Formatted: Font: Italic

	Formatted Table						Formatted: Right: -0.08"	
	AP ↓ Jul Sept. Jul Oct.	Jul Sept. Jul Oct.	Jul Sept.	Aug Sept. Jul Oct.	Aug Sept. JulOct.	Jul Oct. Jul Oct.	 c. 25 July), except- nymphs/5 leaves ing the season; *: 	23
	S * 8	× × * *	5 *	× ×	2 % * *	7 ** 7	hyl (appro. <i>S. titanus</i> d grapevir ication dur ication dur	
	N _{WGV} 6 4	9	4	4 ω	4 ω	m 0	irifos-metl density of ads of wil kers' appl kers'	23
	29 29 29	17 20	19	23 23	16 20	24	I Chlorpi y; STN: rof marl r of marl	
	6	14	9	20	220	$120 \\ 110$	ume) and of stud metres 1 : numbe	
	STN 0.05 0.14	0.01 0.01	0.31	0.18 0.08	0.01 0.03	0.05	rox. 26 J Y s: yeau tance in ards; N _m ards; N	
	Y _s 2010 2011	2010 2011	2010	2010 2011	2010 2011	2011	am (app planting; mum dis in viney in viney	
	Y _P 2004	2008	1970	1981	2004	2008	hiametox year of 1 mini: mini of traps of traps	
	Sv 2780	1500	1800	2800	2550	8600	sd with T 1 m ² , Y _F ; (006). D _F : number	
l sites and marker -applications.	CvVariety Barbera	Grignolino	Freisa	Barbera	Barbera	Barbera, Grignolino, Ruché	pevine. All vineyards (Vin.) were treate the same dates; Sv: size of vineyards, in ential sampling plan (Lessio & Alma, 2 re were 2 separate stands of WGV); Nv, luring the season.	
features of the experimenta	<u>Coordinates (°N; E)</u> 44.965299; 8.252597	44.965215; 8.252018	44.946083; 8.247651	44.970248; 8.252081	<u>44.968798; 8.249197</u>	44.962938; 8.260826	f vineyards and stands of wild grateated twice with Etofenprox on 1 vineyard, calculated with a seque on stands of WGV (in site D ther P: application period of markers d	23
1. Main	Vin. A-1	A-2	В	C-1	C-2	D	onsisted o that was t in the in the r of traps to milk, A.	
Table	Site		В	C		D	Sites c vin. B per pla number egg. **	

Marker	Year	<u>N</u>	<u>T</u>	Independent	b	s.e	t	P 🔸	Formatted: Centered
				variable					Formatted Table
Egg	2010	<u>5</u>	<u>24</u>	Intercept	0.83	0.13	6.27	0.00 •	Formatted: Centered
				Time	-0.01	0.01	-0.63	0.54 🔺	Formatted: Centered
		_		Rainfall	-0.00	0.01	-0.91	0.38 •	 Formatted: Centered
	2011	<u>8</u>	<u>17</u>	Intercept	1.06	0.14	7.47	0.00 •	 Formatted: Centered
				Time	-0.01	0.01	-0.69	0.52 •	 Contered
				Rainfall	-0.01	0.01	-0.70	0.51 •	 Formatted: Centered
Milk	2011	7	2	Intercept	-0.15	0.13	-1.21	0.29 •	Formatted: Centered
		—	_	Time	0.04	0.01	2.99	0.04 🔺	Formatted: Centered
				Rainfall	-0.01	0.01	-0.94	0.40 🔺	Formatted: Centered
•									Formatted: Centered

Table 2. Results of weighted least square (WLS) regression of marked *S. titanus* as a function of rainfall and time.

Dependent variable: rate of marked *S. titanus* (previously arcsin square root transformed) collected on traps placed on wild grapevine (WGV) at each observation, without considering differences between experimental sites; N: number of observations during the season; T: number of traps observed; independent variables: rainfall occurred (mm) and time elapsed (days) from between marker's application on WGV and insects' collection; weight variable: total insects captured (marked + unmarked) on traps placed on WGV at each observation.

 State
 Comparison
 <thComparison</th>
 Comparison
 Comparison

year	site	males		females		Sex ratio (m/f)		Homogeneity of		
ĺ		total	marked	total	marked	total	marked	F	df	, P
2010	A*	276	115	549	188	0.50	0.61	1.10	1,21	0.31
	B*	255	85	4065	86	0.06	0.99	0.05	1,7	0.83
	C*	12	4	151	51	0.08	0.08	0.81	1,21	0.38
2011	A*	755	455	1377	739	0.55	0.62	0.17	1,21	0.68
	C*	298	197	761	406	0.39	0.49	1.88	1,23	0.18
	D*	150	92	386	171	0.39	0.54	0.18	1, 11	0.68
	D**	150	25	386	72	0.39	0.35	2.84	1,11	0.12

Dependent variable: rate of marked *S. titanus* males and females (<u>marked/total</u>) previously arcsin square root ransformed); independent variable: distance from treated WGV. *: egg; **: milk; df: degrees of freedom.

year	site	intercept	slope	\mathbb{R}^2	Р	r _{0.5}
2010	A*	8.27	0.05	0.56	< 0.05	13.86
	B*	9.51	0.03	0.48	< 0.05	23.10
	C*	73.43	0.04	0.61	< 0.05	17.33
2011	A*	55.69	0.05	0.80	< 0.05	13.86
	C*	4.19	0.02	0.84	< 0.05	34.66
	D*	29.13	0.01	0.34	< 0.05	69.31
	D**	6.2	0.01	0.12	< 0.05	69.31

Table 4. Results of exponential regression of marked *S. titanus* adults as a function of minimum distance from wild grapevine (WGV).

Dependent variable: <u>percentage of marked S. titanus captured during the whole season at the same minimum distance</u> from treated wild grapevine (WGV), weighted by the number of traps placed at the same distanceper trap; independent variable: minimum distance from treated wild grapevine (WGV) (see text for details). *: egg; **: milk; $r_{0.5}$: mean dispersal index (in metres).

Formatted: Right: -0.12"

-					
year	site	interpolation	ME	RMSE	
		method			
2010	A*	IDW	-1.27	7.85	
	A*	KB	0.70	6.51	
	B*	IDW	-1.06	5.58	
	B*	KB	0.70	5.73	
	C*	IDW	-0.72	1.51	
	C*	KB	0.22	1.20	
2011	A*	IDW	-4.48	42.90	
	A*	KB	-0.88	14.23	
	C*	IDW	-2.38	14.12	
	C*	KB	0.31	12.71	
	D *	IDW	-1.54	15.26	
	D *	KB	2.32	19.26	
	D **	IDW	-0.39	6.18	
	D **	KB	0.21	2.70	

Table 5. Results of cross-validation analysis on the interpolation maps of marked *S. titanus* adults.

*: egg; **: milk; IDW: Inverse Distance Weighting; KB: Kernel interpolation with Barriers; ME: Mean Error; RMSE: Root Mean Square Error.

Figure captions

Fig. 1. Captures of *Scaphoideus titanus* adults on stands of wild grapevine (WGV) and in vineyards within the different experimental sites, and rate of marked specimens (*: egg; **: milk). A: 2010; B: 2011.

Fig. 2. Cumulative distribution frequencies Frequencies (F) and cumulative frequencies (CF) of marked *Scaphoideus titanus* adults (CF marked) as a function of minimum distance (Dmin) from treated stands of wild grapevine (WGV) in the different experimental sites: A: site A (vineyards A-1 and A-2 + 1 WGV); B: site B (vineyard B + 1 WGV); C: site C (vineyards C-1 and C-2 + 1 WGV close to C-1); D: site D (vineyard D + 2 WGV); *: egg; **: milk.

Fig. 3. Interpolation maps of marked *Scaphoideus titanus* captures in site A. IDW: inverse distance weighting; KB: kernel interpolation with barriers. A: IDW, 2010; B: IDW, 2011; C: KB, 2010; D: KB, 2011. Dots represent the position of yellow sticky traps (sampling points).

Fig. 4. Interpolation maps of marked *Scaphoideus titanus* captures in site B. IDW: inverse distance weighting; KB: kernel interpolation with barriers. A: IDW, 2010; B: KB, 2010. Dots represent the position of yellow sticky traps (sampling points).

Fig. 5. Interpolation maps of marked *Scaphoideus titanus* captures in site C, IDW: inverse distance weighting; KB: kernel interpolation with barriers. A: IDW, 2010; B: IDW, 2011; C: KB, 2010; D: KB, 2011. Dots represent the position of yellow sticky traps (sampling points).

Fig. 6. Interpolation maps of marked *Scaphoideus titanus* captures in site D, obtained with Inverse distance weighting (IDW) or kernel interpolation with barriers (KB). A: IDW, egg,

2011; B: IDW, milk, 2011; C: IDW, egg + milk, 2011; D: KB, egg, 2011; E: KB, milk, 2011;F: KB, egg + milk, 2011. Dots represent the position of yellow sticky traps (sampling points).

























Figure 6