- 1 The original publication is available at <u>www.springerlink.com</u>
- 2 <u>http://dx.doi.org/10.1007/s10682-010-9391-4</u>
- 3 Evolutionary Ecology (2011) 25:155-169

Are the best dispersers the best colonizers? Seed mass, dispersal and establishment in *Carduus* thistles

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16 Running title: Seed mass, dispersal and establishment in Carduus

- 17 **Key words**: colonization; establishment; germination; seed mass; dispersal-establishment
- 18 trade-off; wind dispersal

19 Abstract

20 The negative correlation between dispersal and establishment appears to be well documented 21 in the plant literature; smaller seeds tend to disperse better but germinate less well, and produce smaller seedlings. However, because dispersal capacity is often quantified using 22 proxies, such as the settling velocity of wind-dispersed seeds, little is known about the exact 23 24 shape of this negative correlation, and how it is modified by other plant traits and 25 environmental conditions. We studied the dispersal-establishment correlation in two winddispersed thistles (Carduus nutans and Carduus acanthoides). We applied a mechanistic wind 26 27 dispersal model (WALD) to seeds released under a range of environmental conditions, and 28 tested germination and seedling growth under standardized conditions in a greenhouse. 29 Dispersal distance and establishment (germination and seedling growth) were not 30 significantly correlated, although in both species, smaller seeds dispersed farther, and showed lower germination and lower seedling growth rates. This apparent paradox can partly be 31 32 explained by the significant influence of other factors such as release height and environment 33 (wind and vegetation), which explained more variation in dispersal than did terminal velocity. 34 Another potential explanation is the variation in seed traits: germination is strongly positively 35 related to seed mass, weakly positively related to plume loading, but not significantly related 36 to terminal velocity. This weakening of the correlation with germination is due to additional 37 layers of trait (co)variability: for instance, seed mass and pappus size are positively correlated, 38 and thus big seeds partially compensate for the negative effect of seed mass with larger pappi. 39 Our mechanistic approach can thus lead to a better understanding of both potentially opposing 40 selection pressures on traits like seed mass, and diluting effects of other seed, plant and 41 environmental factors.

42 Introduction

43 Seed size is one critical factor affecting several aspects of the plant colonization process. This
44 trait may have either positive or negative effects on two key early steps in colonization,
45 dispersal and establishment, and thus represents a likely trade-off between different

- 46 colonization stages. In wind-dispersed plants seed size, or more specifically, seed mass, may
- 47 affect dispersal negatively by increasing the seed settling velocity (Greene and Johnson 1993;
- 48 Greene and Quesada 2005). Conversely, it has been demonstrated in many plants (not just 49 wind-dispersed species), that seed mass tends to positively affect germination ability and

1 seedling growth (Turnbull et al. 1999; Turnbull et al. 2004). Thus, by increasing seed mass, 2 wind dispersed plants may trade dispersal success for germination and establishment success. 3 This has been pointed out repeatedly in studies of dispersal and establishment within and 4 among species (Strykstra et al. 1998; Soons and Heil 2002; Debain et al. 2003; Jakobsson and 5 Eriksson 2003). In some cases the relationships may be different, e.g. for animal-dispersed 6 seeds where seed size and dispersal distance is not necessarily negatively related (Coomes 7 and Grubb 2003), and for secondary wind dispersal where the relationship is unimodal 8 (Schurr et al. 2005). This may change or eliminate the negative correlation. Moreover, even 9 when seed mass has significant and opposing effects on dispersal and establishment, they 10 themselves may not be negatively related (Soons and Heil 2002; Debain et al. 2003). This may be due to low variability in commonly used dispersal proxies such as terminal velocity 11 12 (Soons and Heil 2002; Debain et al. 2003), in which case a negative correlation will not be detected using a purely empirical approach. Therefore, to understand and resolve this apparent 13 14 paradox, a mechanistic analysis of the dispersal-establishment relationship with realistic 15 variation affecting dispersal and germination is required. This is an important step towards understanding the implications for evolution of seed size in relation to dispersal and 16 17 establishment.

In this paper, we start to address this potentially fruitful line of inquiry. Specifically, we test the conditions for the existence of negative dispersal-establishment correlations using empirical studies of dispersal, germination and seedling growth for two *Carduus* thistles. Using mechanistic statistical models for the dispersal of the two species, coupled with statistical models of germination and establishment, we address the components of the dispersal-establishment relationship to investigate how the overall putative trade-off is affected by species traits and the environment.

25 Material and methods

26 A conceptual model

Our starting point for this analysis is a conceptual model of how plant traits and environment interact to influence dispersal and establishment (Fig. 1). By "dispersal" we mean the movement of a seed from its mother plant to its destination location. Thus dispersal is only the first step of colonization, which also includes establishment, i.e. germination, survival and plant growth. In this paper we focus on the physical displacement involved in dispersal and on the germination of dispersed seeds and early seedling survival and growth.

33 In the conceptual model several factors appear as "predictor" and "response" variables 34 for different relationships. We carried out a set of empirical studies of dispersal, germination 35 and seedling growth as dependent on seed traits to test these basic relationships and their 36 effects on dispersal and early establishment for a pair of wind-dispersed thistles, Carduus 37 nutans and C. acanthoides. We focus mostly on the biological links in the empirical studies, 38 while the meteorological relationships between environmental variables (wind at 39 measurement height, vegetation height, release height and wind between release height and ground) were modelled as described below. 40

41 The species

42 Our study species are two economically important invasive weeds: *Carduus nutans* L. and C. acanthoides L. (musk and plumeless thistle; Asteraceae). These thistles are originally from 43 44 Eurasia, but have spread to most other continents (Holm et al. 1979). The life-history and 45 demography of the species is well-known from several studies (Desrochers et al. 1988; Shea 46 and Kelly 1998; Shea et al. 2005; Shea et al. 2006; Jongejans et al. 2007b; Jongejans et al. 2008). Both species are primarily found in pastures and disturbed areas and seem to do 47 48 particularly well on fertile soils over limestone (Desrochers et al. 1988). They often establish 49 after disturbance (Jongejans et al. 2007b) Both species are short-lived monocarpic perennials, and colonize exclusively by seed. Carduus nutans has on average slightly larger seeds (3.0-50

4.0 mm long) than *C. acanthoides* (2.5-3.5 mm). *Carduus nutans* also often grows taller than *C. acanthoides* and produces more seeds per flower head, but fewer flower heads per plant
(Desrochers et al. 1988; Skarpaas and Shea 2007). In Central Pennsylvania, flowering of *C. nutans* is usually completed by late June, and the peak dispersal season is in July; for *C. acanthoides* the peak dispersal season is July-August.

6 A mechanistic dispersal model

7 In natural environments, the dispersal distance of a seed depends on environmental factors in 8 addition to seed mass and derived traits such as settling velocity, and hence the dispersal-9 establishment relationship may be modified by the environment. Mechanistic models of wind dispersal typically include variables such as seed release height, horizontal wind speed and 10 11 various aspects of air instability, in addition to terminal velocity (Kuparinen 2006). Many of 12 these models involve complex simulations, but the WALD analytical model of long-distance 13 dispersal has been found to predict dispersal well for a number of wind-dispersed species 14 (Katul et al. 2005), including the Carduus thistles considered here (Skarpaas and Shea 2007). 15 This model is derived from Lagrangian fluid mechanics theory and predicts an inverse Gaussian (Wald) distribution of dispersal distances *r* (Katul et al. 2005, eqn 5b): 16

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$$p(r) = \left(\frac{\lambda}{2\pi r^3}\right)^{\frac{1}{2}} \exp\left(-\frac{\tau(r-\mu)^2}{2\mu^2 r}\right).$$
 (1)

18 The location parameter μ (the mean) and the scale parameter τ are related to species and 19 environment characteristics by

$$20 \qquad \mu = \frac{HU}{F} \tag{2}$$

$$21 \qquad \tau = \left(\frac{H}{\sigma}\right)^2,\tag{3}$$

where *H* is the seed release height, *F* is seed settling velocity, *U* is the hourly mean horizontal wind velocity between *H* and the ground, and σ is a turbulent flow parameter reflecting wind variation due to vegetation structure and weather conditions (Katul et al. 2005). σ is derived from the horizontal wind speed under the assumptions that the dissipation rate can be estimated from surface roughness and friction velocity, and that the friction velocity is proportional to the vertical turbulence (see Skarpaas and Shea 2007, Appendix).

28 According to this model, the expected dispersal distance for a single seed is μ (the 29 mean of the inverse Gaussian distribution; Evans et al. 2000), which is given by the standard 30 ballistic equation (equation 2). The variance depends on both μ and τ (Evans et al. 2000). The probability that a seed reaches a certain distance is given by 1-CDF(r), where CDF(r) is the 31 32 cumulative distribution function of the Wald model, i.e. the integral of the probability density 33 function (equation 1). The distribution of dispersal distances of an entire population of seeds (the dispersal kernel) also depends on the variability of the parameters underlying μ and τ 34 35 during the release period. To account for variability in terminal velocity, release height and 36 wind conditions for extended periods of time (more than one hour), the WALD model can be 37 integrated over these variables to obtain the dispersal kernel k(r) (Skarpaas and Shea 2007):

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$$k(r) = \iiint p(H)p(U)p(F)p(r)dHdUdF$$

39 where p(H), p(U) and p(F) are the probability densities of H, U and F, respectively, and p(r)40 is the WALD model (equation 3). We used this integrated model to estimate distributions of

41 dispersal distances on the basis of independent measurements of environment (wind,

42 vegetation height) and plant traits (seed release height, terminal velocity) in the experiments

43 described below.

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1 Dispersal experiments

2 Eight dispersal experiments were carried out in Central Pennsylvania during the dispersal 3 season (July-October) in 2003 and 2004 (Table 1). Seeds from the dispersal experiments in 4 2004 were used for seed trait and germination studies (see below). As the experiments 5 involved the release of seeds of invasive species, the experiments were carried out at sites 6 where the species already occurred (Allen and Shea 2006). Seeds were drawn from pools of 7 seeds collected from several plants (> 5) in two populations of each species near the study 8 sites. C. nutans seeds were collected in the Elliotsburg area, Perry County, PA; C. 9 acanthoides seeds were collected in the State College area, Centre County, PA. For each 10 species we selected two sites with different vegetation heights corresponding to the natural 11 habitats of the species: one site with low vegetation (pasture) and one with high vegetation 12 (wasteland). Maximum vegetation heights (flowering stalks) were 0.54 \pm 0.25 m and 1.27 \pm 13 0.49 m at the low and high vegetation sites respectively (mean \pm SD of measurements at 20 14 random points at each site). At each site we carried out two dispersal experiments to cover a 15 range of wind speeds (Table 1). The experiments were all carried out over a period of 2-4 16 hours in the late morning or early afternoon on sunny days.

17 We measured field wind speed and direction using an automated mobile weather 18 station with an anemometer mounted at 2 m above the ground. In the two experiments carried 19 out in 2003, horizontal wind speeds were measured using a Young cup anemometer. For the 20 six experiments in 2004 we used a 3D ultrasonic anemometer (CSAT3, Campbell Scientific, 21 Utah, USA) to measure wind speeds in three dimensions (including updrafts). Horizontal 22 wind speeds for 2004 were calculated as the vector sum of the two horizontal components. In 23 both years, wind speeds were stored every 10 s using a CR-10X datalogger (Campbell 24 Scientific). Wind speed and turbulence between the release height and the ground was 25 estimated using the logarithmic wind profile and a mechanistically based turbulence approximation involving horizontal wind speed and standard meteorological parameters (the 26 27 von Karman and Kolmogorov constants and surface roughness parameters), as described in 28 the appendix of Skarpaas and Shea (2007).

In each experiment we released 40-150 dispersal units (filled achenes with intact 29 30 pappi) one at a time. The dispersal units were released at random heights in the range 0.30 -2.30 m (corresponding roughly to flower head heights on the plants), pre-determined by a 31 random number generator. Each dispersal unit was released by hand at the beginning of a 10-32 33 second recording interval. As soon as the procedure was completed for one dispersal unit, we 34 repeated the whole procedure with the next dispersal unit. We followed each dispersal unit 35 and measured the distance from the release point to the point where it first landed, or, in a few 36 cases, was lost from sight. Although we recorded the last distance at which the lost dispersal 37 units were observed, they were not included in any analyses in this study, because 38 measurements of traits and establishment were carried out on seeds collected from the 39 dispersal experiments.

40 Seed traits

41 The dispersal units from the dispersal field experiments carried out in 2004 were collected 42 and stored individually in vials to preserve the pappi for measurements in the lab. The 43 maximum diameter of the pappus, and the maximum width and length of the achene ('seed' 44 hereafter) were measured using a digital micrometer. Settling velocity (terminal velocity) was 45 measured by dropping each intact dispersal unit through an air-tight tube and timing the 46 descent through 1.16 m using a stopwatch. To minimize bias from acceleration, the dispersal 47 units were allowed to accelerate for 7 cm. To minimize measurement error, five replicate 48 measurements for each dispersal unit were averaged. Finally, the mass of each seed with the 49 pappus removed was measured using an electronic balance.

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Germination and seedling establishment 1

2 After lab measurements were completed for all seeds, a seed germination trial was conducted 3 between October 20 and November 28, 2004, using all the seeds from dispersal studies 2, 4, 6 4 and 7 (Table 1): 110 seeds of Carduus nutans; 170 of C. acanthoides; 280 in total. Each seed 5 was randomly assigned a position on one of six trays in the greenhouse. The seeds were 6 planted in bare soil, 0.5 cm below the surface, which has been shown to be optimal for these 7 species (McCarty et al. 1969). The six trays were placed in a greenhouse with a temperature 8 range of 18-27°C, artificial light 12 hrs/per day (7:00am—7:00pm) and 30-80% air humidity. 9 The trays were watered regularly. The planted seeds were checked daily and the date of each 10 germination was recorded, as well as the survival of germinated seedlings. Once a week (from 11 November 7 until November 28, 2004), we recorded the maximum diameter of the rosette and 12 the length of the longest leaf for each seedling. Analysis 14

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We tested the relationships among species characteristics (seed mass, pappus diameter, plume loading, release height, settling velocity), environmental variables (vegetation height and wind speeds) and dispersal and establishment (germination and early plant growth) using generalized linear regression models (GLM). In the figures we present results for the four studies where we have data on dispersal, seed traits and establishment (studies 2, 4, 6 and 7 in Table 1), but include additional analyses in the text for the full dispersal data set (all studies in Table 1) where only dispersal-related variables are involved. Settling velocity F has been found to be linearly related to the square root of plume loading L (Andersen 1993; Meyer and

21 22 Carlson 2001):

$$23 \qquad F = \alpha \sqrt{L} \,.$$

 α is a constant and the plume loading L is given by (Greene and Johnson 1990): 24

$$25 \qquad L = \frac{M}{A},\tag{6}$$

26 where M is seed mass (mg) and A is pappus area (mm²) assuming a circular plume (Matlack 27 1987). On a logarithmic scale, dispersal distance, seed settling velocity and seed mass should 28 be linearly related (eqs. 2, 5 and 6). Thus, in the statistical analysis these variables were log 29 transformed. Plume loading was square root transformed when used as a predictor for settling 30 velocity (eq. 5). For the WALD model parameters we obtained independent estimates (not fitted to dispersal distances). The relationship between germination and seed mass were 31 32 analyzed using logistic regression. All analyses were carried out in R (R Development Core 33 Team 2009).

Results 34

35 Large (heavy) seeds had bigger pappi in both species (Fig. 2A; linear regression, P < 0.001). However, an increase in pappus area was not sufficient to compensate for the increased mass 36 37 of the seed: plume loading increased linearly with seed mass in both species (Fig. 2B; linear 38 regression, P < 0.001). Carduus nutans had significantly heavier seeds (Table 2), larger pappi 39 than C. acanthoides (Fig. 2A, linear regression, P < 0.001), but lower plume loadings (Fig. 40 2B, linear regression, P < 0.001). Terminal velocity, the main dispersal trait of the achene-41 pappus unit, increased linearly with the square root of plume loading in both species (Fig. 2C; 42 linear regression, P < 0.001). Terminal velocity was lower in C. nutans, but not significantly 43 so (Table 2).

44 Dispersal distances were strongly affected by plant traits and environmental factors, 45 but there was no significant difference between the species (Fig 3; linear regression, P >0.195). Seeds with higher terminal velocities dispersed shorter distances (Fig. 3A; linear 46 47 regression, P < 0.001). Seed release height (plant height), on the other hand, affected dispersal 48 distances positively (Fig. 3B; linear regression, P < 0.001). This was also true for the full

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dispersal data set including all 8 experiments listed in Table 1 (linear regression, P < 0.001). As expected, dispersal distances also increased with the ambient wind speed (Fig 3C, linear regression, P < 0.001; full dispersal data set: P < 0.001). Of the factors that explained a significant proportion of the variation in dispersal distance (ANOVA, P < 0.05; for the seeds where all measurements were available) wind speed explained the most (45%), followed by terminal velocity (16%), vegetation height (7%) and seed release height (2%).

7 The wind speed experienced by each individual dispersal unit varied considerably 8 between dispersal experiments in high and low vegetation (see also Marchetto et al. 2010b). 9 The resulting distributions of dispersal distances were significantly skewed towards shorter 10 distances in high vegetation for both species (Fig. 4; Kolmogorov-Smirnov one-sided tests, P < 0.001). The WALD model predicted the dispersal distance distributions extremely well for 11 12 Carduus nutans in low vegetation, where there was no significant difference among the observed and predicted distributions (Kolmogorov-Smirnov two-sided test, P = 0.653). In the 13 14 three other cases the model predictions deviated slightly from the observations at intermediate 15 distances (Kolmogorov-Smirnov two-sided tests, P < 0.034), but the model nevertheless predicted the distributions well in both low and high vegetation considering that predictions 16 17 were made on the basis of independent measurements of the predictors.

18 While seed mass affected dispersal negatively (Fig 5A), it affected establishment positively. The probability of germination increased significantly with seed mass (Fig. 5B; 19 20 logistic regression, P < 0.001). Of the 110 Carduus nutans seeds 15 germinated (13.6%), while 25 of the 170 C. acanthoides seeds germinated (14.7%). There was no significant 21 22 difference between the species in germination (logistic regression, P = 0.164). Although some large seeds germinated earlier than small seeds, seed mass did not significantly affect 23 germination time in any of the species (linear regression, P > 0.100). Nor did it affect seedling 24 25 survival; all seedlings survived until the end of the experiment. However, seed mass did affect 26 early seedling growth in both species positively, both in terms of leaf growth (Fig. 5C) and rosette growth (data not shown, but similar pattern as leaf growth; linear regression, P < P27 28 0.001). By both growth measures, C. acanthoides seedlings grew significantly faster than C. 29 *nutans* seedlings for comparable seed masses (Fig. 5C; linear regression, P < 0.001).

30 Despite the fact that seed mass affected dispersal negatively and establishment 31 positively, there was no statistically significant negative relationship between dispersal and establishment in our data. The probability of germination was predicted well by seed mass 32 33 (see above), but increasingly worse by plume loading (logistic regression coefficient c =34 0.138, P = 0.058), terminal velocity (c = 3.07, P = 0.215) and dispersal distance (log r; c = -35 0.041, P = 0.739; Fig. 6A). Controlling for wind speed (log U) and release height (log H) made the relationship between dispersal and germination stronger but still not statistically 36 37 significant (logistic regression coefficient c = -0.236, P = 0.191). Similarly, seedling leaf and 38 rosette growth were not significantly related to the observed dispersal distance (linear 39 regression coefficients 0.016 and 0.004 respectively; P = 0.851 and 0.933; Fig. 6B and C).

40 Even though dispersal distance and germination were not significantly negatively 41 related, there may still be opposing selection pressures on seed mass since seed mass affected 42 dispersal distance negatively and establishment positively. Under the assumption that these 43 separate and opposing effects could cause a negative correlation between the ability to arrive and the ability to establish in a new location, we explored the predicted optimal seed mass in 44 45 either species (Fig. 7). Note that this hypothetical picture assumes that there is no interaction effect of seed mass and vegetation height on the probability of establishment (i.e. that small 46 and large seeds are affected similarly by vegetation height). The predicted seed mass, where 47 effects on dispersal and germination are optimally balanced, is strongly modified by 48 49 environmental conditions. For example, the optimal seed mass for colonizing safe sites more 50 than 2 m away from the mother plant is drastically reduced by high vegetation, which 1 effectively lowers the wind speed experienced by the seed. The reduction in the optimal seed

2 mass is stronger for *C. acanthoides* than for *C. nutans* under comparable conditions.

3 **Discussion**

4 In a direct comparison of dispersal and germination we found no statistical support for a 5 negative relationship between dispersal and establishment in our data. However, at the same 6 time our results suggest evolutionarily relevant opposing effects of seed mass on dispersal and establishment for both Carduus nutans and C. acanthoides. This has also been demonstrated 7 8 for many other species (Strykstra et al. 1998; Soons and Heil 2002; Debain et al. 2003; 9 Jakobsson and Eriksson 2003). This apparent paradox, which has also been observed in some 10 other studies (e.g. Soons and Heil 2002; Debain et al. 2003), can be explained by low variability in commonly used dispersal proxies such as terminal velocity. Our study also 11 raises other likely explanations. One possible explanation is the significant influence of other 12 13 factors such as release height and environment (wind and vegetation) on dispersal. In our 14 study these factors explained more of the variability in dispersal distance than did terminal velocity. Similarly, strong effects of the environment were reported by Nathan et al. (2002) 15 16 who found that relatively heavy seeds were uplifted above the canopy (i.e. likely to be longdistance dispersed). Another possible explanation for the lack of a clear negative relationship 17 18 between dispersal and establishment is the variation in seed traits and their measurement: 19 germination is strongly positively related to seed mass, less clearly but still positively related 20 to plume loading, but not significantly related to terminal velocity and dispersal distance. This 21 dilution of effects of seed traits on germination is partly because additional layers of 22 variability in traits and measurements reduce our ability to see a relationship with a limited 23 sample size, and partly because of trait co-variability. For instance, seed mass is positively 24 correlated with pappus size. As pointed out by e.g. Sakai et al. (1998), heavy seeds are not 25 poor colonizers if the weight is compensated for by bigger dispersal structures such as wings 26 or plumes. For the Carduus thistles tested here, heavier seeds had larger pappi, but not sufficiently large to compensate for the increase in plume loading and terminal velocity. 27

28 In our analyses, all the relationships in the pathways from seed mass to dispersal and 29 to establishment are statistically significant and the seed mass-mediated effects are sufficiently large to cause more than five-fold differences in the probability of dispersal 30 31 beyond the mean dispersal distance (approximately 2 m) and in the probability of germination 32 (Fig. 7). This suggests that there is a mechanistic basis for a negative dispersal-establishment 33 relationship for both *Carduus* species; the best dispersers may not be the best colonizers, even though the direct correlation between dispersal and establishment appears to be overwhelmed 34 35 by variability in environmental factors in the particular context of our studies. In more constant environments (predictable winds, small range of release heights, etc.), seed mass-36 37 mediated effects will likely be relatively more important.

38 The mechanisms involved, and the shape of the dispersal-establishment relationship, 39 seem to be somewhat different in the two species. C. acanthoides had higher terminal 40 velocities than C. nutans for any given seed mass. This is mostly due to higher plume 41 loadings in C. acanthoides, but other factors, such as lower plume density may also affect 42 terminal velocity (Greene and Johnson 1990; Marchetto et al. 2010a). Regardless of the 43 underlying cause, the predicted optimal seed mass for colonization was larger for C. nutans than for C. acanthoides in similar environments (Fig. 7). Interestingly, observed seed mass is 44 45 even smaller than predicted by our models for both species in low vegetation and for C. 46 nutans in high vegetation, which suggests that the ruderal existence and long-distance 47 dispersal may be very important for these species.

The environment strongly influenced the dispersal-establishment relationship for both
 species in our study. As expected, wind had a strong positive effect on dispersal distances.
 Vegetation height reduces wind speed at dispersal height, and thereby reduces dispersal

1 distances. Seeds were also sometimes intercepted by vegetation in the high-vegetation 2 experiments (low outliers in Fig 3). For both species the loss of dispersal capacity is dramatic 3 in high vegetation / low wind, which in turn affects the hypothesized optimal seed mass for 4 colonization (Fig 7). This general conclusion can be (and has been) drawn on the basis of seed 5 characteristics only (terminal velocity and germination), as the expected dispersal distance in 6 the common ballistic equation (equation 2; also the mean of the WALD model) is inversely 7 related to terminal velocity. However, estimating the exact shape of the relationship under 8 varying environmental conditions requires a probability distribution of dispersal distances, 9 provided by the mechanistic dispersal model (WALD) in this study. Using this model we can 10 estimate the shape of the dispersal-establishment relationship for any of the seed traits and environmental variables included in the model. 11

12 This being said, one must keep in mind that any model is a simplification of nature. This goes for the WALD model as well as for the conceptual model (Fig. 1) in this study. 13 14 Although the WALD model has proved to be a good model for several species at the scales at 15 which we can measure (Katul et al. 2005; Skarpaas and Shea 2007), it may not be able to represent extreme long-distance dispersal (Katul et al. 2005). This will affect our ability to 16 17 represent the dispersal-establishment relationship correctly if suitable habitat and safe sites 18 are far apart. In nature, there are also additional possible connections among the variables that 19 we ignore in our analysis because of the experimental design. For instance, in our dispersal 20 experiments seeds were released at random with respect to wind speeds, but natural seed 21 release is strongly affected by wind, humidity and other environmental conditions (Skarpaas 22 et al. 2006; Jongejans et al. 2007a). Moreover, vegetation height and seed release height (a function of plant height) are probably positively correlated in many natural settings (except 23 for e.g. pastures in which cattle graze around thistles), but in our experiments seeds were 24 25 released at random heights independently of vegetation height. Finally, vegetation structure, 26 including vegetation height, will also most likely affect germination and seedling survival and growth through its effect on light and water availability. In our experiments, germination and 27 28 seedling growth tests were carried out under standardized favourable conditions in a 29 greenhouse. This allows the direct analysis of the dispersal-establishment relationship on the 30 same seeds, while also increasing the statistical power in comparison to realistic conditions 31 (where a worst-case scenario would be no germination at all). This means that our exploration 32 of consequences for the dispersal-establishment relationship does not include effects of 33 vegetation on germination and seedling growth. There is some evidence that establishment 34 and seedling growth are reduced in dense and tall vegetation, suggesting better recruitment in 35 open microsites (Jongejans et al. 2007; Shea unpublished data), but we do not know how this relates to seed size. If tall vegetation reduces germination in all seeds, the optimum in Figure 36 37 7 would be the same, but if small seeds are affected disproportionately (see e.g. Coomes and 38 Grubb 2003), the optimum seed mass would be higher.

39 The evolutionary outcome in any particular case may deviate from the relationship 40 depicted in Figure 7 for a variety of reasons. Even if there is a negative correlation between 41 dispersal and establishment, this is not sufficient to imply causality or the existence of a trade-42 off (Knops et al. 2007). We did not control for genetic effects (and have no information about seed masses of parents), so here we are dealing with correlations of phenotypic traits for 43 44 which the genetic basis is unknown. As far as we know, the heritability of seed size in 45 Carduus has not been investigated, but studies of a range of other plant species suggest that seed size may be affected by parental genes as well as parental environments (e.g. Stratton 46 1989; Mazer and Wolfe 1992; Wheelwright 1993). Furthermore, the seed mass predictions in 47 this study are made on the level of individual seeds and not on the level of mother plants: the 48 49 fitness of mother plants, i.e. the relative number of expected offspring over a lifetime, also 50 depends on other trade-offs such as growth vs. reproduction (Metcalf et al. 2008) and seed

1 number vs. seed size (Jakobsson and Eriksson 2003), in addition to environmental factors 2 such as post-dispersal seed predation (Jongejans et al. in prep.) and the availability of safe 3 sites (Geritz et al. 1984; Bullock et al. 1995; Bullock 2000). The panels in Figure 7 apply 4 within homogeneous patches of high or low vegetation. Widely dispersed seeds may 5 encounter different conditions for establishment: if they are better than in the source patch 6 (e.g. less competitive), any negative dispersal-establishment relationship may be cancelled 7 out; if they are worse, it may be even stronger. Moreover, the dispersal-establishment 8 correlation may also be modified by processes at later stages in the life cycle: for instance, the 9 disadvantage of *C. acanthoides* in the initial colonization by seed (dispersal and germination) 10 may be countered by faster seedling growth (Fig 5C).

Thus a number of factors, both species-specific and environmental in origin, will 11 12 introduce variability that will affect the evolutionary outcome in any particular case. Simplifications are necessary to separate the essence from the noise, and a mechanistic 13 14 modelling approach has the potential to include, and integrate over, relevant sources of 15 variability. A complete mechanistic analysis of all of the relevant factors is beyond the scope of this paper, but we consider this a useful first step. A mechanistic approach will help to 16 17 understand the seed mass-dispersal-establishment paradox, and also to study more complex 18 relationships than just positive, negative or non-existent correlations between dispersal and 19 establishment. This makes it possible to go beyond simple detection and description of the 20 dispersal-establishment relationship and has allowed us to explore its mechanistic dependence on species traits and environmental conditions in detail. Such a mechanistic approach will 21 22 lead to a better understanding and quantification of potentially opposing selection pressures 23 on traits like seed mass, as well as of the diluting effects of other seed, plant and 24 environmental factors.

25 Acknowledgements

We are grateful to Jeff Buterbaugh, Paul Chen, Matt Clark, Wilmer Garman, Brian Jones, Pete LeVan, Melanie Northrup, Emily Leichtman, Emily Rauschert, Christina Saylor, Zeynep Sezen and Laura Warg for assistance in field and lab studies, and to two anonymous reviewers for valuable comments. This work was funded by the National Science Foundation (grant no. DEB-0315860 and DEB-0614065 to KS), the Norwegian Research Council (grant no. 161484/V10 to OS) and the Netherlands Organization for Scientific Research (NWO veni-grant 863.08.006 to EJ).

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- 30

Figures 1



2 3 Figure 1. Conceptual model of modeled and hypothetical relationships among plant traits, 4 environment, dispersal and establishment. Solid lines indicate positive relationships; dashed 5 lines indicate negative relationships. Gray boxes refer to figures illustrating results for each 6 connection, except for the meteorological relationships, which are not tested in this study, but 7 modelled using standard methods (see Methods). 'Ambient wind speed' is the horizontal wind 8 speed at measurement height (2 m); 'Wind speed' is the wind speed as perceived by the 9 dispersal unit, estimated as the average horizontal wind speed between release height and the 10 ground. As measures of 'Establishment' we use germination or seedling growth, as specified 11 for each analysis.

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Figure 2. Relationships among dispersal-related seed traits in *Carduus*. Observations and fitted regression models for *Carduus nutans* (filled circles; solid lines) and *C. acanthoides* (open circles; dashed lines). In C the single regression line applies to both species; they were not significantly different. R^2 values: (A) 0.83, (B) 0.96, (C) 0.96.



9 Figure 3. Relationships among *Carduus* seed traits, environmental conditions and dispersal.

10 Observations for Carduus nutans (filled circles) and C. acanthoides (open circles) and

- regression models for both species (lines). R^2 values: (A) 0.18, (B) 0.38, (C) 0.13.

1 2 3 4

Figure 4. Observed (bars) and predicted distributions of dispersal distances using the integrated WALD model (lines) for Carduus nutans and C. acanthoides in low and high vegetation (a few observations > 12 m are omitted for clarity; see Fig. 3). Mean wind speeds 5 6 are given for each study at measurement height (2 m; U₂) and between release height and ground (U_h).

1 2 Figure 5. Effects of seed mass on (A) dispersal, (B) germination and (C) leaf growth in 3 Carduus nutans (filled symbols) and C. acanthoides (open symbols). In A, observations from 4 low-vegetation studies are indicated by circles and high-vegetation studies by triangles. B 5 shows the observed (bars) and predicted (line) probability of germination as a function of seed 6 mass (logistic regression) for C. nutans (black bars) and C. acanthoides (gray bars). C shows observed and predicted seedling growth as a function of seed mass (linear regression through 8 the origin), measured as the average daily growth of the longest leaf (left) for C. nutans (filled 9 circles, solid lines) and C. acanthoides (open circles, dashed lines).

7

 $\begin{array}{c} 11 \\ 12 \end{array}$ Figure 6. Relationships between observed dispersal and (A) germination, (B) leaf growth and

13 (C) rosette growth in Carduus nutans (filled symbols, black bars) and C. acanthoides (open symbols, gray bars). In B and C observations from low-vegetation studies are indicated by 14 15 circles and high-vegetation studies by triangles.

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1 2 3 Figure 7. Modelled relationship between dispersal and establishment (germination and seedling survival), and the effect of the environment, assuming homogeneous effects of 4 vegetation height on establishment across seeds of all sizes. Each panel shows the probability 5 of dispersing > 2 m (dashed line, predicted by the WALD model; Equation 1), the probability 6 of establishment (dotted line, predicted by the logistic regression model; Fig 5) and the 7 resulting probability of colonization > 2 m from the mother plant (solid line; product of 8 dispersal and establishment probabilities) as a function of seed mass. Means (gray vertical 9 line) and ranges (95% confidence intervals; gray shading) of observed seed mass are indicated 10 in each case. Note that the evolutionary optimal seed mass may also depend on several other 11 factors (see the text for details).

Table 1. Dispersal experiments for Carduus nutans and C. acanthoides at sites with low and high vegetation (pastures and wastelands respectively) in Central Pennsylvania 2003 and 2004. Number of seeds released (N) and horizontal (U_z) and vertical (W) wind speeds (means, SD in parentheses) at 2 m above ground. Seeds from experiment 2, 4, 6 and 7 were used for seed trait and germination trials.

5								
6		Date	Site	Species	Vegetation	N	$U_2 ({\rm ms}^{-1})$	$W_2 ({\rm ms}^{-1})$
7	1	29 July 2003	Garman farm, Elliotsburg	C. nutans	low	150	1.00 (0.69)	_*
8	2	23 July 2004	Garman farm, Elliotsburg	C. nutans	low	70	1.33 (0.72)	0.01 (0.10)
9	3	1 Sep. 2004	Allen & Commerce, Carlisle	C. nutans	high	70	1.07 (0.51)	-0.10 (0.13)
10	4	22 Sep. 2004	Allen & Commerce, Carlisle	C. nutans	high	40	0.97 (0.50)	-0.10 (0.20)
11	5	16 Sep. 2003	Haller farm, State College	C. acanthoides	low	150	2.99 (0.83)	_*
12	6	17 Aug. 2004	Haller farm, State College	C. acanthoides	low	100	1.66 (0.67)	-0.10 (0.12)
13	7	17 Aug. 2004	Big Hollow, State College	C. acanthoides	high	70	1.03 (0.56)	-0.08 (0.20)
14	8	4 Oct. 2004	Big Hollow, State College	C. acanthoides	high	54	1.21 (0.68)	-0.07 (0.30)
15					-			

* Vertical wind speeds were not measured in 2003.

Table 2. Traits of the achene-pappus dispersal unit (mean and SD) for Carduus nutans and C. acanthoides, with sample sizes (N) and P-values for a two-sided t-test for the difference between them.

21		C. nutai	C. acanthe	C. acanthoides		
22	Trait	Mean (SD)	N	Mean (SD)	N	Р
23	Seed mass (mg)	2.1 (1.0)	96	1.7 (0.6)	142	0.001
24	Plume diameter (mm)	29.6 (4.7)	107	22.9 (2.7)	163	< 0.001
25	Plume area (mm ²)	706.9 (212.2)	107	419.3 (96.5)	163	< 0.001
26	Plume loading (g mm ⁻²)	3.26 (2.27)	96	4.27 (1.95)	139	< 0.001
27	Terminal velocity (m s ⁻¹)	0.35 (0.11)	68	0.39 (0.12)	25	0.184