Dung beetles as secondary seed dispersers in a temperate grassland

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Abstract

The two-phase dispersal event in which dung beetles move seeds after endozoochory is often assumed to be advantageous for plant regeneration. Because seeds are expected to end up in favourable and safe germination sites, it is considered as an example of directed dispersal. However, literature so far is restricted to tropical rain forest ecosystems, while data for temperate regions are lacking. In this study, the effect of dung beetles on seedling establishment of endozoochorically dispersed seeds is evaluated for a temperate grassland ecosystem. We performed a field experiment in which cages excluded dung beetles from horse and cattle dung samples with mixed-in grass seeds. Seed germination from these samples was significantly higher than that from samples which were accessible to dung beetles. This indicates that the effect of dung beetles on short-term seedling establishment was negative, which contrasts with the patterns found for large-seeded species used in tropical studies. This is most likely attributed to the lack of roller species and the larger depth at which tunneling \textit{Geotrupes} species bury seeds.

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Introduction

Acting as dispersal agents, animals may take plant seeds to non-random microhabitats that are well-suited for establishment and growth. This process, referred to as ‘directed dispersal’, was distinguished by Howe and Smallwood (1982) as one of three major advantages of seed dispersal, and is assumed to be more common than formerly thought (Wenny, 2001). Directed dispersal is considered to be a key step in diplochory (Vander Wall & Longland, 2004), in which seeds are dispersed in two successive phases, each involving a different dispersal agent. According to Vander Wall and Longland (2004), the benefits offered by the different phases differ: second phase dispersers (e.g. ants, rodents, corvids) take seeds to microsites that are more suitable for establishment than those reached by phase one dispersers (e.g. wind, herbivores). One particular case of diplochory occurs when dung beetles act as secondary dispersers of seeds contained in vertebrate dung.

Within the dung beetles (coprophilous species within the Scarabaeidae subfamilies Aphodiinae, Geotrupinae and Scarabaeinae [classification following Janssens, 1949, 1960]), three functional groups are distinguished: dwellers, tunnelers and rollers (Cambefort & Hanski, 1991). Aphodiinae are predominantly dwellers: these small-sized beetles (generally less than 10 mm in length) eat their way through the dung and deposit their eggs without constructing a nest or chamber. Geotrupinae and most Scarabaeinae are tunnelers: they dig a more or less vertical tunnel beneath a dung pat and move the dung to the shaft base. Many Scarabaeinae, specified as rollers, construct a dung ball and transport this over a distance prior to burying it (telecoprid nidification sensu Bornemissza, 1969). Dung beetles do not eat seeds (Andersen & Feer, 2005), so tunnelers and rollers may contribute to directed dispersal by moving and burying seeds along with the dung.

So far, studies on secondary seed dispersal by dung beetles and its influence on plant regeneration almost exclusively focused on tropical rain forest ecosystems. In a review, Andersen and Feer (2005) concluded that burial has a positive impact through lower seed predation and a negative impact through decreased seedling emergence. Little is known, however, about the net outcome of these antagonistic effects. Only Andersen and Levey (2004) really followed the fate of seeds placed in dung until seedling establishment in a (Central Amazonian) rain forest ecosystem. Their results, concerning 11 large-seeded tree species and howler monkey dung, indicated that seeds buried by dung beetles were on average twice as likely to become seedlings than unburied seeds. In the only study investigating dung beetle mediated seed dispersal outside tropical regions we know about, Wicklow, Kumar, and Lloyd (1984) found a subtropical roller species (Kohlmann, 1991) to facilitate seedling establishment of a prairie grass species in North America.

More studies are needed to ascertain if this positive effect also holds for other climate regions and ecosystems (Andersen & Feer, 2005), because it is reasonable to assume that important differences in the dung beetle communities or the plant species involved may lead to deviant patterns. For instance, at lower latitudes dung beetle communities are dominated by rolling or tunneling Scarabaeinae, whereas the northern temperate regions are characterized by a dominance of the dwelling genus Aphodius (Aphodiinae), normally accompanied by one or a few species of tunneling Geotrupinae (Hanski, 1991).

In this study, we aim to test the hypothesis that dung beetle activity has an overall positive effect on successful germination of seeds present in dung in a temperate coastal dune ecosystem. A field experiment was constructed, in which the effect of dung beetle presence on short-term seedling establishment from horse and cattle dung was tested. This ‘main experiment’ was supplemented by some supplementary experiments, i.e. controls to evaluate the main setup in detail and a study of the dung beetle fauna present within.

Materials and methods

Study area and local dung beetle assemblage

The field experiment was set up in the nature reserve ‘Westhoek’ (Belgium, 51°04’50”N–2°34’19”E), consisting of over 340 ha of coastal dune landscape. It is characterized by a spatially heterogeneous mosaic of open dunes, grey dunes, grassland, shrub and woodland. Since 1996, part of the area is grazed by cattle and several equine breeds, restoring extensive grazing previously maintained by livestock. In a recent study of the associated dung beetle fauna, 15 species were found (Struyve, 2002), 12 of which belong to the dweller genus Aphodius Illiger. Onthophagus similis Scriba, Geotrupes niger Marsham and Geotrupes spiniger Marsham were the tunneler species found. No roller species were present in the area. Hence, functionally, the overall dung beetle fauna can be considered as typical of a temperate ecosystem (Hanski, 1991).

Main experiment

At each of two grassland sites within the study area, the effect of dung beetle activity on seed germination was assessed using the same experimental randomized block design. The sites consisted of preliminary mown grasslands located within livestock-excluding fences, the
surroundings of which were grazed by cattle as well as ponies.

One site is distinctly moister than the other: from September 2005 to April 2006, ground water fluctuated between 0.2 and 1.0 m beneath the surface at the moist site and between 1.5 and 2.1 m at the dry site (De Becker, Van Daele, Huybrechts, Provoost, & Leten, unpublished).

Perpendicular to the prevailing moisture gradient at each site, six strips of 1 m width were delimited, which contained (among others [i.e. controls, see below]) eight sod-cut plots (50 × 50 cm, 25 cm inter-plot spacing), on which treated dung samples were deposited, the treatments being allocated at random within a strip (block).

Dung was collected from horses (Konik) and cattle (Galloway) that were fed a seed-free diet in captivity. In September 2005, two-litre samples were deposited on the field plots, and simultaneously, 100 seeds, either from *Agrostis capillaris* L. or *Poa pratensis* L., were mixed into each sample. Dispersules were on average 1.60 mm in length for *A. capillaris* and 2.78 mm for *P. pratensis* (n = 20 each, seeds plus glumes). Cosyns, Claerbout, Lamoot, and Hoffmann (2005) found both grasses to be viably present in dung of cattle, horses and ponies in the ‘Westhoek’ reserve. Period of seed set is August to October for the former, and August for the latter (Grime, Hodgson, & Hunt, 1988).

After deposition in the field, half of the dung samples were caged (‘closed cage’), making the dung inaccessible to dung beetles (1.25 mm mesh), while the others were equipped with an ‘open cage’, covering the top and the southern side of the plot and leaving all other lateral sides open, enabling dung beetles to gain access to the samples easily.

During September and October 2005 – the months in which dung samples were subject to dung beetle colonization – precipitation rates were normal, but temperatures were exceptionally high (in September, mean temperature was 16.0 °C instead of 14.2 °C; October: 14.4 °C instead of 10.6 °C; recorded at ± 20 km off: Malcorps, 2005a, b).

About 73 days after deposition, in November–December 2005, the number of seedlings of either *A. capillaris* or *P. pratensis* was counted on each dung pat once. Seedlings were removed after counting. An additional count was conducted in the following spring (April 2006), but very few seedlings could be added. Visual traces of dung beetle activity were recorded, in particular, tunneling by *Geotrupes*, which was recognized by the presence of tunnel entrances surrounded by bare sand.

**Germination controls**

To test whether the obtained dung was free from seeds, a greenhouse control experiment was performed in which germination from six untreated dung samples per herbivore type (2 L) was monitored. These were deposited on a sand–peat substrate mixture, subjected to a 16-h light:8-h dark regime, and watered manually (September–December 2005).

In order to test the used grass seeds for their viability, we determined germination of 6 times 100 seeds per species in two controls: once in the field, once in the greenhouse. In both, seeds were sown in sand–peat substrate (which, in the field, was spread out within cageless sod-cut plots; September–December).

Moreover, the effect of both ‘open’ and ‘closed cage’ constructions on germination conditions was tested in the greenhouse by sowing seeds in the same substrate, with six replicates for every combination of cage type and plant species (September–December).

**Dung beetle fauna**

To assess the species composition of the dung beetle community in the study area, six horse dung and six cattle dung samples were deposited at both sites and contiguously encircled by five pitfall traps per sample (65 mm diameter, ± 80 mm spacing between). Half of these samples were fitted with ‘open cages’ while the other remained cageless, in order to determine whether the dung beetle fauna visiting the ‘open cages’ was representative of the cageless, i.e. natural situation.

During a 63-day period (mid-September to mid-November 2005), all Scarabaeidae were gathered, identified up to the species level (Janssens, 1960) and classified as tunneler (*Geotrupes* spp., *Onthophagus* spp.) or dwellers (*Aphodius* spp.).

**Data analysis**

In the main experiment, Dung type, Plant species, Site and Cage type were subsequently used as fixed factors in two separate Mixed Models, in each case with seedling numbers (‘successful germination’) as the dependent variable. In the first, ‘open’ and ‘closed cages’ were the Cage type levels, while in the second analysis the ‘open cages’ group was split up in two: with and without traces of tunnelers. In both models, the correction for location of a plot according to the prevalent moisture gradient was made by incorporating the position of the strip it was in as a random factor (AR(1)-structured; Littell, Milliken, Stroup, & Wolfinger, 1996). We started from the full model, and used a backward procedure by stepwise eliminating non-significant contributions to reach the reduced models presented here.

To analyse the dung beetle fauna gathered in the pitfalls, we used General Linear Models to evaluate densities (instead of Mixed Models, as the corrections for location proved not to contribute significantly) and
an ordination technique to evaluate composition. In the linear models, the total number of dung beetles per sample and the numbers of dwellers and tunnelers were used as dependent variables. The fixed factors Cage type, Dung type, Site (and interactions) were used as independent variables. Species composition was analysed by detrended correspondence analysis (DCA), following suggestions of McCune and Grace (2002) on the determination coefficient ($r^2$). Rare species (occurrence in less than 3 out of 24 samples) were removed in advance, and densities were converted to relative numbers (percentage individuals of species $x$ occurring in sample $y$). We then tested whether the coordinates on the axes that represented the highest proportion of variation, were significantly different between the levels within each factor, using Wilcoxon two-sample tests.

Except for the DCA, which was done using Pc-Ord 4.26 (McCune & Mefford, 1999), statistical analyses were executed with the SAS software package (SAS Institute, 2003).

**Results**

**Dung beetle fauna**

Using the pitfall traps distributed among the 24 dung samples (12 per site), 650 Scarabaeidae individuals were caught, belonging to 12 species and 3 genera. The three most abundant species were *Aphodius contaminatus* Herbst (46% of all captured individuals), *A. foetens* Fabricius (27%) and *G. spiniger* Marsham (11%). The only other tunnelers, *O. similis* Scriba (1%) and *G. niger* Marsham (<0.2%), were far less numerous.

Numbers of tunnelers were significantly higher for the dry site. The opposite was true for dwellers: higher numbers were found at the moist site. The total number of dung beetles did not differ between the two sites (Table 1; Fig. 1). Apparently, at the moist site the open cages had a repulsive effect on some dweller species (as opposed to the dry site), resulting in a significant interaction of these factors in the dweller, and concomitantly, the total beetle analysis. Also, when visiting horse dung, tunnelers seemed to prefer the open-caged samples, whereas visiting cow dung, cageless pats were preferred. Moreover, the three-way interaction proved significant.

In the DC-analysis, only the first axis represented a substantial portion of the variance ($r^2 = 0.361, \text{gradient length} = 3.241$). The Wilcoxon two-sample test on the sample coordinates of this axis supported the significant segregation between sites ($Z = -2.859, p = 0.004$; Fig. 2). This supported the use of Site as a fixed factor in the analysis of the main experiment: the one site represented a high tunneler presence, the other a low tunneler presence. Moreover, species composition differed between cattle and horse dung ($Z = -2.051, p = 0.040$; Fig. 2), due to apparent preferences of *Aphodius* species (e.g. *A. sphacelatus* Panzer was found almost exclusively in the former, *A. contaminatus* in the latter).

![Fig. 1. Mean number (+S.E.) of dung beetle individuals caught in the pitfalls during the study period, separated in functional groups (dwellers plus tunnelers, and both separately). n.s., (*) and ** indicate $p > 0.10$, $p > 0.05$ and $p < 0.01$, respectively, for differences between sites.](image)

**Table 1.** Numbers of cases ($N$), degrees of freedom (df), $F$-statistics ($F$) and $p$-values ($p$) of the General Linear Models testing for the effects of Cage type, Dung type and Site on the total number of individuals caught per dung sample.

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>Dwellers + tunnelers</th>
<th>Dwellers</th>
<th>Tunnelers$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$N_1$, $N_2$</td>
<td>df</td>
<td>$F$</td>
</tr>
<tr>
<td>Cage type</td>
<td>12, 12</td>
<td>1</td>
<td>1.45</td>
</tr>
<tr>
<td>Dung type</td>
<td>12, 12</td>
<td>1</td>
<td>2.28</td>
</tr>
<tr>
<td>Site</td>
<td>12, 12</td>
<td>1</td>
<td>2.11</td>
</tr>
<tr>
<td>Cage type × Dung type</td>
<td>12, 12</td>
<td>1</td>
<td>1.13</td>
</tr>
<tr>
<td>Cage type × Site</td>
<td>12, 12</td>
<td>1</td>
<td>6.91</td>
</tr>
<tr>
<td>Dung type × Site</td>
<td>12, 12</td>
<td>1</td>
<td>0.23</td>
</tr>
<tr>
<td>Cage type × Dung type × Site</td>
<td>12, 12</td>
<td>1</td>
<td>13.01</td>
</tr>
</tbody>
</table>

$^a$The horse dung sample on the dry site showing the highest tunneler number skewed the residual distribution to be non-normal (Shapiro–Wilk’s test, $p < 0.001$). Nonetheless, the GLM is not altered when excluding this outlier (with regard to (non-)significant factors).
Densities of dung beetles did not differ between dung samples that were equipped with open cages and cageless samples (Table 1), nor did the species composition \((Z = 0.664, p = 0.507; \text{Fig. 2})\). Germination controls

The sites themselves apparently influenced germination differently: within the closed cages on the dry site, it was lower compared to closed cages on the moist site \((F = 11.39, \text{df} = 1, p = 0.002, \text{full model on closed cages data})\).

To a certain extent, the used dung was contaminated with seeds: on average, \(16.33 \pm 7.96\) alien seedlings germinated from each horse dung sample \((n = 6)\), of which \(1.67 \pm 1.97\) individuals could be attributed to Poa spp. and \(0.88 \pm 0.98\) to Agrostis spp. From these data, it was calculated that approximately \(2.6\% \) of the Agrostis-seedlings and \(4.0\% \) of the Poa-seedlings counted in samples of the main experiment actually concerned contaminations.

From the cattle dung samples, \(1.33 \pm 1.21\) alien seedlings emerged \((n = 6)\), none of which were Poa or Agrostis species. Since the contamination degree was less than \(5\% \) for horse dung, and \(0\% \) for cattle, its effect was not taken into account in further analyses.

Seeds of A. capillaris and P. pratensis were sufficiently viable, as shown in the germination controls with seed deposited in sand–peat substrate in a greenhouse \((A.c. 91.17 \pm 2.86\%; P.p. 84.50 \pm 6.53\%; n = 6 \text{ each})\) and in field conditions \((A.c. 61.58 \pm 7.28\%; P.p. 62.00 \pm 8.21\%; n = 12 \text{ each})\).

Cage type ('open' or 'closed cage') had no effect on germination success in controlled greenhouse conditions \((F = 0.095, \text{df} = 1, p = 0.760)\).

Main experiment

For A. capillaris and P. pratensis sown in the dung samples, germination varied greatly, from 2 to 66
seedlings per dung sample. There was also large variation in the degree to which dung beetles processed the dung: some samples were strongly fragmented and exhibited many dung beetle traces, while others were left intact.

Showing no significant contributions, four- and three-way interactions were stepwise eliminated from the full models, resulting in the reduced models presented in Table 2. Using all samples in the analysis, successful germination turned out to be significantly higher under closed cages than under open cages (Table 2; Fig. 3). Relative to this contrast, the difference between closed cages and the open-caged samples showing tunneler traces increased, whereas the difference with those showing no traces decreased (Table 3; Fig. 3). Thus, animals visiting open cages clearly brought about a substantial decline in seedling establishment, and this is especially true for tunnelers. Also, in both approaches, successful germination turned out to be higher on the moister than on the drier site (Table 2; Fig. 3). Overall, *P. pratensis* germinated better than *A. capillaris*, and germination from horse dung was higher than from cattle dung (Table 2), which is likely due to differential nutrient requirements from the plant species, and dung textures (loose vs. compact), respectively.

**Discussion**

Our results show that dung-visiting fauna had a negative effect on short-term, *in situ* germination for the two small-seeded species in a temperate grassland situation, and this was especially true for tunnelers. Since cage type did not affect germination conditions, and the open cages allowed access of a dung beetle community representative of the study area, both in numbers and composition, this effect can be assumed representative of natural situations.

However, apart from this direct effect on seedling establishment, dung beetles may contribute to the soil seed bank: seeds may remain dormant in the soil and germinate successfully later on. If this applies to a substantial number of seeds, the interaction could prove to be positive after all. Thus, whether the event under study here can indeed be considered as directed and beneficial (Vander Wall & Longland, 2004; Wenny, 2001), depends on the outcome on the long term. Indeed, seeds of *P. pratensis* are known to stay viable for more than 20 years, even at great burial depths (Goss, 1924).

Nonetheless, on the short-term scale, our experimental results clearly contrast the positive outcome found by Andresen and Levey (2004), who conducted the only field study following dung beetle mediated dispersal up to the point of seedling establishment until now. Evidently, differences in the dung beetle community, predator guild and/or plant species involved led to deviant patterns.

First, one of the possible dung beetle effects listed by Andresen and Feer (2005) is the reduction of seed clumping in a dung pat, and hence of seedling competition. However, this implies that seed movement by dung beetles also includes a horizontal component, which only is obvious in telecoprid nidification. Since rollers are abundant in tropical regions (cf. Andresen & Levey, 2004) but absent from typical assemblages of northern temperate regions (Hanski, 1991), this presumed positive effect does not occur in the latter.

Second, dung processing may also affect the microclimate that seeds experience. In the case of tunnelers, buried seeds may, for instance, experience more humidity and less light relative to ground level. This may positively as well as negatively affect the probability of germination, depending on the particular requirements of each species (cf. Wicklow et al., 1984). Because of dweller activities, seeds may end up in a more favourable mixed dung–soil substrate (see Holter, 1977).
For *A. capillaris* and *P. pratensis*, microclimatic effects were not considered in detail here, although a comparison between germination successes in dung (closed cages) and in sand–peat substratum hints dung to inhibit germination for both species.

Third, seed burial decreases the risk of seed predation as compared to unburied seeds. Indeed, seed removal by rodents after endozoochory is known for large-seeded plant species of tropical forests (Vander Wall & Longland, 2004) and, anecdotally, of temperate forests (e.g., Bermejo, Traveset, & Willson, 1998 [from bear faeces]; Page, Swihart, & Kazacos, 2001 [raccoon faeces]). In the case of ungulates ranging in temperate grassland systems, however, this as yet remains to be quantified. Although these animals mainly disperse dry, inconspicuous, usually small-sized fruits or seeds (Cosyns et al., 2005; Malo & Suarez, 1995; Pakeman, Digneffe, & Small, 2002), suchlike seeds are known to suffer high post-dispersal predation rates, too (Hulme & Benkman, 2002). For instance wood mice (*Apodemus sylvaticus*) and harvest mice (*Micromys minutus*) – both known to occur within the ‘Westhoek’ reserve (Smeers, 2001) – are known to forage for grass seeds in Western-European grasslands (Edwards & Crawley, 1999). These, as well as any seed or seedling predating group – e.g., insects, birds… – could be responsible for the losses witnessed under open cages. However, the experimental setup did not allow to quantify any of their contributions.

Last, Andresen and Feer (2005) state that “deep burial can also have a negative effect on seedling recruitment by preventing elongating seedlings from reaching the surface”. To evaluate this, we must consider the behaviour of *G. spiniger* Marsham, by far the most abundant tunneler in this study. This species constructs brood burrows consisting of a vertical shaft leading to a few horizontal tunnel-like brood chambers, each of the latter being filled with dung to make a horizontal sausage-shaped brood mass (about 100–150 mm in length and ca. 28 mm in diameter [Ampe, 2003; Klemperer, 1979]) which provides food for the future larva. These brood chambers are found between 10 and 30 cm depth (as measured within the study area by Ampe, 2003), depths at which immediate germination of the dung seed content is unlikely to occur (e.g. Grundy, Mead, & Burston, 2003). The statement of tunnelers having a negative effect on germination is unequivocally shown by our results: in a comparison of both analyses, the difference between the cage types increased when excluding the samples not visited by tunnelers (Fig. 3). Moreover, the between-site differences in tunneler numbers seemed to contribute to the between-site differences in germination: on the moist site, germination was less hampered by tunneler activity than on the dry site (significant interaction of Site and Cage type [interpretation is only meaningful for the analysis including all cages]; Table 2; Fig. 3). Indeed, tunnelers (almost exclusively *G. spiniger*) were more numerous at the drier site than at the moister site, which is most likely due to hydrological conditions: ground water level comes within reach of the shaft depths during the developmental stage of *G. spiniger* (September–March; Kühne, 1995) at the moist site, while it does not at the dry site. The contrasting pattern of dweller numbers can subsequently be attributed to the competitive hierarchy underlying the assemblage (see Hanski & Cambefort, 1991).

In conclusion, the negative short-term effect of tunnelers witnessed in this study is most likely attributed to deep burial. Whether burial ‘rescues’ seeds from being predated (cf. tropical studies) by means of seed bank enrichment, or is deleterious, remains to be quantified. Also, within one region, contrasting small-scale patterns may occur. If *Geotrupes* species were absent, and small *Onthophagus* species were present, for instance, the overall effect on immediate establishment might as well be positive since the latter bury dung to depths that might prove favourable to seedlings. However, since the dominance of *Aphodius* species is complemented with one or a few deep-tunnelers in many habitats, e.g. *Typhoeus typhoeus*, Linné in heathlands (Brussaard, 1985) and *Geotrupes stercorosus* Scriba in deciduous forests (Rembialkowska, 1982), the effect of dung beetle activity described here, might prove to hold elsewhere.

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