

This is a postprint version of the following published document:

Planchuelo, Greg; Catalán, Pablo; Delgado, Juan Antonio. (2016). Gone with the wind and the stream: dispersal in the invasive species *Ailanthus altissima*. *Acta Oecologica*, v. 73, pp.: 31-37.

DOI: <https://doi.org/10.1016/j.actao.2016.02.006>

© 2016 Elsevier Masson SAS. All rights reserved.



This work is licensed under a [Creative Commons Attribution-NonCommercialNoDerivatives 4.0 International License](https://creativecommons.org/licenses/by-nc-nd/4.0/).

Manuscript Number: ACTOEC-D-15-00380R1

Title: Gone with the wind and the stream: Dispersal in the invasive species *Ailanthus altissima*

Article Type: Full length article

Keywords: Anemochory; Hydrochory; Primary dispersal; Repeatability; Samara; Secondary dispersal

Corresponding Author: Mr. Greg Planchuelo,

Corresponding Author's Institution: Technische Universität Berlin

First Author: Greg Planchuelo

Order of Authors: Greg Planchuelo; Pablo Catalán; Juan A Delgado

Abstract: Dispersal is a key process in plant invasions and is strongly related to diaspore morphology. Often, dispersal comprises more than one step, and morphologies adapted to a primary dispersal mechanism can aid or detract from a secondary one. The aim of this work was to assess the relationship between primary wind dispersal and secondary water dispersal in *Ailanthus altissima*, an invasive tree species. Wind and water dispersal potential and their association with the morphological characteristics of samaras were assessed under controlled conditions to ensure the repeatability of the measurements. We found a direct positive relationship between primary wind and secondary water dispersal in *A. altissima*. The main morphological characteristics of the samara that affected the success of the two types of dispersal were side perimeter and mass. However, a possibility of dispersal specialisation exists, as one morphological characteristic (samara width) affects wind dispersal negatively but water dispersal positively, and dispersal potential and samara morphology have been shown to differ across individuals.

Dear Editor,

Here we present a revision of manuscript Ref N° ACTOEC-D-15-00380 that has been accepted for publication in *Acta Oecologica*. The following changes have been made:

- Suggestions regarding language and expression have been taken into consideration and English has been corrected and polished in detail by Kelanie Radvin, a professional American proof reader specialised in ecology.
- All minor suggestions have also been corrected and polished along the following highlights:
 - Lines 71-72: Description of the soil is now more explanatory.
 - Lines 115-121: Water flow is now explained in detail by making a difference between water flow across the whole channel profile and water speed at the surface. This clears the incongruences regarding samaras going seemingly too fast.
 - Lines 122-130: The intervals of measurements for floating samaras have been made clearer.
 - Figure 3: All trendlines have now the same colour for better visualisation.

We hope our revised version will be received favourably and look forward to hearing from you in the near future. Thank you very much for your time and kind regards,



Greg Planchuelo

greg.planchuelo@gmail.com - Department of Ecology, Technische Universität Berlin

Highlights

We study the relationship between primary wind and secondary water dispersal in *Ailanthus altissima* in relation to its samara morphology.

There is a direct positive relationship between primary and secondary dispersal in this species.

The main morphological traits of the samara that determine both types of dispersal are side perimeter and mass.

Dispersal potential and samara morphology varies between different individuals.

Our findings suggest the possibility of specialization of dispersal modes in *Ailanthus altissima* into different environments.

1 **Gone with the wind and the stream: Dispersal in the**
2
3
4 **invasive species *Ailanthus altissima***
5
6

7 **Greg Planchuelo ¹ · Pablo Catalán ² · Juan Antonio Delgado ³**
8
9

10
11
12
13
14
15
16
17
18
19
20
21
22
23
24 ¹ G. Planchuelo (corresponding author)

25
26 Department of Ecology, Technische Universität Berlin, Rothenburgstrasse 12, 12165

27
28 Berlin, Germany

29
30 e-mail: greg.planchuelo@gmail.com

31
32 Telephone: 0049 17698828226
33
34
35

36 ² P. Catalán
37

38 Interdisciplinary group of complex systems, Department of Mathematics, Universidad

39
40 Carlos III, Madrid, Spain
41
42

43
44 ³ J A. Delgado
45

46 Department of Ecology, Universidad Complutense, C/José Antonio Nováis 2, 28040

47
48 Madrid, Spain
49
50
51
52
53

54
55 *All authors have approved the final version of this article and declare to have participated*
56
57 *equally in its design, accomplishment and preparation.*
58
59
60
61
62
63
64
65

1 **Abstract** Dispersal is a key process in plant invasions and is strongly related to diaspore
2 morphology. Often, dispersal comprises more than one step, and morphologies adapted to a
3 primary dispersal mechanism can aid or detract from a secondary one. The aim of this work was
4 to assess the relationship between primary wind dispersal and secondary water dispersal in
5 *Ailanthus altissima*, an invasive tree species. Wind and water dispersal potential and their
6 association with the morphological characteristics of samaras were assessed under controlled
7 conditions to ensure the repeatability of the measurements. We found a direct positive
8 relationship between primary wind and secondary water dispersal in *A. altissima*. The main
9 morphological characteristics of the samara that affected the success of the two types of dispersal
10 were side perimeter and mass. However, a possibility of dispersal specialisation exists, as one
11 morphological characteristic (samara width) affects wind dispersal negatively but water dispersal
12 positively, and dispersal potential and samara morphology have been shown to differ across
13 individuals.

14 **Keywords:** Anemochory, Hydrochory, Primary dispersal, Repeatability, Samara, Secondary
15 dispersal

16 Introduction

17 A successful invasion comprises three stages: dispersal of diaspores, formation of a new self-
18 sufficient population and, finally, the spread of this new population to nearby habitats (Leung et
19 al., 2002; Williamson and Fitter, 1996). Therefore, diaspore dispersal mechanisms are crucial
20 processes in understanding plant invasions, and as such, their study is key to managing plant
21 invasions effectively (Leung et al., 2002).

22 Seed dispersal syndromes define the dispersal strategy of a diaspore and have been studied in
23 depth on numerous occasions (Howe and Smallwood, 1982; Van der Pijl, 1982; Wheelwright and
24 Orians, 1982). Diaspores may be dispersed ballistically, in which case fruits spring open abruptly
25 and explosively release them. Animals can also disperse diaspores that are partly edible or that
26 attach to them by means of hooks or sticky surfaces. Wind-dispersed diaspores, on the other hand,
27 commonly have light structures that can act as wings, plumes or balloons, thus decreasing fall
28 velocity and increasing dispersal distance (Augspurger, 1986; Matlack, 1987). Diaspores can also
29 be dispersed by water, if they are able to float and resist water damage (Säumel and Kowarik,
30 2013).

31 Diaspore dispersal is not always a single-step process, and multiple vectors (animals, wind or
32 water) may be involved (Vander Wall et al., 2005). While primary vectors move diaspores away
33 from the parent plant, secondary vectors can dramatically increase the transport distances
34 (Nathan et al., 2008; Säumel and Kowarik, 2013). Although if there is promising research in
35 estimating dispersal distance (Soons et al., 2004; Tackenberg, 2003; Tackenberg et al., 2003), the
36 relationship between dispersal and diaspore morphology is not yet completely understood
37 (Higgins et al., 2003), as it is a complex multi-scale process that may involve different vectors
38 (Nathan et al., 2008). Furthermore, diaspore morphologies adapted for a primary dispersal
39 mechanism can indirectly favour or dampen secondary dispersal mechanisms (Hintze et al., 2013;
40 Kowarik and Säumel, 2008).

41 Many invasive tree species have diaspores adapted for wind dispersal (Burrows, 1986). The
42 morphological adaptations for wind dispersal can also render diaspores well adapted for

43 dispersing through water, since features such as a low mass or a high surface area are suited to
44 both (Nilsson et al., 2010; Säumel and Kowarik, 2013). Wind can move diaspores long distances
45 (Thomson et al., 2011; Vittoz and Engler, 2007), and those that fall along a watercourse could
46 potentially disperse even further (Poschlod and Bonn, 1998; Säumel and Kowarik, 2013). The
47 relationship between wind and water dispersal remains uninvestigated, as is the influence of
48 diaspore morphology on each dispersal mechanism (Higgins et al., 2003). Nonetheless, selective
49 pressure will likely benefit plants that are good at both methods of dispersal. Since wind dispersal
50 is multi-directional, it can enable propagules to land in a wide range of new habitats, while water
51 dispersal increases the probability of propagules landing in a suitable area (i.e. close to a water
52 source) and can amplify wind dispersal transport distances by a factor of at least 20 (Säumel and
53 Kowarik, 2013).

54 Here we evaluate the relationship between dispersal and diaspore morphology in the invasive
55 tree *Ailanthus altissima* (Miller) Swingle. This species uses wind as primary and water as
56 secondary dispersal vector (Kowarik and Säumel, 2008). It has also been reported to spread along
57 roads, railways and water courses (Kowarik and Säumel, 2007; Merriam, 2003), where the
58 relative relevance of the two mechanisms should vary. In this study, we consider the following: 1)
59 the relationship between wind dispersal potential and water dispersal potential in *A. altissima*, 2)
60 the role of samara morphology in both types of dispersal and 3) differences in the dispersal
61 capabilities of individual *A. altissima* trees.

62
63

64 **Materials and methods**

65 *Study area and species*

66 Field work was carried out on the campus of the Complutense University of Madrid. This is an
67 urban campus located in the city of Madrid (Central Spain, 40° 27' 4" N, 3° 43' 35" W, at 645 m
68 above sea level). The climate is Mediterranean, semi-arid and continental with a mean annual
69 temperature and rainfall of 14.6°C and 530 mm, respectively. Soils are characterised as luvisols
70 (FAO-UNESCO soil classification system) and are siliceous, sandy and nutrient-poor with a slightly
71 acidic pH (Nombela G, 1994). Structurally, they are highly developed soils with good air diffusion
72 and low waterlogging capabilities.

73 *A. altissima* is a tree from the Simaroubaceae family native to China that is currently widespread
74 across all continents except in Antarctica. It is classified as a "noxious weed" and invasive species
75 in many regions for its rapid growth, allelopathic effects, extensive root system and ability to
76 reproduce quickly via diaspores and clonal growth (Kowarik and Säumel, 2007; Lawrence et al.,
77 1991). The plant grows 8–18 m tall, with females producing up to 325,000 samaras per year (Bory
78 and Clair Maczulajtys, 1981). These samaras are adapted to wind dispersal and have one seed in
79 the centre of each wing. Diaspores rotate along their axis and are rigid and sturdy (Kowarik and
80 Säumel, 2007) which enables a variety of flying methods, with autorotation being the most
81 common (Lentink et al., 2009; Yasuda and Azuma, 1997). Water dispersal has also been reported
82 in this species (Kaproth and McGraw, 2008; Kowarik and Säumel, 2008).

83 *Sampling, measurements and analysis*

84 In January 2013, we randomly selected seven female *A. altissima* trees growing spontaneously in
85 open spaces in the campus. We collected roughly 50 samaras from each tree, discarded the
86 damaged ones, and retained 242 samaras for our measurements (40 samaras per tree except for
87 two trees, from which only 19 and 23 samaras were used). Samaras were weighed to the nearest
88 0.1 mg and individually stored in paper bags until morphological and dispersal measurements
89 were taken.

90 Samara morphology was described from a frontal and a side view (Figure 1). Pictures of
 91 individual samaras were taken from a tripod placed at a fixed distance and using a fixed focal
 92 length. We included a scaled ruler as a reference to calculate distances with Adobe Photoshop CS6
 93 and Image J v1.47. In this way, we measured specific morphological attributes of samaras that are
 94 related to wind and/or water dispersal. They included frontal area, side area and frontal
 95 perimeter, which are closely related to the wind and water dispersal potential of samaras as the
 96 total surface area is a function of these values (Nilsson et al., 2010; Säumel and Kowarik, 2013);
 97 and samara width, which is closely related to the autorotation and flotation potential of the
 98 samara (Lentink et al., 2009). We also measured other morphological variables to obtain a more
 99 complete description of the samaras' morphology (Table 1).

Table 1 Average measurements \pm standard deviations and range of the variables used to characterise samaras of *A. altissima* (n=242). Measurement units are shown in parentheses.

Variable description	Average \pm SD	Range
<i>Morphology</i>		
Frontal Area (cm ²)	2.957 \pm 0.581	1.397–4.398
Frontal Perimeter (cm)	10.529 \pm 0.966	7.797–13.321
Width (cm)	1.225 \pm 0.181	0.698–1.764
Length (cm)	4.371 \pm 0.415	3.293–5.391
Side Area (cm ²)	1.259 \pm 0.284	0.700–2.664
Side Perimeter (cm)	9.506 \pm 0.920	7.095–11.85
Side Height (cm)	0.635 \pm 0.127	0.323–1.233
Mass (mg)	34.3 \pm 7.6	20.70–59.10
<i>Dispersal</i>		
Descent Velocity (m/s)	1.106 \pm 0.215	0.703–1.705
Drifting Velocity (m/s)	0.530 \pm 0.011	0.481–0.554
Floating Time (days)	2.074 \pm 0.589	1.0–4.0

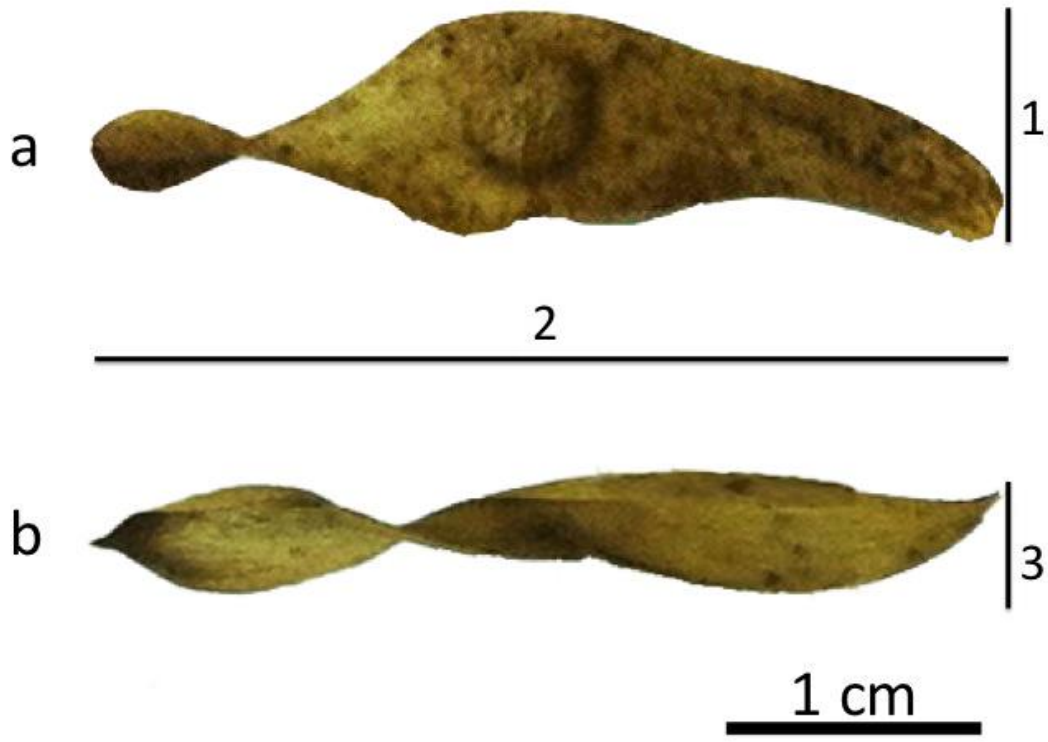


Figure 1 Some morphological measurements of the samaras from a frontal view (a) and side view (b): width (1); length (2); side height (3).

We estimated samara wind dispersal potential by measuring their average descent velocity (Greene and Johnson, 1993; Landenberger et al., 2006). This trait is inversely related to dispersal distance (Greene and Johnson, 1989; Nathan et al., 2011; Tackenberg, 2003), and it was quantified by dropping the samaras inside an airtight and sealed chamber (Greene and Johnson, 1993) from a height of 2.0 m. Each samara was dropped in the same manner three times, and the time it took to reach the ground was recorded with a stop watch (Greene and Johnson, 1993; Landenberger et al., 2006). Descent velocity was calculated as height divided by time to reach the ground. Average descent velocity for all 242 samaras was 1.106 ± 0.215 m/s (range 0.703–1.705 m/s; Table 1).

The water dispersal potential of the samaras was inferred from their drifting velocity and floating time, which respectively relate to their ability to drift downstream and the length of time they can be transported by flotation.

115 To measure drifting velocity we built a polypropylene channel, square in cross section, 4 m long,
116 0.09 m wide and 0.05 m high. Water flow was empirically measured to be 0.108 L/s (roughly 0.25
117 m/s) across the whole channel profile and 0.591 m/s at the water surface. We recorded the time it
118 took for each samara to traverse the 4 m channel and calculated drifting velocity as channel length
119 divided by the time recorded. Each samara was released from the same place in the same position
120 three times; between measurements they were allowed to dry in the open air for a week. Average
121 drifting velocity for all 242 samaras was 0.530 ± 0.011 m/s (range 0.481–0.554 m/s; Table 1).

122 To measure the floating capabilities of samaras, we placed them in individual water containers
123 with 55 ml of distilled water. We then placed the containers in an orbital shaker at 150 rpm for 12
124 min to allow the floating samaras to become fully impregnated with water. In this way we
125 eliminated differences in floating time due to the way the samara lands in the water (i.e., whether
126 the samara breaks the water surface tension). Finally, we placed the containers into an airtight
127 chamber which was observed at 24 h intervals and registered the time it took for the samaras to
128 sink. This process was repeated three times, allowing the samaras to dry for a week between
129 measurements. Average floating time for all 242 samaras was 2.074 ± 0.589 days (range 1–4 days;
130 Table 1).

131 We performed three different measurements of each dispersal variable to test the repeatability
132 and consistency of our protocol and, at the same time, to generate for each samara an average
133 value to be used in the statistical analyses. Throughout the course of this experiment, samaras
134 were not painted, coloured, written on, modified or altered in any way. They were placed inside
135 labelled paper bags for individual identification.

136 *Statistical analysis*

137 The consistency and repeatability of our dispersal measurements were evaluated in terms of the
138 intraclass correlation coefficient (ICC) for the repeated measurements made on each samara. The
139 relationship between wind and water dispersal was assessed by Pearson correlation analyses. To
140 explore the potential effect of each morphological variable on dispersal estimates, we used single
141 regression analysis. Then, the best models relating dispersal estimates and the morphological

142 attributes of samaras were obtained using an information-theoretical approach and identity as
143 the link function, since dispersal estimates were normally distributed. The selection of the most
144 parsimonious set of parameters was based on fit to the data and number of variables of the model,
145 according to the Akaike information criterion corrected for small sample sizes (AICc; Johnson and
146 Omland, 2004). Although the lowest AICc indicates the model that best fits the data (Hosmer and
147 Lemeshow, 1989), models with $\Delta AICc \leq 2.0$ are considered equally informative (Burnham and
148 Anderson, 2002). Wald statistics were used to assess the significance of regression coefficients in
149 the selected models. Collinearity amongst independent variables was assessed with the variance
150 inflation factor (VIF). To test for differences in samara morphology between individual *A. altissima*
151 trees, MANOVA and principal component analysis (PCA) were used. Finally, to test for differences
152 in dispersal potential between individual trees, MANOVA was used followed by univariate
153 ANOVAs and Tukey's HSD tests. All analyses were performed with SPSS v21 (IBM).

154

155 **Results**

1 156 The ICC results show that, except for drifting velocity, our dispersal measurements were
2
3 157 consistent across repetitions (Fleiss and Cohen, 1973; Lew and Doros, 2010). The lack of
4
5 158 consistency in measurements of drifting velocity could be mainly due to subtle changes in water
6
7
8 159 velocity during the three experimental trials, rather than being the effect of repeated rewetting of
9
10 160 samaras, as no trend was found when we later performed the floatability measurements.
11
12

13 161

16
17 **Table 2** Intraclass correlation coefficient (ICC) results and average
18 measurements of each repetition for wind and water dispersal. ICC
19 results are calculated from the individual values of each of the 242
20 samaras from each repetition.
21
22

	Descent Velocity (m/s)	Drifting Velocity (m/s)	Floating Time (days)
ICC	0.8	0.3	0.7
Repetition 1	1.07 ±0.22	0.56 ±0.13	2.16 ±0.53
Repetition 2	1.01 ±0.24	0.54 ±0.14	1.95 ±0.60
Repetition 3	1.11 ±0.23	0.50 ±0.14	2.14 ±0.50

23
24
25
26
27
28
29
30
31
32
33
34 162

35
36
37
38 163 Pearson correlation results showed a significant inverse relationship between descent velocity
39
40 164 and floating time ($p=0.001$; $r=-0.206$), meaning that samaras with slower descent rates will have
41
42 165 longer flotation times (Fig. 2). On the other hand, there was a positive relationship between
43
44 166 descent velocities and drifting velocities of samaras ($p=0.05$; $r=0.189$; Fig. 2). Finally, there was no
45
46
47 167 statistically significant relationship between drifting velocities and floating times ($p=0.48$).
48
49

50 168

51
52
53 169

54
55
56 170

57
58
59
60
61
62
63
64
65

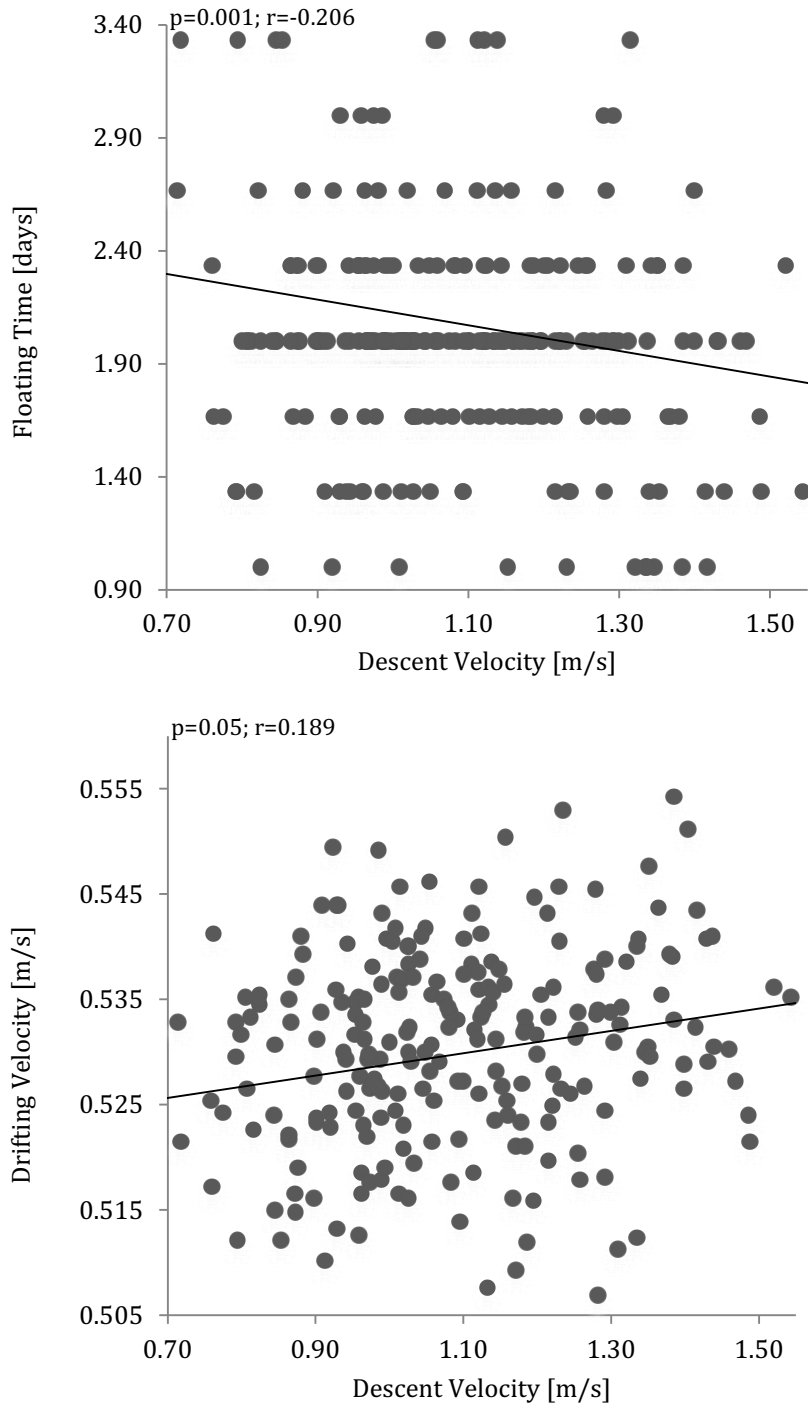


Figure 2 Relationship between wind and water dispersal in *A. altissima*.

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48 171
49
50
51 172
52
53
54 173
55
56
57 174
58
59
60
61 175
62
63
64
65

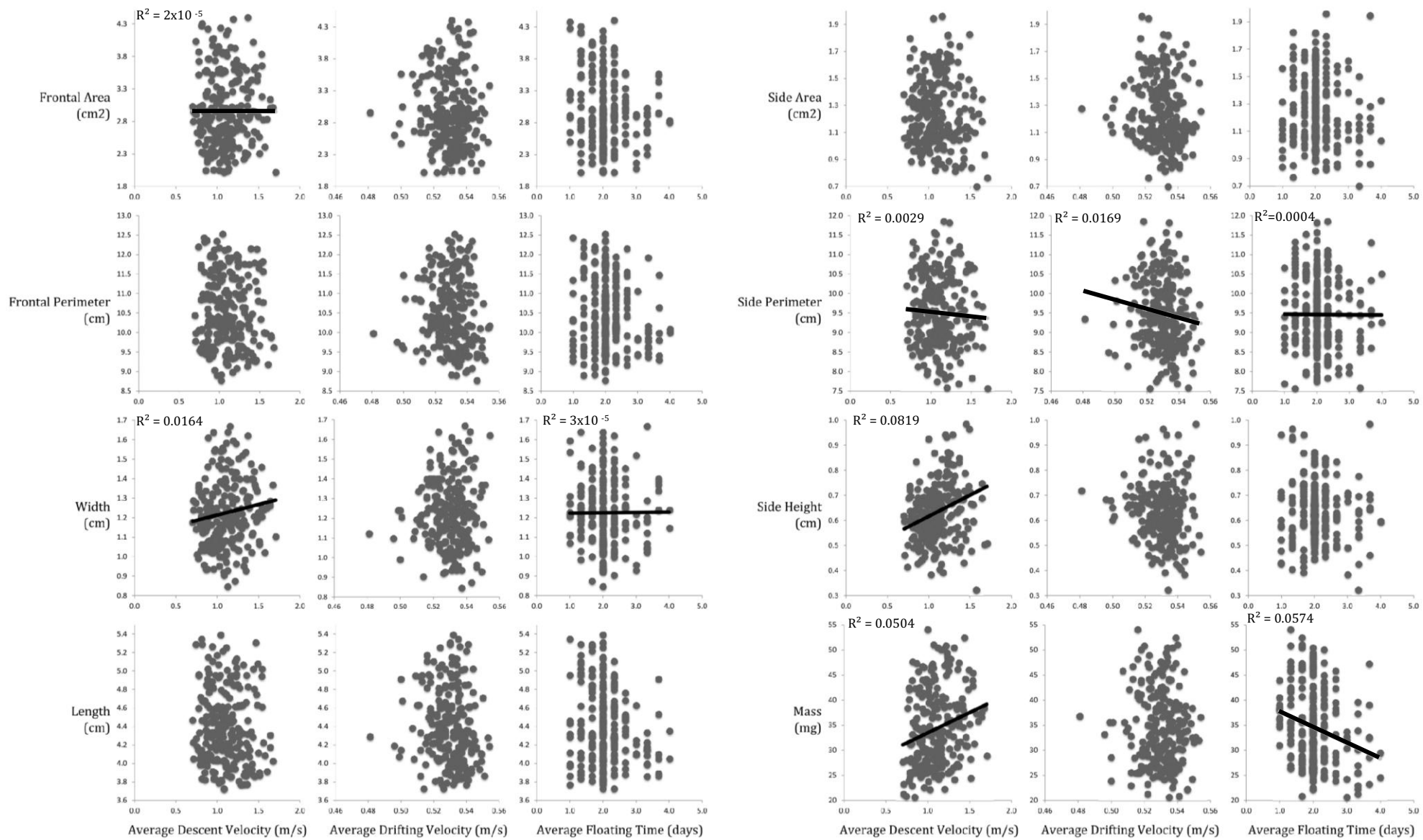


Figure 3 Relationship between samara morphology and dispersal potential in *A. altissima*. Trendlines are shown for significant relationships (n=242).

199 Figure 3 shows the relationship of each morphological variable and dispersal capabilities. Mass
200 had the largest positive effect on descent velocity, followed by width and side height, while
201 frontal area and side perimeter had negative effects on descent velocity. Moreover, drifting
202 velocity was only influenced by side perimeter, which had a negative effect. Finally, the flotation
203 time of samaras was inversely related to mass and, to a lesser extent, positively related to side
204 perimeter and width. It must be highlighted that all of those linear regressions were significant
205 because of the large number of points, but the R^2 values were extremely low.

206 The results of the best subset procedure are shown in Table 3. For descent velocity, six models
207 with $\Delta AICc \leq 2.0$ were obtained, but using the criteria of model complexity, two models with five
208 variables each were considered as the most parsimonious ones (Table 3). Both included frontal
209 area and side perimeter (with negative coefficients), side height and mass (with positive
210 coefficients), and differed in the inclusion of samara width or side area (with positive and
211 negative coefficients; Table 4). There were more than ten models selected for drifting velocity
212 ($\Delta AICc \leq 2.0$), but two of them were considered to be the most parsimonious because of their
213 lower number of variables. In fact, both only selected one variable: the side perimeter or the
214 length of the samara (Table 3), and both were negatively related to drifting velocity (Table 4).
215 Finally, although seven models for floating time presented $\Delta AICc \leq 2.0$, just one of them was
216 considered the most plausible model because of its lower number of variables (Table 3). Floating
217 times of samaras were negatively correlated to mass and positively to width and side perimeter
218 (Table 4). Ordinary linear regression models performed with the same subset of variables
219 showed a relevant improvement of prediction power in relation to that of single variables, but
220 adjusted R^2 values were still low (Table 3).

221
222
223
224

Table 3 Model selection showing which combinations of morphological traits best explain each samara dispersal variable. Models displayed are the ones with the lowest number of variables as long as their $\Delta AICc < 2$.

Model	Predictors	AICc	Adjusted R ² ^a
<i>Descent Velocity</i>			
A	1 5 6 7 8	-803.719	0.239
B	1 3 6 7 8	-803.574	0.239
<i>Drifting Velocity</i>			
A	6	-2196.939	0.013
B	4	-2196.679	0.012
<i>Floating Time</i>			
A	3 6 8	-284.417	0.130

Predictors: (1) frontal area; (2) frontal perimeter; (3) width; (4) length; (5) side area; (6) side perimeter; (7) side height; (8) mass.
^aThe adjusted R² values were calculated by means of ordinary least squares regression to better determine the predictive value of each model.

Table 4 Parameter estimates for the predictor variables included in the best models for descent velocity, drifting velocity and floating time.

Variable	Estimate	Standard Error	Wald test	P
<i>Descent Velocity - Model A</i>				
Intercept	1.367	0.147	86.53	<0.001
Frontal Area	-0.139	0.033	17.81	<0.001
Side Area	-0.134	0.059	5.10	0.024
Side Perimeter	-0.069	0.021	11.31	0.001
Side Height	0.471	0.115	16.82	<0.001
Mass	0.020	0.003	45.96	<0.001
<i>Descent Velocity - Model B</i>				
Intercept	1.312	0.156	70.44	<0.001
Frontal Area	-0.172	0.037	21.65	<0.001
Width	0.208	0.094	4.95	0.026
Side Perimeter	-0.086	0.019	21.13	<0.001
Side Height	0.371	0.103	12.99	<0.001
Mass	0.018	0.003	38.08	<0.001
<i>Drifting Velocity - Model A</i>				
Intercept	0.544	0.007	5919.79	<0.001
Side Perimeter	-0.002	0.001	4.15	0.042
<i>Drifting Velocity - Model B</i>				
Intercept	0.544	0.007	57040.60	<0.001
Length	-0.003	0.002	3.89	0.049
<i>Floating Time</i>				
Intercept	0.823	0.434	3.59	0.058
Side Perimeter	0.196	0.054	13.11	<0.001
Mass	-0.046	0.007	39.23	<0.001
Width	0.798	0.244	10.68	0.001

1 226 The variance inflation factor (VIF) ranges from 1 to 5 (frontal area=4.66; frontal perimeter=2.02;
2
3 227 width=2.22; length=3.88; side area=2.04; side perimeter=2.83; side height=1.52; mass=3.67),
4
5 228 indicating a weak correlation among the independent variables (Belsley et al., 2005).
6
7

8
9 229 Samara morphology was significantly different among the studied trees of *A. altissima*
10
11 230 (MANOVA, $F_{48,1126} = 11.97$, $p < 0.0005$, Wilks' lambda = 0.483). The PCA resulted in two factors
12
13 231 with eigenvalue > 1, explaining 71.2% of the variance. The main principal component was
14
15 232 positively related ($p < 0.01$ in all cases) to mass ($r = 0.885$), frontal area ($r = 0.876$), side
16
17 233 perimeter ($r = 0.827$), width ($r = 0.660$) and side height ($r = 0.425$). The second component was
18
19 234 related ($p < 0.01$ in all cases) positively to side height ($r = 0.751$) and mass ($r = 0.170$) and
20
21 235 negatively to width ($r = -0.321$) and frontal area ($r = -0.247$). Within the samara trait space,
22
23 236 individual trees were segregated on the first axis according to the size of the samaras, whereas
24
25 237 on the second axis they were segregated mainly according to their side perimeter (Fig. 4).
26
27 238 These results show that there were no large differences in samara morphology within individual
28
29 239 trees, but that across individuals there may be great differences. Nevertheless, some individuals
30
31 240 had a broader spectrum of samara morphologies than others (see e.g. plant 2 versus plant 4 in
32
33 241 Fig 4).
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

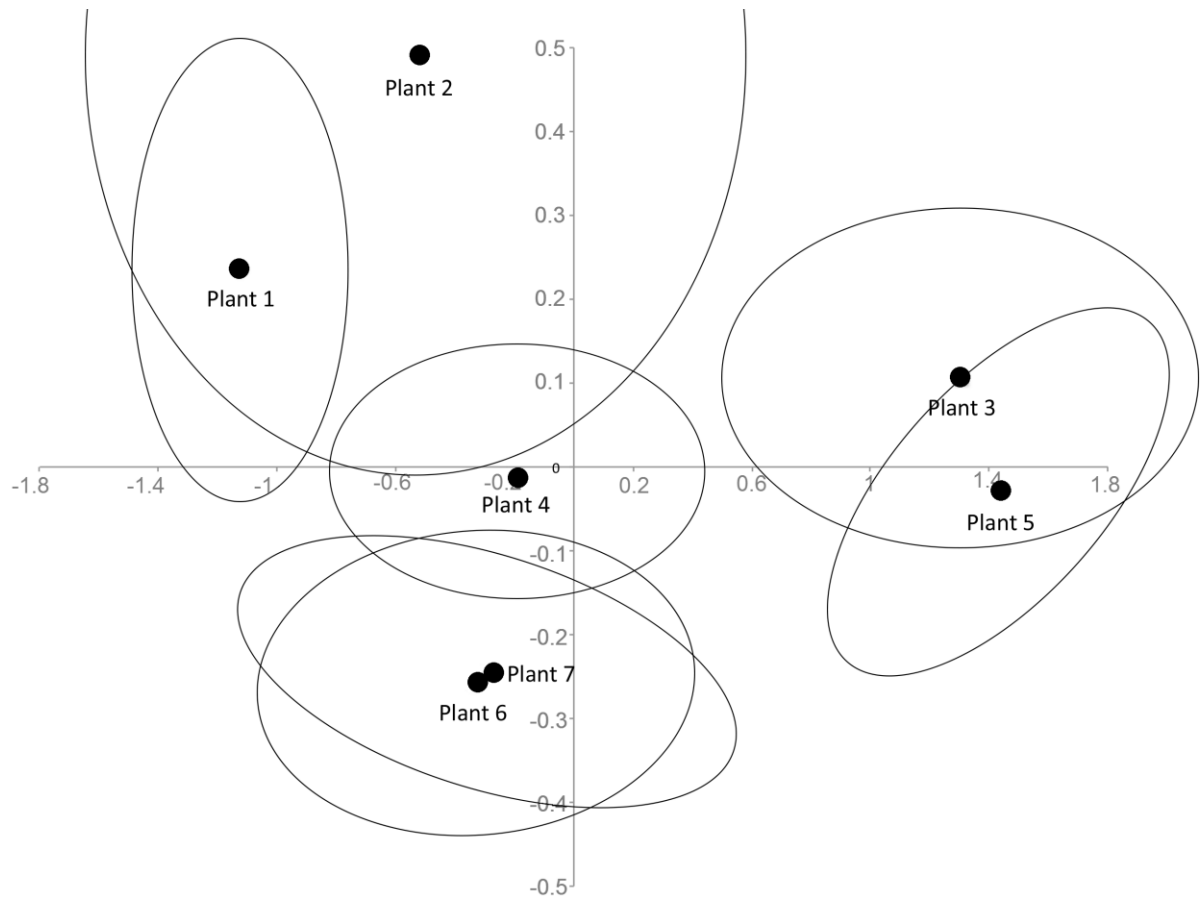


Figure 4 Distribution of individual trees (and their 95% confidence ellipses) in the samara morphology space.

Dispersal capacity was also significantly different among the studied trees of *A. altissima* (MANOVA, $F_{18,660} = 23.76$, $p < 0.0005$, Wilks' lambda = 0.032). Further univariate ANOVAs tell us that individuals differ in the three types of dispersal (Table 5). These results suggest that different dispersal capacities are expected depending on the individual (Fig. 5).

Table 5 Univariate ANOVA results for wind and water dispersal.

	p value	$F_{6, 235}$
Descent Velocity	<0.0005	11
Drifting Velocity	<0.0005	16
Floating Time	<0.0005	5

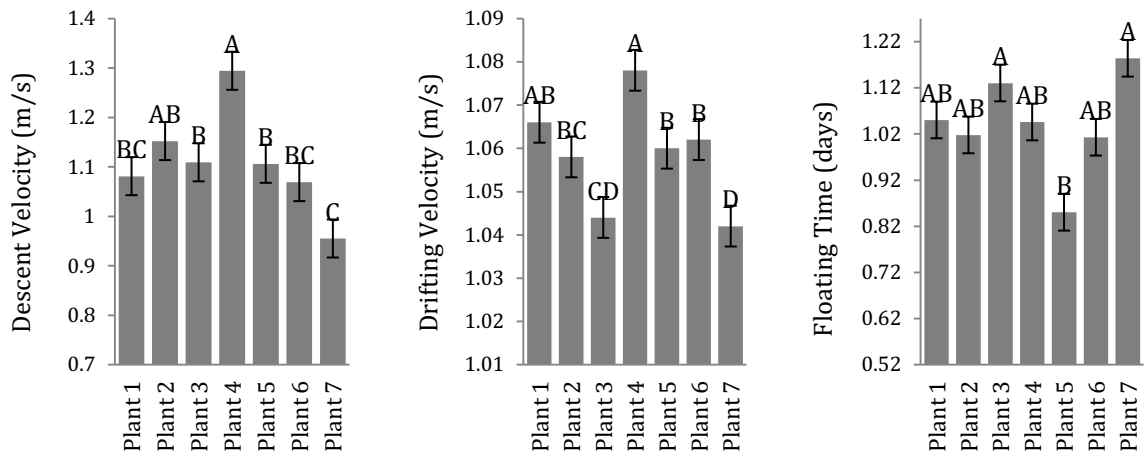


Figure 5 Average values and standard deviations of descent velocity, drifting velocity and floating time per plant. Tukeys HSD's homogeneous plant subsets are ordered by descending mean values and labelled with letters. For all plants n=40 except plants 2 and 3 (n=19 and n=23 respectively).

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

249

250

251 Discussion

1 252 Our results suggest that wind dispersal as a primary vector and water dispersal as a secondary
2
3 253 vector are positively correlated in *A. altissima*. Samaras with slower descent velocities also
4
5 254 showed longer floating times. It has previously been reported that plants whose primary
6
7 255 dispersal method is wind may also be effectively dispersed by water (Soomers et al., 2012). Our
8
9 256 results showed such a positive relationship at the level of individual samaras within a single
10
11 257 species. From an evolutionary standpoint, this positive relationship between wind and water
12
13 258 dispersal suggests that there may be a selective pressure to simultaneously improve both
14
15 259 dispersal potentials in some anemochorous species. This is further supported if we take into
16
17 260 account that both kinds of dispersal are governed, at least in *A. altissima*, by similar
18
19 261 morphological characteristics of the samaras.
20
21
22
23

24 262 Diaspore mass is negatively correlated to both kinds of dispersal, as reported for many wind-
25
26 263 and water-dispersed species (Nilsson et al., 2010; Säumel and Kowarik, 2013). The greater the
27
28 264 mass, the shorter the distance the diaspore will be able to fly (Greene and Johnson, 1993) and
29
30 265 the faster it will sink in water. Diaspore mass has been reported to evolve quickly in wind-
31
32 266 dispersed species (Cheptou et al., 2008), but there was a wide variation in this trait in the
33
34 267 studied population of *A. altissima* (Table 1). These results suggest that there is not a strong
35
36 268 selective pressure on diaspore mass to increase dispersal distance. Alternatively, variation in
37
38 269 samara mass in the studied population could be the result of two counteracting selective
39
40 270 pressures: 1) to reduce samara mass and, therefore, increase dispersal distance; and 2) to
41
42 271 increase samara mass and produce heavier, more competitive seedlings (Delgado et al., 2009).
43
44
45
46
47

48 272 On the other hand, there was a positive correlation between descent and drifting velocities,
49
50 273 suggesting a negative correlation between wind and water dispersal in moving water. This
51
52 274 suggestion will need further research however, since in our study drifting velocities were
53
54 275 inconsistent between repetitions, and the effects of side perimeter or length of the samara on
55
56 276 this measure of dispersal were very weak. In addition, although side perimeter was positively
57
58 277 related to wind dispersal and negatively to drifting velocity, it was also positively related to
59
60 278 floating time, so there was not a consistent opposite effect of this variable on wind and water
61
62
63
64
65

279 dispersal. Furthermore, in field conditions, the weak effect of side perimeter or samara length on
280 drifting velocity might have little effect in determining water transport due to the combined
281 effects of current, waves and wind (Wang et al., 2015). As a consequence, differences in dispersal
282 could depend mainly on river characteristics (Säumel and Kowarik, 2013).

283 Samara width clearly affected dispersal via wind and water differently in *A. altissima*. Although it
284 was one of the less important characteristics in determining both water and wind dispersal
285 capabilities, it opens up a possibility for differentiation in dispersal strategies: narrower
286 samaras dispersing better by wind than by water. Therefore, samara morphology could suffer
287 opposite selection pressures depending on the relevance of each dispersal mode in each habitat
288 type (Venable, 1985).

289 It must be highlighted that, although several morphological characteristics were significantly
290 related to dispersal capabilities in *A. altissima*, there was a large variability in all these cases (i.e.
291 low R^2 values). In other words, every independent variable is a poor predictor of the dispersal
292 capability of the samara. These results are surprising since morphological variables were, and
293 still are, widely used as surrogates of seed dispersal capabilities in both interspecific (Hintze et
294 al., 2013; Smith et al., 2015) and intraspecific comparisons (Bartle et al., 2013; Delgado et al.,
295 2009). The use of diaspore characteristics to assess dispersal potential seems to be promoted
296 because they are easily measurable traits rather than because they are highly correlated to
297 dispersal (Hintze et al., 2013). Obtaining direct dispersal measures is an overwhelming task and
298 thus beyond the scope of many studies, however, single characteristics should not be relied on
299 when inferring dispersal capabilities from diaspore morphology. The simultaneous use of
300 several morphological characteristics substantially increased prediction power as shown in our
301 results.

302 We found that individual plants under the same mesoclimatic conditions had different samara
303 morphologies and consequently differed in their dispersal potential by wind and by water. In
304 addition, strong differences in samara characteristics (area per unit of weight and total weight)
305 have already been described for nearby stands of this plant, indicating that they differ in
306 invasion potential (Delgado et al., 2009). This could be due to genetic or maternal environmental

307 effects even in shared mesoclimatic conditions (Galloway, 2005; Mousseau and Fox, 1998). From
308 the perspective of an invasive species, this broad spectrum of plant traits and dispersal
309 capabilities might be beneficial as it could enable successful settlements in a wide range of
310 habitats (Constán-Nava and Bonet, 2012; Williamson and Fitter, 1996).

311 In conclusion, our results suggest that primary wind dispersal and secondary water dispersal
312 are usually positively correlated in *A. altissima* since most morphological characteristics of
313 samaras affect both dispersal modes in the same way. Samaras with low mass and a large side
314 perimeter had larger dispersal potentials both by wind and by water. The width of samaras in
315 contrast affected the two types of dispersal in opposite ways, allowing differentiation in the
316 dispersal strategies of this invasive species. Furthermore, variation in samara morphology
317 within individuals of *A. altissima* was quite low, supporting the specialization of individuals in
318 different dispersal modes. For instance, it could be suggested that *A. altissima* trees producing
319 narrower samaras would be favoured if they occur close to watercourses, whereas those
320 producing wider samaras would be favoured in open areas or roadsides. To what extent
321 variations in the morphology of samaras and, consequently, their dispersal capabilities could be
322 due to adaptation to different environments could be an interesting topic for future research.

323

324

325 Acknowledgements

326 Funded by the Community of Madrid and the European Social Fund through the program REMEDINAL 3.

327 **Literature**

328 Augspurger, C.K., 1986. Morphology and dispersal potential of wind-dispersed diaspores of neotropical
1 329 trees. American journal of Botany, 353-363.
2

3
4 330 Bartle, K., Moles, A.T., Bonser, S.P., 2013. No evidence for rapid evolution of seed dispersal ability in range
5 331 edge populations of the invasive species *Senecio madagascariensis*. Austral Ecology 38, 915-920.
6

7
8 332 Belsley, D.A., Kuh, E., Welsch, R.E., 2005. Detecting and Assessing Collinearity, Regression Diagnostics. John
9 333 Wiley & Sons, Inc., pp. 85-191.
10

11
12 334 Bory, G., Clair Maczulajtys, D., 1981. Production, dissemination et polymorphisme des semences
13 335 d'*Ailanthus altissima* (Mill.) Swingle, Simarubacees.(Production and different types of seeds in relation
14 336 with dissemination in *Ailanthus altissima* (Mill.) Swingle,(Simarubaceae).). Rev. Gen. Bot 88, 1049-1050.
15
16

17
18 337 Burrows, F., 1986. The aerial motion of seeds, fruits, spores and pollen. Seed dispersal, 1-47.
19

20 338 Cheptou, P.-O., Carrue, O., Rouifed, S., Cantarel, A., 2008. Rapid evolution of seed dispersal in an urban
21 339 environment in the weed *Crepis sancta*. Proceedings of the National Academy of Sciences 105, 3796-3799.
22
23

24 340 Constán-Nava, S., Bonet, A., 2012. Genetic variability modulates the effect of habitat type and
25 341 environmental conditions on early invasion success of *Ailanthus altissima* in Mediterranean ecosystems.
26 342 Biological Invasions 14, 2379-2392.
27
28
29

30 343 Delgado, J.A., Jimenez, M.D., Gomez, A., 2009. Samara size versus dispersal and seedling establishment in
31 344 *Ailanthus altissima* (Miller) Swingle. Journal of environmental biology / Academy of Environmental
32 345 Biology, India 30, 183-186.
33
34

35
36 346 Fleiss, J.L., Cohen, J., 1973. The equivalence of weighted kappa and the intraclass correlation coefficient as
37 347 measures of reliability. Educational and psychological measurement.
38
39

40 348 Galloway, L.F., 2005. Maternal effects provide phenotypic adaptation to local environmental conditions.
41 349 New Phytologist 166, 93-100.
42
43

44 350 Greene, D., Johnson, E., 1989. A model of wind dispersal of winged or plumed seeds. Ecology 70, 339-347.
45

46
47 351 Greene, D., Johnson, E., 1993. Seed mass and dispersal capacity in wind-dispersed diaspores. Oikos, 69-74.
48

49 352 Higgins, S., Nathan, R., Cain, M., 2003. Are long-distance dispersal events in plants usually caused by
50 353 nonstandard means of dispersal? Ecology 84, 1945-1956.
51
52

53 354 Hintze, C., Heydel, F., Hoppe, C., Cunze, S., König, A., Tackenberg, O., 2013. D 3: the dispersal and diaspore
54 355 database–baseline data and statistics on seed dispersal. Perspectives in Plant Ecology, Evolution and
55 356 Systematics 15, 180-192.
56
57
58

59 357 Howe, H.F., Smallwood, J., 1982. Ecology of seed dispersal. Annual review of ecology and systematics, 201-
60 358 228.
61
62
63
64
65

- 359 Kaproth, M.A., McGraw, J.B., 2008. Seed viability and dispersal of the wind-dispersed invasive *Ailanthus*
360 *altissima* in aqueous environments. *Forest science* 54, 490-496.
- 1
2 361 Kowarik, I., Säumel, I., 2007. Biological flora of Central Europe: *Ailanthus altissima* (Mill.) Swingle.
3
4 362 *Perspectives in Plant Ecology, Evolution and Systematics* 8, 207-237.
5
- 6 363 Kowarik, I., Säumel, I., 2008. Water dispersal as an additional pathway to invasions by the primarily wind-
7
8 364 dispersed tree *Ailanthus altissima*. *Plant Ecology* 198, 241-252.
9
- 10 365 Landenberger, R.E., Kota, N.L., McGraw, J.B., 2006. Seed dispersal of the non-native invasive tree *Ailanthus*
11
12 366 *altissima* into contrasting environments. *Plant Ecology* 192, 55-70.
13
- 14 367 Lawrence, J.G., Colwell, A., Sexton, O.J., 1991. The ecological impact of allelopathy in *Ailanthus altissima*
15
16 368 (*Simaroubaceae*). *American journal of Botany* 78, 948-958.
17
- 18 369 Lentink, D., Dickson, W.B., van Leeuwen, J.L., Dickinson, M.H., 2009. Leading-edge vortices elevate lift of
19
20 370 autorotating plant seeds. *Science* 324, 1438-1440.
21
- 22 371 Leung, B., Lodge, D.M., Finnoff, D., Shogren, J.F., Lewis, M.A., Lamberti, G., 2002. An ounce of prevention or a
23
24 372 pound of cure: bioeconomic risk analysis of invasive species. *Proceedings. Biological sciences / The Royal*
25
26 373 *Society* 269, 2407-2413.
27
- 28 374 Lew, R., Doros, G., 2010. Design based on intra-class correlation coefficients. *American Journal of*
29
30 375 *Biostatistics* 1, 1.
31
- 32 376 Matlack, G.R., 1987. Diaspore size, shape, and fall behavior in wind-dispersed plant species. *American*
33
34 377 *Journal of Botany*, 1150-1160.
35
- 36 378 Merriam, R.W., 2003. The abundance, distribution and edge associations of six non-indigenous, harmful
37
38 379 plants across North Carolina. *Journal of the Torrey Botanical Society*, 283-291.
39
- 40 380 Mousseau, T.A., Fox, C.W., 1998. The adaptive significance of maternal effects. *Trends in ecology &*
41
42 381 *evolution* 13, 403-407.
43
- 44 382 Nathan, R., Katul, G.G., Bohrer, G., Kuparinen, A., Soons, M.B., Thompson, S.E., Trakhtenbrot, A., Horn, H.S.,
45
46 383 2011. Mechanistic models of seed dispersal by wind. *Theoretical Ecology* 4, 113-132.
47
- 48 384 Nathan, R., Schurr, F.M., Spiegel, O., Steinitz, O., Trakhtenbrot, A., Tsoar, A., 2008. Mechanisms of long-
49
50 385 distance seed dispersal. *Trends in ecology & evolution* 23, 638-647.
51
- 52 386 Nilsson, C., Brown, R.L., Jansson, R., Merritt, D.M., 2010. The role of hydrochory in structuring riparian and
53
54 387 wetland vegetation. *Biological reviews of the Cambridge Philosophical Society* 85, 837-858.
55
- 56 388 Nombela G, N.A., Bello A, 1994. Structure of the nematofauna in Spanish Mediterranean Continental soils.
57
58 389 *Biol Fertil Soils*, 183-192.
59
60
61
62
63
64
65

- 390 Poschlod, P., Bonn, S., 1998. Changing dispersal processes in the central european landscape since the last
391 ice age: An explanation for the actual decrease of plant species richness in different habitats? *Acta*
1 392 *Botanica Neerlandica* 47, 27-44.
2
- 3
4 393 Säumel, I., Kowarik, I., 2013. Propagule morphology and river characteristics shape secondary water
5 394 dispersal in tree species. *Plant Ecology* 214, 1257-1272.
6
- 7
8 395 Smith, J.R., Bagchi, R., Ellens, J., Kettle, C.J., Burslem, D.F., Maycock, C.R., Khoo, E., Ghazoul, J., 2015.
9 396 Predicting dispersal of auto-gyrating fruit in tropical trees: a case study from the Dipterocarpaceae. *Ecol*
10 *Evol* 5, 1794-1801.
11
- 12
13 398 Soomers, H., Karssenberg, D., Soons, M.B., Verweij, P.A., Verhoeven, J.T.A., Wassen, M.J., 2012. Wind and
14 399 Water Dispersal of Wetland Plants Across Fragmented Landscapes. *Ecosystems* 16, 434-451.
15
- 16
17 400 Soons, M.B., Heil, G.W., Nathan, R., Katul, G.G., 2004. Determinants of long-distance seed dispersal by wind
18 401 in grasslands. *Ecology* 85, 3056-3068.
19
- 20
21 402 Tackenberg, O., 2003. Modeling long-distance dispersal of plant diaspores by wind. *Ecological*
22 403 *Monographs* 73, 173-189.
23
- 24
25 404 Tackenberg, O., Poschlod, P., Bonn, S., 2003. Assessment of wind dispersal potential in plant species.
26 405 *Ecological Monographs* 73, 191-205.
27
- 28
29 406 Thomson, F.J., Moles, A.T., Auld, T.D., Kingsford, R.T., 2011. Seed dispersal distance is more strongly
30 407 correlated with plant height than with seed mass. *Journal of Ecology* 99, 1299-1307.
31
- 32
33 408 Van der Pijl, L., 1982. Principles of seed dispersal in higher plants. *Principles of seed dispersal in higher*
34 409 *plants*.
35
- 36
37 410 Vander Wall, S.B., Kuhn, K.M., Beck, M.J., 2005. Seed removal, seed predation, and secondary dispersal.
38 411 *Ecology* 86, 801-806.
39
- 40
41 412 Venable, D.L., 1985. The evolutionary ecology of seed heteromorphism. *American Naturalist*, 577-595.
42
- 43
44 413 Vittoz, P., Engler, R., 2007. Seed dispersal distances: a typology based on dispersal modes and plant traits.
45 414 *Botanica Helvetica* 117, 109-124.
46
- 47
48 415 Wang, S.-z., Nie, H.-b., Shi, C.-j., 2015. A drifting trajectory prediction model based on object shape and
49 416 stochastic motion features. *Journal of Hydrodynamics, Ser. B* 26, 951-959.
50
- 51
52 417 Wheelwright, N.T., Orians, G.H., 1982. Seed dispersal by animals: contrasts with pollen dispersal, problems
53 418 of terminology, and constraints on coevolution. *American Naturalist*, 402-413.
54
- 55
56 419 Williamson, M.H., Fitter, A., 1996. The characters of successful invaders. *Biological conservation* 78, 163-
57 420 170.
58
- 59
60
61
62
63
64
65

421 Yasuda, K., Azuma, A., 1997. The autorotation boundary in the flight of samaras. Journal of theoretical
422 biology 185, 313-320.

1
2 423
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

Figure 1
[Click here to download high resolution image](#)

