



REVIEW

SPECIAL ISSUE: THE ROLE OF SEED DISPERSAL IN PLANT POPULATIONS: PERSPECTIVES AND ADVANCES IN A CHANGING WORLD

Intrinsic and extrinsic drivers of intraspecific variation in seed dispersal are diverse and pervasive

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Abstract

There is growing realization that intraspecific variation in seed dispersal can have important ecological and evolutionary consequences. However, we do not have a good understanding of the drivers or causes of intraspecific variation in dispersal, how strong an effect these drivers have, and how widespread they are across dispersal modes. As a first step to developing a better understanding, we present a broad, but not exhaustive, review of what is known about the drivers of intraspecific variation in seed dispersal, and what remains uncertain. We start by decomposing 'drivers of intraspecific variation in seed dispersal' into intrinsic drivers (i.e. variation in traits of individual plants) and extrinsic drivers (i.e. variation in ecological context). For intrinsic traits, we further decompose intraspecific variation into variation among individuals and variation of trait values within individuals. We then review our understanding of the major intrinsic and extrinsic drivers of intraspecific variation in seed dispersal, with an emphasis on variation among individuals. Crop size is the best-supported and best-understood intrinsic driver of variation across dispersal modes; overall, more seeds are dispersed as more seeds are produced, even in cases where per seed dispersal rates decline. Fruit/seed size is the second most widely studied intrinsic driver, and is also relevant to a broad range of seed dispersal modes. Remaining intrinsic drivers are poorly

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understood, and range from effects that are probably widespread, such as plant height, to drivers that are most likely sporadic, such as fruit or seed colour polymorphism. Primary extrinsic drivers of variation in seed dispersal include local environmental conditions and habitat structure. Finally, we present a selection of outstanding questions as a starting point to advance our understanding of individual variation in seed dispersal.

Keywords: Crop size; fruit size; interindividual variation; intraindividual variation; seed dispersal effectiveness; seed dispersal traits.

Introduction

Intraspecific variation in seed dispersal has important consequences for individual reproductive success, plant population dynamics, community structure and evolution. For example, intraspecific variation in seed dispersal distances (Janzen 1970), the microhabitat destination of dispersed seeds (Schupp 1988) and the treatment in the mouth and gut (Traveset et al. 2007) affect demography and individual plant fitness through their impacts on the number of seeds dispersed, surviving, germinating and growing as seedlings. As a prominent example, dispersal kernels that include interindividual variation in dispersal distances are not equal to a population-level dispersal kernel based on mean dispersal distances. Including this intraspecific variation can alter the rate of population spread and the extent of gene flow (Schreiber and Beckman 2019; Wyse et al. 2019). Furthermore, individual variation in seed dispersal increases the range of habitats and conditions where seeds are dispersed, increasing the likelihood of the population to persist under unfavourable events (the portfolio effect; Bolnick et al. 2011). Although poorly studied, intraspecific variation in seed dispersal may also influence community assembly, species richness and responses to anthropogenic changes (Snell et al. 2019). See Snell et al. (2019) for a thorough review of the consequences of intraspecific variation in dispersal. However, given the historical focus in seed dispersal studies on population means, there are large gaps in our understanding of intraspecific variation in dispersal. We do not know how pervasive detectable variation in seed dispersal is, what the drivers of individual variation are and to what extent drivers have independent versus interactive effects. To date there only have been scattered efforts to summarize the breadth of our understanding of the drivers of intraspecific variation in seed dispersal.

The phrase 'intraspecific variation in the drivers of seed dispersal' is diffuse and subsumes multiple types of drivers and levels of variation. Decomposing this variation helps structure our thinking about intraspecific variation in dispersal. First, drivers of intraspecific variation in seed dispersal can be categorized as intrinsic variation based on trait expression of individual plants (e.g. fruit crop size, seed size, plant height) and extrinsic variation based on the ecological context of the plant (e.g. fruiting neighbourhood, topography). Further, intraspecific variation can be divided into variation among individuals (interindividual variation) and variation within individuals (intraindividual variation) (Herrera 2017). Most drivers of intraspecific variation in seed dispersal have both an interindividual and an intraindividual component (e.g. fruit size, fruit sugar concentration; even crop size or fruiting neighbourhood of the same individuals vary over time).

When considering drivers of intraspecific variation in seed dispersal, it is important to clarify what aspects and consequences of dispersal are being affected. Seed dispersal effectiveness, or SDE, depends on both the quantity of seeds dispersed (i.e. the immediate outcome of dispersal) and the

quality of dispersal provided to those seeds (i.e. the delayed consequences of dispersal; Schupp 1993; Schupp et al. 2010, 2017; reviewed in Box 1). While SDE is usually viewed as mean quantity multiplied by mean quality, these means are derived from a sample of individuals that likely differ substantially in both the quantity and the quality of dispersal. Beyond SDE, the probability of long-distance dispersal (LDD) can vary intraspecifically, which in turn contributes to population spread and gene flow. In this review, we focus mostly on seed movement, largely because that is what we have the most information on. However, we address consequences for seedling establishment or recruitment where relevant information is available.

In this paper, we provide a broad but not exhaustive review of the drivers of intraspecific variation in the quantity, and to a lesser extent, the quality components of seed dispersal (see Table 1 for a summary). We emphasize intrinsic drivers and interindividual variation because of our interest in individual fitness, defined as the contribution of an individual to future generations (Sæther and Engen 2015) (see Herrera 2009, 2017 for a focus on intraindividual variation in plant traits). However, we also consider intraindividual variation in traits because it can scale up to affect interindividual variation in dispersal. Further, intraindividual variation is not independent of interindividual variation. Lastly, we consider simple intraspecific variation in traits because much relevant work focuses on population-level trait variation without considering its apportionment into intra- and interindividual components.

We have several goals with this review. First, we illustrate the breadth of drivers of interindividual variation in seed dispersal. Second, we use diverse examples to illustrate the broad geographic and taxonomic scope of interindividual differences in seed dispersal, to assess how consistently they occur and to explore the range of impacts on seed dispersal processes. Third, we briefly discuss the barriers to fully understanding these drivers and their effects.

Intrinsic Variation: Plant Traits

Fruit crop size

Crop size (i.e. seed production) varies substantially among individuals and populations within a season and across years (e.g. Norghauer et al. 2011; Norghauer and Newbery 2015). Crop size is probably the most widely studied and best-supported driver of interindividual variation in the quantity of seeds dispersed. A strong positive relationship between crop size and the number of seeds dispersed and in the number of seeds being dispersed long distance is expected in abiotic dispersal modes such as anemochory and hydrochory and in the biotic dispersal mode epizoochory (Clark et al. 1998; Table 1). However, the expectation is less clear with endozoochory, synzoochory

Box 1. The Consequences of Seed Dispersal: Seed Dispersal Effectiveness and Long Distance Dispersal

Seed dispersal effectiveness, or SDE, can be defined ideally as the contribution a seed disperser makes to the production of new reproductive adults of a plant it disperses, whether the ‘disperser’ is a frugivorous bird, a seed-caching rodent or the wind (Schupp 1993; Schupp et al. 2010, 2017). $SDE = \text{quantity} \times \text{quality}$, where quantity is the number of seeds dispersed and quality is the probability that a dispersed seed successfully produces a new adult. However, in practical terms empirical studies are generally restricted to quantifying the contribution to some earlier relevant stage such as seedling establishment rather than new adults.

From the perspective of this review, the quantity of seed dispersal is straightforward and well-studied: the number of seeds dispersed. Quality, on the other hand, is influenced by a number of attributes of dispersal that arise repeatedly in this review. Three particularly important and frequent attributes of dispersal that arise when considering intraspecific variation in seed dispersal are briefly highlighted below.

Distance dispersed: The distance seeds are dispersed from the parent can affect the quality of dispersal in several ways. The most widely recognized consequence is increased survival by escaping from distance- and density-dependent seed and seedling enemies that concentrate attack beneath and near adult conspecifics (e.g. Janzen 1970; Connell 1971; Howe et al. 1985; Schupp 1988; Comita et al. 2014). Longer dispersal distances may also increase the chances of reaching unpredictably located suitable sites (Norghauer et al. 2011). Lastly, longer distance dispersal is important for gene flow and colonization of new sites (Nathan 2006; García et al. 2007; Jordano 2017). This attribute of dispersal is applicable to all modes of dispersal, biotic and abiotic.

Dispersal destination: Where in the landscape a seed is deposited can be described by the biotic and abiotic environments the potential recruit faces (Schupp and Fuentes 1995)—the seedscape (Beckman and Rogers 2013)—and these environments interact with the seed/seedling to determine its fate. There is substantial evidence that the habitat or microhabitat in which a seed is deposited, whether by a bird defecating, a rodent caching or a floating seed landing, has a large influence on seed and seedling fate (e.g. Schupp 2007; Zhang et al. 2013; Young and Kelly 2018). Further, whether seeds are deposited widely scattered or in high density clumps at latrines, sleeping trees or favourite processing sites influences seedling competition and susceptibility to density-dependent natural enemies independent of distance from the parent (e.g. Schupp et al. 2002).

Treatment in the mouth and gut: For animal-vectored dispersal, the first critical distinction is whether all seeds are being treated gently and dispersed physically intact or whether some-to-many are broken or damaged (Schupp 1993). Secondly, for intact seeds it can matter whether the seed is (i) dropped after some of the pulp has been picked off and consumed, (ii) swallowed and either regurgitated or spit out clean or (iii) swallowed, passed through the digestive track, and defecated. These different pathways can result in differences in germination (e.g. Rodríguez-Pérez et al. 2005; Reid and Armesto 2011; Haurez et al. 2018) and in post-dispersal interactions with seed predators and secondary dispersers (e.g. Fricke et al. 2016; Pan et al. 2016; Guerra et al. 2018).

and myrmecochory, where animal dispersers make foraging decisions in resource-heterogeneous environments where these dispersers can be satiated (e.g. Manasse and Howe 1983; Hampe 2008; Table 1) or prematurely leave feeding trees in order to mix diets with complementary resources (Whelan et al. 1998; Morán-López et al. 2018a). Nonetheless, expected patterns with respect to crop size have been proposed for endozoochory. Howe and Estabrook (1977) developed two models based on specialized (model 1) versus opportunistic (model 2) frugivore/seed dispersal systems. They suggested that the number of seeds dispersed should increase with fruit availability for both types of species, although the number dispersed should plateau for model 1 plant species that depend on specialized dispersers that tend to involve relatively few species and become satiated. They further predicted that the effect of crop size on the proportion of the available seeds dispersed would differ for model 1 and model 2 species. For model 1 species, the proportion of seeds dispersed was expected to initially increase with crop size but would reach a peak at some intermediate crop size due to disperser satiation and then drop with ever larger fruit crops. In contrast, for model 2 species they predicted that the proportion of the seed crop dispersed would increase with increasing crop size, perhaps stabilizing at a constant proportion at larger crop sizes, but not decreasing. However, the dichotomy between specialized and opportunistic dispersal systems is not generally accepted at this point (Schupp et al. 2010), leading other authors (e.g. Carlo et al. 2007) to propose a general expectation that the number of seeds dispersed should increase with increasing crop size. In fact, this is considered one major driver of the development of frugivory hubs, where hub individuals in the network (those

with the largest fruit crops) receive more dispersal services than expected, leaving non-hub individuals with little dispersal services (Carlo et al. 2007).

This last prediction appears to be supported by studies mostly of endozoochory that demonstrate that as crop size increases, visitation rate by avian (e.g. Saracco et al. 2005; Ortiz-Pulido et al. 2007; Guerra et al. 2017) and mammalian (e.g. Guitián and Munilla 2010) dispersers increases, which translates into an increased quantity of seeds dispersed (Table 2). For example, *Prunus mahaleb* fruit crop size explained 80 % of seeds dispersed in a population in southern Spain (Jordano and Schupp 2000). With respect to the proportion of seeds dispersed, results to date show no consistent relationship (Table 2). These patterns suggesting a general increase in the number but not the proportion of seeds dispersed with increasing crop size are supported by a meta-analysis that found positive bird-mediated selection on fruit crop sizes as measured by both visitation rate and the quantity of seeds dispersed, but no selection on the proportion of seeds dispersed (Palacio and Ordano 2018).

Crop size can also affect the quality component of SDE and the probability of LDD. Increasing crop size lifts the entire dispersal kernel, resulting in more seeds in the tail of the distribution and thus more LDD and increased population spread and gene flow (Clark et al. 1998). Increasing crop size also results in more seeds dispersing farther in a local dispersal context, increasing the chances of reaching suitable sites (Norghauer et al. 2011) and surviving distance- and density-dependent mortality (Janzen 1970; Connell 1971). Although empirical evidence is limited, crop size can also affect the quality of endozoochorous dispersal by altering disperser behaviour and disperser assemblages of

Table 1. An overview of how intrinsic variation in plant traits influences seed dispersal quantity and quality. Quantity is indicated by solid arrows, while dashed arrows are for quality. Grey arrows indicate uncertainty (i.e. we assume this relationship to be true but no studies have explicitly measured this), and vertical lines without arrowheads indicate a lack of relationship. Representative references are included, however this is not meant to be an exhaustive list. See text for more details.








	Abiotic dispersal			Biotic dispersal			
	 Anemochory (wind)	 Hydrochory (water)	 Ballistic (explosive)	 Endozoochory (ingested)	 Epizoochory (attached outside)	 Synzoochory (carried away)	 Myrmecochory (ants)
Crop size	Larger crops have more seeds dispersed (Norghauer et al. 2011) ↑	Larger crops have more seeds dispersed ↑	Larger crops have more seeds dispersed ↑	Larger crops have more seeds dispersed ↑ Larger crops have higher visitation rates No impact on the proportion of seeds dispersed (Palacio and Ordano 2018) ↓	Larger crops have more seeds dispersed ↑	Removal rates slow (Greenberg and Zarnoch 2018) or increase (Pesendorfer and Koenig 2016) with crop size Caching rates and seed survival in caches increase with crop size (Pesendorfer et al. 2016, Zwolak et al. 2016) ↑	Larger crops have more seeds dispersed (Boulay et al. 2007) ↑
Fruit and seed size	Lighter seeds travel farther (Larios and Venable 2015) ↓	Lighter seeds travel farther (Delefosse et al. 2016) ↓ Heavier seeds survive longer in the water (Manasse 1990) ↑	Inconsistent patterns between seed mass and dispersal distance (Lisci and Pacini 1997; Rezvani et al. 2010) ↔ Larger fruits had longer dispersal (Jacobs and Lesmeister 2012) ↑	Both larger and smaller seeds are preferred (see Table 3) ↔ Variation in fruit/seed size can increase the diversity of disperser assemblages ↑↑		Larger seeds are preferred, dispersed further and more likely to be cached (see Table 3) ↑↑	The largest and smallest seeds are dispersed (Manzaneda et al. 2009) ↔
Fruit and seed morphology	Seeds travel shorter distances with greater wing loading (Sinha and Davidar 1992, Skarpaas et al. 2011) ↓ Shorter, fatter seeds traveled farther (Gravuer et al. 2003) ↑			Larger fruit/seed size select for fewer dispersers (greater specialization) (McKey 1975, Tutin & Fernandez 1994)	More complex diaspores are dispersed more frequently and dispersed further in disturbed areas and grassland (Monty et al. 2016) ↑↑		
Height	Seeds released from taller plants travel farther (Zhu et al. 2016) ↑		Seeds released from taller plants travel farther (Gómez 2007) ↑	Taller or smaller trees have a higher percentage of fruit removed (Ortiz-Pulido et al. 2007, Crestani et al. 2019) ↓	Fruits attached to different parts of the animal, travelled different distances (Liehrmann et al. 2018) ↔		
Rewards and deterrents				Higher sugar or calorie content preferred (Levey 1987; Palacio et al. 2014) ↑ Fruit with greater pulp mass are preferred (Wheelwright 1993; Luna et al. 2016) ↑ Higher ratio of reward:seed preferred (Manasse and Howe 1983; Sobral et al. 2010, although see Crestani et al. 2019) ↑		Preference for lower tannin concentration in acorns (Shimada et al. 2015; Wang & Yang 2015) ↓	Ants preferred elaiosomes richer in oleic acid (Boulay et al. 2007) ↑

Table 2. Examples of studies reporting the relationships between interindividual variation in plant crop size and both the number of seeds removed (#) and the proportion of the seed crop removed (prop.). Arrows denote shape of the relationship between crop size and the variable, with a dot representing no information.

Species	Form of the relationship with increasing crop size		Reference
	# seeds removed	Prop. seeds removed	
Tropical endozoochorous tree			
<i>Casearia corymbosa</i>	↗	↘	Ortiz-Pulido et al. (2007) (#), Howe and Vande Kerckhove (1979) (prop.)
<i>Eugenia uniflora</i>	↗	•	Blendinger and Villegas 2011
<i>Guarea glabra</i>	↗		Howe and De Steven (1979)
<i>Virola nobilis</i>	•		Manasse and Howe (1983)
<i>Virola surinamensis</i>	↗	↗	Moreira et al. (2017)
Tropical endozoochorous shrub			
<i>Erythroxylum havanense</i>	↗	•	Gryj and Domínguez (1996)
<i>Miconia fosteri</i>	↗	↗	Blendinger et al. (2008)
<i>Miconia irwinii</i>	↗	•	Guerra et al. (2017)
<i>Miconia serrulata</i>	↗	↗	Blendinger et al. (2008)
Temperate endozoochorous tree			
<i>Olea europaea</i>	↗		Alcántara et al. (1997)
<i>Prunus mahaleb</i>	↗	↘	Jordano (1995)
Temperate endozoochorous shrub			
<i>Crataegus monogyna</i>	↗	•	Sallabanks (1993)
<i>Prunus virginiana</i>	↗		Parciak (2002)
<i>Sambucus pubens</i>	↗	•	Denslow (1987)
Temperate synzoochorous tree			
<i>Quercus lobata</i>	↗	•	Pesendorfer and Koenig (2016)

individual plants. For example, as *Vassobia breviflora* crop size increased, disperser residence time in the canopy decreased, increasing the probability of seed dispersal away from the parent rather than seeds processed in situ (Palacio et al. 2017). Increasing crop size also increased fruit consumption by legitimate dispersers (gulpers) without affecting consumption by pulp consumers, altering the realized dispersal assemblage and increasing overall dispersal quality (Palacio et al. 2017). Lastly, with a population-wide increase in *Fagus sylvatica* crop size (masting), there was an increase in survival of seeds cached by *Apodemus flavicollis*, a clear increase in the quality of dispersal (Zwolak et al. 2016; Table 1). On the other hand, seed survival in caches can be lower under trees that produced large seed crops (Schubert et al. 2018); thus, spatial variation in crop size might have different effects than temporal variation (masting).

Fruit and seed size

Fruit and seed size variation is likely the second most widely studied driver of interindividual variation in seed dispersal. Fruit and seed size vary within individuals, but also among individuals, years and populations (Sobral et al. 2013; González-Varo and Traveset 2016; Herrera 2017). In a study of 39 species from 46 populations, on average 62 % of seed size variation was within individuals while 38 % was among individuals, though individual species varied substantially (Michaels et al. 1988). Thus, fruit and seed size variation can influence animal disperser decisions regarding which plants to forage in (interindividual) and then which fruits to consume (intraindividual) (Vanthomme et al. 2010; Effiom et al. 2013). Furthermore, mean fruit size of individuals can be highly heritable, indicating potential selection response (Wheelwright 1993; Galetti et al. 2013).

Many studies have demonstrated size-based fruit or seed selection by dispersers, suggesting a potentially important

role for fruit/seed size in driving interindividual variation in the quantity component of SDE, although actual patterns of selection are not consistent and appear to depend on the plant and animal species involved (Table 3a). It is generally thought that fruit/seed size-based selection is a function not so much of fruit/seed size, but rather by the fruit/seed size relative to the disperser size. For example, for endozoochorous birds that swallow fruits whole, it is widely believed that fruit selection is driven by fruit diameter and bird gape width (e.g. Wheelwright 1993; González-Varo and Traveset 2016). Similarly, it is thought that seed size selection by synzoochorous seed dispersers is related to the ratio of seed to disperser size (Muñoz and Bonal 2008a).

The extent to which fruit/seed size selection contributes to interindividual differences in the quantity of seeds dispersed is unclear. Dispersers may select among individual plants based on mean traits or among individual fruits independent of the mother plant. While some studies demonstrate that frugivores select among fruiting plants based on mean fruit or seed size (e.g. Howe and Vande Kerckhove 1981; Wheelwright 1993; Alcántara et al. 1997; Martínez et al. 2007), others demonstrate that at the population level, individual fruits are selected based on their sizes (e.g. Parciak 2002; Hernández 2009; Larrinaga 2010). Thus, even strong selection of fruits based on size need not lead to differential selection of individual plants based on fruit size. For example, dispersers of *P. mahaleb* strongly selected fruits based on size, but this was almost entirely driven by selection of smaller fruits within an individual plant's fruit crop, while there was inconsistent and weak selection among individual plants based on fruit size (Jordano 1995). Because multiple traits associated with selection by dispersers may be correlated with seed size, the degree to which selection is driven by fruit or seed size, rather than a correlated trait is unclear

Table 3. Examples of studies on fruit or seed size selection by animal dispersers. In endozoochory (a) selection always denotes preferential removal/dispersal. In synzoochory (b) we consider the relationship between seed size and various quantitative (removal/dispersal) and qualitative (consumption, caching %, caching distance, seedling production) metrics of SDE. For myrmecochory (c), the only study of which we are aware, is presented as in endozoochory (a).

Plant species	Animal dispersers	Selection for	Reference
(a) Endozoochory			
<i>Corema album</i>	<i>Oryctolagus cuniculus</i>	Smaller	Larrinaga (2010)
<i>Viburnum opulus</i>	<i>Erithacus rubecula</i> , <i>Turdus philomelos</i>	Smaller	Hernández (2009)
<i>Prunus mahaleb</i>	Birds	Smaller	Jordano (1995)
<i>Prunus virginiana</i>	Birds	Smaller	Parciak (2002)
<i>Viola nobilis</i>	Birds	Smaller	Howe and Vande Kerckhove (1981)
<i>Crataegus monogyna</i>	<i>Turdus migratorius</i>	Larger	Sallabanks (1993)
<i>Crataegus monogyna</i>	<i>Turdus</i> spp.	Larger	Martínez et al. (2007)
<i>Ocotea tenera</i>	Birds	Larger	Wheelwright (1993)
<i>Henriettea succosa</i>	Birds	Larger	Crestani et al. (2019)
<i>Olea europaea</i> var. <i>sylvestris</i>	Birds	Larger (1 of 2 years)	Alcántara et al. (1997)
(b) Synzoochory			
<i>Quercus ilex</i>	<i>Garrulus glandarius</i>	Trees with smaller acorns, but larger individual acorns (removal, one of two habitats)	Morán-López et al. (2015a)
<i>Carapa procera</i>	<i>Myoprocta acouchy</i>	Larger (removal, caching %, and caching distance)	Jansen et al. (2004)
<i>Myrcianthes coquimbensis</i>	Rodents	Larger (removal and caching %)	Luna et al. (2016)
<i>Quercus rubra</i>	Rodents	Larger (caching %)	Wróbel and Zwolak (2017)
<i>Pinus armandii</i>	<i>Apodemus latronum</i> and <i>Apodemus chevrieri</i>	Larger (removal, consumption, and caching %)	Wang and Ives (2017)
<i>Astrocaryum mexicanum</i>	<i>Heteromys desmarestianus</i>	Larger (consumption); no effect (caching); smaller (caching distance, cache survival)	Brewer (2001)
<i>Quercus ilex</i>	Rodents	No effect (removal); larger (caching %, caching distance, cache survival)	Gómez et al. (2008)
<i>Quercus serrata</i>	Rodents	No effect (removal and caching); larger (caching distance, cache survival)	Xiao et al. (2004)
<i>Quercus ilex</i>	<i>Apodemus sylvaticus</i> and <i>Mus spretus</i>	No effect (removal by <i>A. sylvaticus</i>); smaller (removal by <i>M. spretus</i>)	Muñoz and Bonal (2008a)
<i>Quercus robur</i>	<i>Garrulus glandarius</i>	Intermediate-sized (removal)	Bossema (1979)
<i>Carapa oreophila</i>	Rodents	No effect (removal, consumption, caching %, and caching distance)	Yadok et al. (2018)
<i>Pittosporopsis kerrii</i>	Rodents	Larger (removal); intermediate (caching %, caching distance, seedling production)	Cao et al. (2016)
(c) Myrmecochory			
<i>Helleborus foetidus</i>	Ants	Smaller and larger (removal)	Manzaneda et al. (2009)

(Jordano 1984; Martínez et al. 2007). For example, the four main avian dispersers of *Rubus ulmifolius* in southern Spain differed in the distribution of seed sizes dispersed, but seed size, seed number, pulp/seed ratio and percent pulp co-varied, making it difficult to determine which trait or traits were being selected (Jordano 1984).

Fruit and seed size can also drive intraspecific variation in the quantity and quality of dispersal in other ways. For gape-limited, endozoochorous birds, intra- and interindividual variation in fruit diameter can affect the proportion of a plant's fruit crop that a disperser can swallow. For instance, in a *Myrtus communis* population in southern Spain, some individuals produced large fruits that only *Turdus merula* and *T. philomelos* could swallow and disperse. However, other individuals in the same population produced smaller fruits that were completely available to these species and partially available to *Sylvia atricapilla*, *Erithacus rubecula* and, in the case of one individual, the smallest disperser,

S. melanocephala (González-Varo and Traveset 2016). Thus, the realized disperser assemblages of individual plants varied from two to five species. Moreover, realized disperser assemblages of individual plants varied across years due to changes in fruit size.

Such among-individual and among-year variation in realized disperser assemblages can affect interindividual variation in dispersal outcomes. First, variation in the number of animal species feeding on an individual plant likely affects the quantity of seeds dispersed (Schupp et al. 2010). Second, interindividual variation in realized disperser assemblages is expected to drive interindividual variation in LDD, gene flow and the quality of dispersal because disperser species differ in their dispersal kernels, treatment in the mouth and gut, and microhabitat destination of seeds (Jordano and Schupp 2000; García et al. 2007; Schupp et al. 2010). Species-specific preferences in microhabitat and fruit/seed size can also result in microhabitats accumulating different seed size distributions (Obeso et al. 2011).

Lastly, seed size affects whether a seed is swallowed and passed through the digestive system versus being dropped, spat out or regurgitated, which affects both treatment in the mouth and gut and dispersal distances (Corlett and Lucas 1990; Jordano 1992; Kunz and Linsenmair 2008).

Fruit and seed size also affect synzoochorous and myrmecochorous dispersal (Table 3b). In general, larger seeds tend to be dispersed more rapidly and farther, and are more likely to be cached than smaller seeds; in contrast, no obvious pattern links seed size and the probability of surviving in a cache. The actual outcome of the interaction may be more related to the ratio of seed to disperser size rather than seed size alone (Muñoz and Bonal 2008a). However, three species of rodents varying 4-fold in mass all preferentially selected and dispersed larger fruits of the Chilean desert shrub *Myrcianthes coquimbensis* (Luna et al. 2016).

While there is abundant evidence that synzoochorous dispersers select and handle individual seeds based on size, there are fewer studies documenting dispersers selecting on mean seed size among individual plants. The large Japanese wood mouse (*Apodemus speciosus*) preferentially dispersed seeds of individual *Q. serrata* trees with larger mean acorn size (Shimada et al. 2015). Similarly, *Apodemus* spp. disproportionately dispersed and cached seeds from *Pinus armandii* individuals with larger mean seed mass (Wang and Ives 2017), although the greater probability of their seeds being consumed cancelled the benefits of increased dispersal. By contrast, mean seed size of the Queen palm (*Syagrus romanzoffiana*) had no influence on tree selection by squirrels (Alves et al. 2018), and *Garrulus glandarius* preferentially fed on *Quercus ilex* trees with smaller acorns (Morán-López et al. 2015a). Thus, although evidence is limited, there is potential for seed size to contribute to interindividual variation in the quantity and quality of seed dispersal by synzoochorous dispersers.

Although the consequences of within-individual variation in plant traits have not been considered frequently in ecology (e.g. Herrera 2017), in addition to selection based on individual or mean fruit/seed size, mutualistic dispersers may select among individual plants based on the extent of intraindividual variation in fruit or seed size. In a latitudinal study of *Crataegus monogyna* seed dispersal by *Turdus* spp. in Europe, birds selected against intraindividual fruit size variation in populations with lower variation and selected for intraindividual fruit size variation in populations with higher variation (Sobral et al. 2013). Similarly, *A. speciosus* not only selected individual *Q. serrata* trees with larger acorns, but also selected individual trees with a greater variability in acorn weight (Shimada et al. 2015).

Seed size also affects abiotic seed dispersal. In the seagrass *Zostera marina*, settling rate increases with seed size, suggesting smaller seeds disperse farther (Delefosse et al. 2016). When grazed by the specialist herbivore *Ophraella communa*, the riparian weed *Ambrosia artemisiifolia* produces lighter, more buoyant seeds, demonstrating a clear mechanism for interindividual variation in dispersal (Fukano et al. 2014). In ballistically dispersed species, both seed and fruit size can affect patterns of seed dispersal. In *Oxalis acetosella* (Berg 2000) and *O. corniculata* (Rezvani et al. 2010), dispersal distances increased with seed mass, while in *Mercurialis annua*, dispersal distances decreased with increasing seed mass (Lisci and Pacini 1997). In the only study on fruit size and ballistic dispersal of which we are aware, dispersal distance increased with fruit length in *Erodium cicutarium* (Jacobs and Lesmeister 2012).

It is generally believed that dispersal distances of anemochoric species will decrease as seed mass increases, and

this expectation appears to be well-supported, although the variance explained is generally low. This general pattern has been reported in both tropical and temperate environments as well as across trees, shrubs and herbaceous species (Morse and Schmitt 1985; Bhuyan et al. 2000; Meyer and Carlson 2001; Bullock et al. 2003; Debain et al. 2003; Gravuer et al. 2003; Skarpass et al. 2011), although there are exceptions (e.g. Wyse et al. 2019). Given that seed mass varies both among and within individuals (Sinha and Davidar 1992; Gravuer et al. 2003), seed mass variation may contribute to interindividual variation in dispersal distances. For example, under highly competitive conditions, plants of the wind-dispersed desert annual *Dithyrea californica* produce smaller, lighter seeds that are dispersed farther (Larios and Venable 2015). Given the typical heterogeneous distribution of individuals in populations, it is likely that *D. californica* individuals vary continuously in competitive environments and thus potentially in dispersal ability. Finally, the actual pattern of wind dispersal is driven not simply by seed mass, but by the relationship between seed mass and the dispersal structure (e.g. pappus, Skarpass et al. 2011; and see below under Morphology).

Height: plant, seed abscission, seed attachment

Interspecific studies of the effect of plant height on seed dispersal suggest plant height is a major correlate of dispersal distances and is considerably more important than seed size (Thomson et al. 2011, 2018; Augspurger et al. 2017). However, given that height of reproductive adults varies substantially more among than within species, it need not follow that the more limited interindividual variation in plant height will be a major driver of interindividual variation in seed dispersal distances. Nonetheless, limited empirical evidence suggests that interindividual height variation might be at least a minor driver of interindividual variation in dispersal, at least for abiotically dispersed species. With anemochory, plant (or seed release) height has been shown to be positively related to dispersal distances in trees (Sinha and Davidar 1992) and herbaceous perennials (Sheldon and Burrows 1973; Weiblen and Thomson 1995; Skarpass et al. 2011; Zhang et al. 2011; Zhu et al. 2016; DiTommaso et al. 2018), in some cases relatively strongly (e.g. Zhu et al. 2016). In *Carduus nutans*, height, and therefore dispersal potential, is environmentally plastic, increasing with simulated climate change (Zhang et al. 2011). Interestingly, increasing tree height in *Lophopetalum wightianum* not only led to greater dispersal distances and larger seed shadows, but also more even seed dispersion, potentially decreasing density-dependent mortality (Sinha and Davidar 1992). Norghauer et al. (2011) suggest that tree height does not affect dispersal distances of mahogany in Amazonian forests because all reproductive trees are emergent above the canopy and exposed to winds. Note, however, that much of our understanding comes from controlled releases of seeds at fixed heights in wind tunnels, and that the heterogeneity of the real world likely reduces the explanatory power of plant height. Intraindividual variation in seed release height could further obscure any potential interindividual variation in dispersal distances based on variation in plant height. Interindividual height variation is also important for ballistic dispersal. Increasing height resulted in increasing dispersal distances in the mustards *Erysimum mediohispanicum* (Gómez 2007), *Arabidopsis thaliana* (Wender et al. 2005) and *Lepidium campestre* (Thiede and Augspurger 1996), but not in *O. acetosella* (Berg 2000).

We have less direct evidence that height is important for biotic dispersal. Interspecific studies suggest that seed presentation height can affect the frequency of epizoochorous

dispersal (Hughes et al. 1994; Fischer et al. 1996; Graae 2002; Wessels et al. 2008; Hovstad et al. 2009; Albert et al. 2015), but data for intraspecific effects of height variation are limited. In two studies on small tropical trees, variation in interindividual plant height affected endozoochorous seed dispersal; in *Casearia corymbosa*, plant height very weakly affected fruit removal (Ortiz-Pulido et al. 2007), and in *Henrietta succosa*, dispersers favoured shorter individuals (Crestani et al. 2019). Other studies have documented vertical segregation of frugivore communities in tropical forests (Shanahan and Compton 2001; Poulsen et al. 2002; Lahann 2007; Yoshikawa et al. 2009; Flörchinger et al. 2010), so fruiting conspecifics of different heights may have at least marginally different disperser assemblages, which could affect both the quantity and quality of dispersal.

Rewards and deterrents

Effects of variation in rewards and deterrents on seed dispersal have been studied extensively, though mostly from the perspective of interspecific differences (e.g. Cazetta et al. 2008 for frugivorous bird-dispersed trees; Vander Wall 2010 for rodent-cached trees); effects of intraspecific variation have received much less attention. Nonetheless, available data suggest that interindividual variation in rewards and deterrents may be an important driver of interindividual variation in seed dispersal in some cases.

Intraspecific variation in fleshy fruit seed dispersal driven by intraspecific variation in rewards offered to animal dispersers has been addressed in a variety of ways with some studies focusing on the quality of the reward, and some on the absolute or relative quantity of reward. Interspecific comparisons show that fruit colour is predictive of protein (brightness), sugar (chroma) and lipid (darkness and chroma) concentrations (Cazetta et al. 2012; Schaefer et al. 2014), and that these signals can be effective even if relationships are weak (Albrecht et al. 2018). We have long known of interindividual variation in fruit sugar content and that dispersers can distinguish these differences by taste (Levey 1987). However, it is not known whether dispersers can distinguish intraspecific differences in nutrients based on colour. Nonetheless, dispersers distinguish among plants that differ in rewards in some manner (Crestani et al. 2019). In China, the frequency with which the deer *Muntiacus muntjak* visited fruiting *Choerospondias axillaris* trees was correlated with mean Kcal/fruit (Chen et al. 2001). Similarly, birds selected for *Celtis ehrenbergiana* trees (Palacio et al. 2014) and *Sambucus pubens* shrubs (Denslow 1987) with greater mean sugar concentration of fruits. However, frugivores preferentially fed on *Henrietta succosa* individuals with intermediate sugar concentrations (Crestani et al. 2019). These limited results suggest that variation in fruit energy content is a potential driver of interindividual variation in seed dispersal. We are aware of no evidence that endozoochorous dispersers discriminate among plants based on variation in other nutrients. Individual *Viola nobilis* trees varied substantially in protein, lipid and non-structural carbohydrate content, but this variation did not explain variation among trees in seed dispersal (Manasse and Howe 1983).

There is more extensive evidence that either the absolute or relative quantity of reward is important in fruiting plant selection by dispersers. As noted previously, independent of body size, three rodent species preferentially selected larger fruits of the shrub *M. coquimbensis*; larger fruits had more pulp, and it was the pulp, not the seed, that was consumed (Luna et al. 2016). In *Ocotea tenera*, dispersers selected trees with larger diameter fruits, as noted above, apparently due to the greater pulp mass; fruit diameter explained much of the variation in

total pulp mass ($r^2 = 0.56$) but not in the ratio of seed mass/fruit mass ($r^2 = 0.01$) (Wheelwright 1993). However, more typical are studies demonstrating selection driven by intraspecific variation in relative rather than absolute reward per fruit. In *V. nobilis* (Howe and Vande Kerckhove 1981; Manasse and Howe 1983), *V. calophylla* (Russo 2003) and *C. monogyna* (Sallabanks 1993; Martínez et al. 2007; Sobral et al. 2010), variation in dispersal was explained in part by mean pulp:seed ratio but not by the absolute quantity of pulp. Sometimes this resulted in the selection of smaller fruits (Howe and Vande Kerckhove 1981), sometimes the selection of larger fruits (Martínez et al. 2007; Sobral et al. 2010).

Such interactions between fruit size and either pulp quantity or pulp:seed ratio suggest caution when considering the role of fruit or seed size variation in interindividual variation in seed dispersal. As noted, many studies report fruit selection based on fruit or seed size, and size clearly has an impact on seed dispersal, at a minimum by filtering which dispersers can swallow and disperse the seeds of endozoochorously dispersed species. However, in many cases where dispersers select larger or smaller fruits, we do not know if it is selection based on fruit/seed size or rather selection based on a correlated trait such as absolute or relative quantity of the reward.

Most work on deterrents with fleshy-fruited plants has focused on hypotheses addressing why ripe fruits contain toxins rather than assessing interindividual variation in concentrations and dispersal. In the single field study of which we are aware, mean emodin concentration in *Rhamnus alaternus* fruit pulp was unrelated to fruit removal rate among plant individuals in 1 year but was positively correlated with removal rate in another year (Tsahar et al. 2002). Levey and Cipollini (1998) showed that cedar waxwings (*Bombycilla cedrorum*) feeding on artificial fruits discriminated against 'fruits' containing realistic concentrations of α -solamargine compared to 'fruits' without this glycoalkaloid, but did not discriminate among artificial fruits that differed in α -solamargine concentration. However, potential seed disperser species differ in sensitivity to varying tannin concentrations in artificial fruits (Zungu and Downs 2015). Whitehead and Poveda (2011) reported a potential environmental influence on intraspecific variation in deterrents; in *Hamelia patens*, artificial herbivory of subtending leaves reduced fruit removal in adjacent inflorescences due to reduced palatability, presumably a result of herbivory-induced chemical changes. Plant secondary compounds also can alter gut retention times of seeds, which can affect seed dispersal distances and germination (Tewksbury et al. 2008; Baldwin and Whitehead 2015), although we know little about the degree of natural intraspecific variation, particularly interindividual variation, in secondary compounds, and the consequences of this variation.

In myrmecochorous species, chemical composition of the elaiosome has been studied mostly through interspecific comparisons of seed and elaiosome chemical profiles (e.g. Fischer et al. 2008). Fatty acid composition in *Euphorbia characias* elaiosomes, especially of oleic acid, varies among populations and among individuals within populations, but not within individuals (Boieiroa et al. 2012). In the related *Helleborus foetidus*, this variation is ecologically important (Boulay et al. 2007) as seed-dispersing ants preferentially visited plants with elaiosomes richer in oleic acid. Many consider oleic acid to be an attractant or behavioural trigger rather than a reward because elaiosomes are rich in other critical nutrients, such as amino acids (Fischer et al. 2008). However, it is likely both attractant and reward since it evokes seed harvesting and it is the biosynthetic precursor of the essential nutrients linoleic and linolenic acids (Fischer et al. 2008).

In synzoochorous dispersal systems, evidence presented above that these dispersers frequently preferentially harvest, disperse, disperse farther or cache larger seeds reflects a response to the reward offered since the seed is both the propagule being dispersed and the reward offered for dispersal (Gómez et al. 2019). However, whether it is seed size itself or energy content that drives the decision is unclear. A study with artificial seeds suggests the answer is complicated in that energy was the primary predictor of initial harvest, but size was the primary determinant of post-harvest fate (removed versus *in situ* consumption, distance dispersed, and cached versus consumed after dispersal; Wang and Yang 2014).

Variation in seed defences can simultaneously affect the risk of seed predation and the probability of seed dispersal. A particularly well-studied example of this phenomenon involves mechanical seed defences in synzoochorous lamber pine (*Pinus flexilis*). Cone structure in this species is under a conflicting selective pressure from the red squirrel (*Tamiasciurus hudsonicus*), a predispersal seed predator, and the Clark's nutcracker, a seed disperser (Siepielski and Benkman 2010). Individuals that produce particularly well-defended cones tend to be dispersed by *Peromyscus* mice rather than the Clark's nutcracker (Siepielski and Benkman 2008), resulting in much shorter dispersal distances and different habitat destinations of the seeds.

Endocarp or seed coat thickness as a deterrent has long been considered an evolutionary response to synzoochory, in particular to managing the dual role of seed dispersal and seed predation, and there is substantial interspecific support for the idea that these dispersers select and handle seeds at least partly based on endocarp or seed coat thickness (see Vander Wall 2010). Intraspecific studies on the effects of seed coat thickness are sparse and inconsistent; mean endocarp thickness of the palm *S. romanzoffiana* did not influence tree selection by the squirrel *Guerlinguetus ingrami* (Alves et al. 2018), but *P. flexilis* seeds with thicker seed coats were more likely to be cached by rodents and were dispersed further than seeds with thinner coats (Siepielski and Benkman 2008). Chemical deterrents may also play a role in interindividual dispersal by synzoochorous dispersers. Mice preferred individual *Quercus serrata* trees producing acorns with lower mean and coefficient of variation of tannin concentration (Shimada et al. 2015). Similarly, rodents were less likely to remove acorns produced by *Q. rubra* trees from nitrogen-addition plots, presumably due to changes in chemical composition (Bogdziewicz et al. 2017).

Interpretation of these results is complicated by frequent co-variation of size, nutritional content and concentration of secondary metabolites in fruits and seeds (e.g. Izhaki et al. 2002), as discussed above with co-variation of fruit size and absolute or relative quantity of reward in fleshy-fruited plants. Interestingly, co-variation of these traits can even vary with position of fruits in the plant canopy (Houle et al. 2007). Recent innovative work with artificial fruits has begun to tease apart the disparate roles of size, nutrients and secondary compounds in seed selection by caching rodents. Wang and Yang (2015) manipulated seed size, tannin, fat, protein and starch content in artificial seeds and showed how all these factors affected rodent foraging. Rodents preferentially removed seeds with less tannin; increasing fats, and to a lesser degree, proteins, reduced this negative effect. Seed size, tannins and nutrient content of artificial seeds all affected various stages of the seed dispersal process by rodents, with size and nutrients tending to favour dispersal, and tannins disfavoured dispersal (Wang et al. 2013). In contrast, artificial seed experiments with *Dasyprocta punctata* in Costa Rica suggest

size but not tannin concentration affect seed dispersal decisions (Kuprewicz and García-Robledo 2019). Furthermore, artificial seeds with different characteristics were cached in different microhabitats, which can affect quality of dispersal; larger and more nutritious seeds were most likely to be cached under shrubs (Wang and Corlett 2017).

Morphology

Here we consider forms of morphological variation beyond fruit and seed size that might influence interindividual variation in dispersal. The most apparent cases of morphology driving interindividual variation in seed dispersal are with heterocarpic species that produce diaspores of two or more distinct morphologies differing in dispersal ability. For example, in the annual grass *Bromus tectorum* plants produce caryopses with (complex) and without (simple) sterile florets attached, which differ in dispersal ability because complex diaspores attach better to animal fur (Monty et al. 2016). Numerous plant species produce dimorphic, soft (non-dormant) and hard (dormant) seeds (Baskin and Baskin 2014). Paulsen et al. (2013, 2014) argued that this strategy enables individual plants to benefit both from the antipredation advantages of hard seeds and the dispersal advantages of soft seeds. Hard seeds emit fewer volatiles than soft seeds and are more difficult to detect by granivores such as rodents that rely on olfaction. Because granivores act both as dispersers and predators, detection might be advantageous, but can also result in seed consumption. In heterocarpic species, the relative proportions of different propagule types often vary with environmental conditions. For example, increasing stress can result in either an increase (Imbert and Ronce 2001; Martorell and Martínez-López 2014) or a decrease (Mandák and Pyšek 1999; Lu et al. 2013) in the production of more dispersible morphs. Interestingly, *Calendula arvensis* produce three distinct fruit morphs, one adapted to epizoochory, one to anemochory and one without adaptations for dispersal, although the extent of interindividual variation in the production of different morphs is unknown (De Clavijo 2005).

Beyond simply fruit or seed size, actual dimensions or shape can influence foraging decisions of endozoochorous and synzoochorous dispersers. For example, fruit diameter independent of length or overall mass is thought to be the most important metric of fruit size for frugivorous birds based on how fruits are swallowed (e.g. Wheelwright 1993). Acorn shape also influences preference by the European jay (*G. glandarius*); when diameter was held constant, jays preferred longer acorns; when length was constant, they chose wider acorns; and when mass was constant, they chose longer and slimmer acorns over shorter and wider acorns (Bossema 1979). How widespread such patterns are is unknown.

As noted previously, patterns of anemochorous dispersal are not driven solely by seed mass, but are influenced by the relationship between seed mass and the dispersal structure, which varies intraspecifically. It is generally thought that wing (coma, pappus) loading, frequently measured as fruit mass per unit surface area of the dispersal structure, is the major determinant of dispersal ability in wind-dispersed species. Thus, in trees (Sinha and Davidar 1992; Bhuyan et al. 2000; Debain et al. 2003), shrubs (Meyer and Carlson 2001) and forbs (Morse and Schmitt 1985; Donohue 1998; Skarpaas et al. 2011), increasing wing loading results in shorter dispersal distances or greater falling velocities, implying shorter dispersal distances. Other traits of propagules also lead to intraspecific variation in wind dispersal. Sheldon and Burrows (1973) argued that the fine details of pappus architecture influence dispersal more

than wing loading. In *Zygophyllum xanthoxylon*, a 'shape' index explained a small portion of the variation in dispersal distance (Zhu et al. 2016). For *Liatris scariosa* var. *novae-angliae*, dispersal distances were influenced by achene length (negatively), achene width (positively) and pappus length (positively), all of which differed among populations and among maternal families within populations, indicating likely interindividual variation in seed dispersal distances (Gravuer et al. 2003). *Carduus nutans* respond plastically to experimental drought with decreased wing loading and terminal velocity due to reductions in seed mass without changes in plume characteristics; interestingly, they also showed decreased intraindividual variation in terminal velocity (Teller et al. 2014).

The role of morphological variation in seed dispersal potential for other dispersal modes is poorly understood. In epizoochory, morphological variation has been examined almost exclusively interspecifically (e.g. Tackenberg et al. 2006; Will et al. 2007; Hovstad et al. 2009; Albert et al. 2015). However, limited evidence exists that seeds of epizoochorous species also vary intraspecifically in number, size and shape of appendages and in attachment potential (Gorb and Gorb 2002). We are aware of only one relevant study on hydrochory, where intraspecific variation in the size of the aeriferous mesocarp layers in *Scaevola crassifolia* fruits affected buoyancy (Guja et al. 2014).

Colour polymorphism

Some fleshy-fruited species exhibit fruit colour polymorphism, producing two or more colour morphs, sometimes on different plants and sometimes on the same plant (e.g. Willson and O'Dowd 1989). Selection of particular colour morphs ranges from relatively strong (e.g. *Rubus spectabilis*, Gervais et al. 1999) to weak (e.g. *Rhagodia parabolica*, Willson and O'Dowd 1989) to non-existent (e.g. *M. communis*, Traveset et al. 2001). Selection can be consistent across large geographic areas (e.g. *R. spectabilis*, Gervais et al. 1999) or vary across years and populations (e.g. *Acacia ligulata*, Whitney 2005) or even among individuals of a disperser species (e.g. various birds dispersing *R. spectabilis*, Traveset and Willson 1998). In the only study we know addressing selection among individual plants, the deer *M. muntjak* preferred *C. axillaris* trees with yellow fruits over those producing yellowish-green fruits (Chen et al. 2001). The basis of colour morph selection is unclear, but there is no evidence we are aware of that in colour polymorphic systems morphs differ in size, pulp:seed ratio or major nutrients (Willson and O'Dowd 1989; Traveset and Willson 1998).

Phenology

Interindividual variation in fruiting phenology is widespread in herbaceous forbs (Collier and Rogstad 2004), shrubs (SanMartin-Gajardo and Morellato 2003) and trees (Howe and Vande Kerckhove 1979; Franklin and Bach 2006; Muhanguzi and Ipulet 2012), but this variation may or may not affect seed dispersal. In *C. corymbosa*, later fruiting trees were visited by more species, but this had no real effect on dispersal because the additional frugivore species ate very few fruits (Howe and Vande Kerckhove 1979). *Olea europaea* var. *sylvestris* individuals that ripened fruit earlier were favoured in 1 of 2 years (Alcántara et al. 1997), while *O. tenera* trees ripening fruits earlier had greater and more rapid fruit removal than late-ripening trees (Wheelwright 1993). In *Q. serrata*, acorns produced later in the season were larger with lower tannin concentrations, making them more valuable food items (Takahashi et al. 2011). Phenological variation potentially affects dispersal quality as well. Although not linked to

individual plant fruiting phenology, González-Varo et al. (2019) demonstrated that dispersal quantity and quality changed through the fruiting season of the bird-dispersed *Pistacia lentiscus*.

Extrinsic Variation: Ecological Context

Fruiting neighbourhood

The presence, species identity, density and relative desirability of co-occurring fruiting neighbours can influence interindividual variation in seed dispersal. Some argue that trees compete with neighbours for dispersal and that intraspecific competition should be more intense than interspecific competition, especially in the tropics where it was thought there was little overlap in dispersal assemblages across species (Howe and Estabrook 1977). However, facilitation of dispersal by neighbours is also a possibility if the collective lure of multiple fruiting trees disproportionately attracts dispersers (Morales et al. 2012). In fact, this scenario has been proposed as another driver of hub and non-hub dispersal networks since frugivores are assumed to choose high-quality patches to forage in without considering the number of trees contributing to that patch (Carlo et al. 2007).

As expected, intraspecific competition has been found to reduce the quantity of dispersal of many tropical and temperate trees and shrubs (Table 4). However, many studies also have found intraspecific facilitation of the quantity of seed dispersal across taxa and ecosystems (Table 4). Thus, both intraspecific competition and facilitation have been found to affect the quantity of dispersal in both tropical and temperate systems, but given the relative scarcity of empirical work and its bias to tropical systems, general patterns are not clear. In a multispecies comparison, species fruiting in high densities were more likely to have dispersal reduced by neighbours (competition), whereas species fruiting in low density were more likely to have dispersal increased by neighbours (facilitation), a logical expectation (Albrecht et al. 2015). Further, whether competition or facilitation of dispersal by conspecific fruiting neighbourhoods occurs can be affected by the heterospecific fruiting neighbourhood (Rumeu et al. 2019).

Heterospecific fruiting neighbourhoods might also influence interindividual variation in dispersal quantity and quality, although we have even less empirical evidence for heterospecific than conspecific interactions, and outcomes appear to be complex. Dispersal of the tropical tree *Eugenia uniflora* was unaffected by heterospecific fruiting neighbourhoods (Blendinger and Villegas 2011). Similarly, *Solanum americanum* in monospecific patches and in mixed patches with *Cestrum diurnum* did not differ in the quantity of seeds dispersed; however, in the presence of *C. diurnum*, *S. americanum* seeds were dispersed in smaller seed loads among more defecations, resulting in reduced potential competition and increased number of sites occupied (Carlo 2005). In the south-eastern USA, the native shrub *Morella cerifera* marginally facilitated the dispersal of the invasive shrub *Triadica sebifera* and improved its germination, but inhibited seedling growth (Battaglia et al. 2009). In California oak woodlands, the high-quality disperser California scrub-jay (*Aphelocoma californica*) responded numerically and functionally to *Quercus lobata* with large acorn crops when the dominant *Q. douglasii* had low acorn production, but not when *Q. douglasii* produced abundant acorns (Pesendorfer and Koenig 2017). By contrast, the seed predator acorn woodpecker (*Melanerpes formicivorus*) had a constant response to *Q. lobata* trees independent of background

Table 4. Examples of studies reporting intraspecific competition for dispersers (i.e. reduction in dispersal quantity by conspecific neighbours), intraspecific facilitation of dispersal (i.e. increase in dispersal caused by conspecific neighbours) or no effect of neighbours including a variety of life forms in tropical and temperate regions.

Species	Description	Result	Reference
<i>Schefflera morototoni</i>	Tropical tree	Competition	Saracco et al. (2005)
<i>Viola nobilis</i>	Tropical tree	Competition	Manasse and Howe (1983)
<i>Viola surinamensis</i>	Tropical tree	Competition	Moreira et al. (2017)
<i>Erythroxylum havanense</i>	Tropical shrub	Competition	Gryj and Domínguez (1996)
<i>Attalea butyracea</i>	Tropical palm	Competition	Jansen et al. (2014)
<i>Sambucus pubens</i>	Temperate shrub	Competition	Denslow (1987)
<i>Viburnum recognitum</i>	Temperate shrub	Competition	Smith and McWilliams (2014)
<i>Viburnum dentatum</i>	Temperate shrub	Competition	Smith and McWilliams (2014)
<i>Eugenia uniflora</i>	Tropical tree	Facilitation	Blendinger and Villegas (2011)
<i>Miconia fosteri</i>	Tropical shrub	Facilitation	Blendinger et al. (2008)
<i>Miconia irwinii</i>	Tropical shrub	Facilitation	Guerra et al. (2017)
<i>Geonoma pauciflora</i>	Tropical palm	Facilitation	Pizo and Almeida-Neto (2009)
<i>Tristerix corymbosus</i>	Temperate mistletoe	Facilitation	Morales et al. (2012)
<i>Juniperus communis</i>	Temperate shrub	Facilitation	García et al. (2001)
<i>Quercus ilex</i>	Temperate tree	Facilitation	Morán-López et al. (2015a)
<i>Miconia serrulata</i>	Tropical shrub	No effect	Blendinger et al. (2008)

acorn production. Consequently, *Q. lobata* trees received high-quality dispersal when *Q. douglasii* acorns were sparse, but little dispersal and extensive seed predation when *Q. douglasii* acorns were abundant. Synzoochorous foragers collect seeds both for current consumption and future use, but preferences often differ between consumed and cached items (Lichti et al. 2017), opening the potential for complex, indirect seed–seed interactions. For example, one seed species could provide a preferred short-time food supply and therefore subsidize caching of another species more suitable for long-term storage. Such ‘apparent predation’ (sensu Lichti et al. 2014) was documented between *Quercus robur* and *Q. rubra* in Poland (Bogdziewicz et al. 2019).

Finally, fruiting neighbourhoods can affect dispersal quality as well as quantity. With both endozoochorous and synzoochorous dispersal, higher density fruiting neighbourhoods have been shown to result in shorter dispersal distances (Carlo and Morales 2008; Morales et al. 2012; Jansen et al. 2014).

Habitat structure: broader aspects

Beyond fruiting neighbourhoods, effects of other aspects of habitat structure on seed dispersal have been addressed from local within-patch variation in structure to landscape-scale variation. Here we give a brief overview from the perspective of interindividual variation in seed dispersal, emphasizing smaller scale population-level variation, with scale defined by the dispersal agent. Note that the drivers and consequences of habitat effects on interindividual variation in seed dispersal operate at much larger spatial scales for plants that are dispersed by mobile animals than for those dispersed abiotically.

At the scale of meters to tens of meters, the distances of *P. mahaleb* individuals from nests and rock outcrops affected the composition of the avian disperser assemblages foraging on those plants (Fuentes et al. 2001). Similarly, but at a larger spatial scale, three nearby stands with vegetation differing in vertical structure and species composition differed substantially in seed disperser assemblages foraging on *P. mahaleb* (Gutián et al. 1992). In both cases, differences in assemblages resulted in differences in the quantity of dispersal, the handling of fruits and seeds, and the microhabitat destination of dispersed seeds. In a highly heterogeneous forest, *C. monogyna* individuals growing with dense tree cover dispersed more seeds and over longer distances than did individuals growing with more sparse cover (Herrera

et al. 2011). Similarly, with greater amount and continuity of forest cover, the carnivore *Martes foina* dispersed seeds longer distances (Herrera et al. 2016). Lastly, *Corema album* seeds were dispersed by the same three species in three adjacent habitat patches varying in vegetation structure, but all three species exhibited among-habitat variation in both the quantity and quality of dispersal (Calviño-Cancela and Martín-Herrero 2009).

More discrete habitat patchiness can also drive interindividual variation in dispersal quantity and quality. In *O. europaea* var. *sylvestris*, genetic information on avian dispersers and seed parents revealed major differences in dispersal for trees in remnant forest stands versus isolated trees in adjacent agricultural fields (González-Varo et al. 2017), with forest and isolated trees differing substantially in the assemblage of birds dispersing their seeds and in the destinations of dispersed seeds. In continuous forest, *G. glandarius* dispersed more *Q. ilex* acorns, dispersed them further, and cached them in better microhabitats than when foraging in adjacent open dehesas with only scattered oaks, while within dehesas, trees close to forest or in spatial clumps were more likely to be dispersed (Morán-López et al. 2015a).

Numerous studies have addressed habitat fragmentation effects on seed dispersal. In the Amazonian tree *Duckeodendron cestroides*, dispersed by arboreal and terrestrial mammals, both the quantity and the mean and maximum distance of dispersal were greater in continuous forest than in fragments (Cramer et al. 2007). Similarly, for the bird-dispersed African tree *Leptonychia usambarensis*, compared to continuous forest, fragments had fewer species and individuals of seed dispersers, had fewer seeds removed and had seedlings located closer to parents (Cordeiro and Howe 2003). Fragmentation combined with hunting led to the loss of larger-gaped dispersers and a reduction in seed dispersal of larger fruits, resulting in rapid evolution of reduced fruit and seed size in *Euterpe edulis* (Galetti et al. 2013). Fragmentation can also impact synzoochorous dispersal. In *Astrocaryum aculeatum*, decreasing forest patch area was associated with a higher quantity of dispersal (increased seed removal rate), but lower quality of dispersal (reduced caching and reduced dispersal distances), likely due to changes in rodent community composition (Jorge and Howe 2009). Similar findings have been found in *Q. ilex* dispersed by *Apodemus sylvaticus* (Morán-López et al. 2015b, 2018b). However,

fragmentation does not always negatively affect seed dispersal; forest fragmentation in Poland reduced the number of larger frugivores without decreasing fruit removal (Farwig et al. 2017).

A meta-analysis of primarily tropical fleshy-fruited species (Markl et al. 2012) suggested that fragmentation does not affect visitation or seed removal rates, and only marginally reduces dispersal distances. By contrast, a meta-analysis of a worldwide data set (Fontúrbel et al. 2015) suggested that fragmentation reduced interaction rates (visitation or fruit removal), but not disperser diversity (abundance or species richness); at a major biome level, fragmentation reduced disperser diversity only in temperate zones but reduced interaction rates in both temperate and tropical zones. Additionally, inter- or intraspecific variation in disperser traits such as movement distance, movement frequency and gut retention time of seeds represent one mechanism explaining how fragmentation can positively or negatively affect dispersal distances (fragment entrapment, Jones et al. 2017).

Beyond fragmentation, habitat disturbance, degradation and simplification can impact dispersal quantity and quality. In oaks (*Quercus velutina* and *Q. alba*), timber harvest resulted in 67 % reduction in SDE by rodents, probably due to increased vegetation cover facilitating recovery of cached acorns (Kellner et al. 2016). In a Mediterranean system, habitat degradation reduced the abundance, species richness and movement of avian dispersers, resulting in reduced fruit removal, seed dispersal distances, seed survival and seedling success (Rey and Alcántara 2014). Other studies have shown that increasing forest disturbance can result in decreased likelihood of plants being visited by dispersers (Lehouck et al. 2009; Moreira et al. 2017), as well as decreased species richness of dispersers and reduced dispersal distances (Chatterjee and Basu 2015) and of rates of seed dispersal (Lehouck et al. 2009). These results are compatible with meta-analyses showing degradation to have a greater negative impact than fragmentation on seed dispersal (Markl et al. 2012), and habitat degradation generally reducing abundance and diversity of dispersers (Fontúrbel et al. 2015).

Habitat structure also can impact seed dispersal by wind, by altering the wind speed that initiates seed release, and by damping wind speeds. In *Taeniatherum caput-medusae* and *Tragopogon dubius*, taller surrounding vegetation reduced dispersal distances (Davies and Sheley 2007). Modelling suggests this should be common in herbaceous communities (Soons et al. 2004). Modelling further suggests that forest canopy height heterogeneity influences the likelihood of LDD; seeds released over shorter parts of the canopy encounter greater turbulence and are more likely to be ejected and experience LDD (Bohrer et al. 2008). Lastly, accelerated seed dispersal by wind along linear disturbances in the Canadian oil sands region has been reported (Roberts et al. 2018).

Topography

Movements of frugivorous birds are influenced by subtle variation in topographic relief, which can affect which individual fruiting trees are encountered during foraging and where seeds are deposited (Westcott 1994, 1997); this is likely true for other animal vectors as well. However, little empirical work directly addresses the role of topography in interindividual variation in seed dispersal. In Ecuador, contrasting results were found for two fleshy-fruited shrub species (Blendinger et al. 2008); *Miconia fosteri* on ridges had a greater number and proportion of fruits removed than did those at the bottom of slopes, while *M. serrulata* had a greater number but not proportion of fruits removed on slopes than on either ridges or at the bottom of slopes. For wind

dispersal, modelling suggests that even moderate topographic variability can have large impacts on variation in dispersal distances and directionality (Trakhtenbrot et al. 2014). Finally, slope steepness influences dispersal distances of heavy seeds, which are more likely to roll down downhill (e.g. oak acorns, Ohsawa et al. 2007), and the likelihood of seed dispersal via runoff (De Rouw et al. 2018).

Non-disperser animal communities

Individual plants might also differ in the quantity and quality of animal-mediated seed dispersal due to the actions of third-party players. Predators can indirectly affect seed dispersal through their effects on risk-sensitive foraging of dispersers. Some of these effects are mediated by vegetation structure, with plants in more open and risky places receiving fewer visits by dispersers (Iida 2004; McCabe and Olsen 2015; Kellner et al. 2016). In other situations, animals respond to olfactory, visual or acoustic predator cues, leading to reduced seed removal rates in frugivorous birds (Breviglieri and Romero 2016; Tella et al. 2016; Shave et al. 2018), bats (Breviglieri et al. 2013) and granivorous rodents (Sunyer et al. 2013). In addition, rodents are sensitive to ungulate presence because of trampling risk or disturbance by rooting (e.g. by wild boar *Sus scrofa*); in *Q. ilex*, the presence of ungulates was associated with lower quality seed dispersal by rodents (lower proportion of seeds cached and not recovered) and changes in caching sites (reduced caching under shrubs) (Muñoz and Bonal 2007). Finally, responses to predators and competitors can interact with other traits, such as the presence and concentration of deterrents (McArthur et al. 2012).

Insects frequently infest fruit pulp, seeds or dispersal structures, which can affect seed dispersal. Howler monkeys (*Alouatta caraya*) preferentially feed in *Ocotea diospyrifolia* trees with high fruit infestation by curculionids and low fruit infestation by moths (Bravo 2012). The seed parasitoid wasp *Macrodasyrceras hirsutum* reduces attractiveness of *Ilex integra* berries to frugivorous birds through 'colour manipulation'; infested fruits are less likely to ripen and turn red, decreasing the risk that the fruits will be eaten and wasps killed (Takagi et al. 2012). In synzoochorous dispersal, seed infestation can increase the probability of rejection (Bossema 1979; Muñoz and Bonal 2008b) or of immediate consumption (Steele et al. 1996; Perea et al. 2012), but generally reduces caching rates (Steele et al. 1996; Perea et al. 2012), thus decreasing dispersal quality. However, not all scatter hoarders discriminate between infested and sound seeds, particularly before insect emergence (Gálvez and Jansen 2007; Cheng and Zhang 2011). Note that these synzoochorous examples are based on responses to individual seeds and it is unknown to what degree they translate into selection among trees differing in infestation levels. Insect attack also affects anemochory. For example, *Rhinocyllus conicus* larvae feeding on *C. nutans* receptacles induce callus formation, resulting in inhibited seed release, shortened pappus filaments and reduced dispersal distances (Marchetto et al. 2014). In turn, insect infestation is often affected by masting (Espelta et al. 2008, but see Bogdziewicz et al. 2018), thereby creating another, indirect pathway through which temporal variation in seed output can affect seed dispersal.

Lifetime Fitness: Temporal Complexity

Most of what we know about intraspecific variation in seed dispersal represents a snapshot in time—a frame or two in a potentially long movie of life. While these frames might

accurately represent the fitness outcomes for an annual plant, the majority of plants discussed in this review are long-lived perennials that are interacting with an extremely dynamic world where both intrinsic and extrinsic factors vary through time. Although we are not in a position to evaluate the overall consequences of this variation, it is important to acknowledge the variation exists.

Fruit crop sizes vary between years. Sometimes this variation is relatively subtle and driven by such factors as resource availability or climatic conditions (Jordano 1987; Jordano and Schupp 2000; Wenk and Falster 2015; Davi et al. 2016). Sometimes the variation is extreme, as seen in masting species (Herrera et al. 1994; Wenk and Falster 2015; Davi et al. 2016; Pearse et al. 2016). Different dispersal kernels are necessary to capture mast versus non-mast years (Martínez and González-Taboada 2009), with potentially greater LDD when acorn density is low (Moran and Clark 2012). The fitness impacts of this variation should depend at least partially on how synchronous fruit crop size variation is in the population and community. Fruit crop size also varies over longer, ontogenetic time scales; crop sizes increase with perennial plant age and size, often plateauing at some point and remaining relatively constant until death, sometimes showing declines with senescence late in life (Davi et al. 2016).

Many other intrinsic traits relevant to intraspecific variation in dispersal are temporally dynamic. Fruit/seed size, and most likely such traits as pulp:seed ratio, vary across years (González-Varo and Traveset 2016). Plant height increases ontogenetically (Coopman et al. 2008). Rewards (Lotan and Izhaki 2013) and deterrents (Tsahar et al. 2002) can change from year to year and in some cases even seasonally.

Temporal variation in extrinsic factors, or the ecological context, is perhaps even more extreme. Fruiting neighbourhoods can change from year to year as different individuals and species respond differently to changing resources and climate (Jordano and Schupp 2000). Other aspects of habitat structure around individual plants can change through time due to successional processes and demographic processes (Herrera et al. 1994), as well as anthropogenic impacts (Markl et al. 2012; Fontúrbel et al. 2015). Lastly, interactions with non-disperser animal communities can vary greatly from year to year as a function of, among other drivers, changes in individual crop sizes and in fruiting neighbourhoods, and population fluctuations of other interacting animal species (Schmidt and Ostfeld 2003).

Complexity

Although many exceptions exist, much work on intraspecific variation in seed dispersal has taken a more or less univariate approach; for example, the impact of fruit crop size, fruit size or plant height on dispersal. Alternatively, some address multiple traits affecting dispersal and quantify the relative importance of each and the presence or absence of interactions. In a recent study using an individual-plant-based network analysis of frugivory, locations of individual *H. succosa* trees within the network were determined by a combination of plant height, fruit size and sugar concentration, with shorter individuals with larger fruits and intermediate sugar concentration being most central (Crestani et al. 2019). Nonetheless, the true complexity of dispersal is often overlooked. In this review we have also taken primarily a univariate approach, which we argue has value, especially at our early stage of understanding the drivers of interindividual variation in seed dispersal. However, it is critical to understand that we do not believe that this is really how the world exists.

We noted the difficulties of knowing what animal seed dispersers base their harvesting decisions on when so many potentially important traits co-vary: fruit size, absolute and relative quantity of reward, seed number and size, nutrients, toxins and more (Jordano 1984; Izhaki et al. 2002). For example, do frugivores select fruits to harvest based on size *per se* or on the underlying variation in pulp:seed ratio (Howe and Vande Kerckhove 1981; Martínez et al. 2007; Sobral et al. 2010)? Such complexities surely exist in other dispersal systems as well. For example, in anemochorous plants, the size of the dispersal structure increases with seed mass, but generally not sufficiently to maintain a constant wing loading (Meyer and Carlson 2001; Debain 2003; Skarpaas et al. 2011; but see Wyse et al. 2019). Co-variation of seed release height and seed terminal velocity (Teller et al. 2014), and of abscission force and terminal velocity (Teller et al. 2015) have also been reported. It is highly likely that co-variation of traits relevant to seed dispersal is as extensive with wind dispersal as with frugivory.

Complexity also arises in animal-dispersed species because foraging animals often make foraging decisions hierarchically (Côtés and Uriarte 2013). For example, foraging frugivores must first select the foraging patch, then choose the individuals to feed in, and then choose which fruits to harvest from that plant. In addition, multiple cues may be used hierarchically at any single stage of this process. For example, experiments with the large fleshy-fruited shrub *C. monogyna* elegantly demonstrated hierarchical selection by *Turdus migratorius* of individual trees in which to forage. First, birds preferred trees with larger crop sizes, but if crop sizes were constant, they preferred plants with larger fruits, and, finally, if fruit size was constant, they preferred plants with greater pulp:seed ratios (Sallabanks 1993).

Understanding variation in seed dispersal is further complicated by the concomitant interindividual variation in seed dispersers, including sexual dimorphism, ontogenetic changes, interindividual variation in specialization and unique animal personalities (Zwolak 2018). For example, our discussion of fruit size variation in *M. communis* and its effect on fruit availability to different seed dispersers was based on measured intraindividual and interindividual variation in fruit diameters but only mean gape width for the dispersal agents (González-Varo and Traveset 2016). Interpretations could be different if interindividual variation in the seed disperser species was also incorporated. More generally, interindividual variation in plants and dispersers interact and it might be difficult to understand one without understanding the other (Côtés and Uriarte 2013). Plants almost certainly respond at the individual level to variation in how seed dispersers interact with them; these eco-evolutionary feedbacks mean that intraspecific variation is important in both sides of the interaction, perhaps even intensifying the individual-level variation in both players (compare with Siepielski and Benkman 2010).

Further complexity is likely in particular dispersal systems, such as for example with diplochorous dispersal, where dispersal is accomplished by a sequence of steps that involve different dispersal agents such as primary dispersal by a frugivorous bird and secondary dispersal by a rodent (Vander Wall and Longland 2004). We predict that, all else being equal, diplochorous dispersal systems would have even greater interindividual variation in seed dispersal success than non-diplochorous systems given that variability arising during the second phase of dispersal is building on variability created during the first phase of dispersal. For example, as discussed previously, intraspecific variation in seed size can affect selection by both frugivorous birds and rodents, sometimes in the same way and sometimes not.

Lastly, all of these trait-based dispersal drivers are playing out in an extraordinarily heterogeneous environment, varying continuously in habitat structure, fruiting neighbourhoods, wind conditions and more.

Where Are We Now and Where Do We Need to Go?

While we show substantial evidence that drivers of intraspecific variation in seed dispersal are diverse and pervasive, we also reveal large gaps in our understanding, partly due to a paucity of research directly addressing intraspecific, especially interindividual, variation in seed dispersal, and partly due to the complexity of interactions among drivers. Our understanding is limited further by the existing empirical work's focus on the quantity of seed dispersal, with much less consideration of the quality of dispersal or LDD. Of particular interest are the intrinsic trait-based drivers that can respond to natural selection. The best-supported and best-understood intrinsic driver of interindividual variation in seed dispersal is crop size; with more seeds produced, more seeds are dispersed. Crop size is also likely the most widespread driver, being relevant to most if not all forms of dispersal. Though less well supported and less well understood, fruit/seed size is likely the second most widespread intrinsic driver. Again, it seems to be relevant to a broad range of seed dispersal modes. However, when it comes to animal-mediated dispersal we do not have a good understanding of the ultimate cause of size-based fruit or seed selection—is it fruit/seed size *per se*, or some co-varying trait such as pulp:seed ratio? Remaining intrinsic drivers are even more poorly understood, though apparently range from widespread but weak, such as plant height, to sporadic and variable in strength, such as colour polymorphism. For extrinsic drivers, a variety of studies have addressed the impact of fruiting neighbourhoods on interindividual variation in seed dispersal, but we do not understand well when to expect competition for dispersers and when to expect facilitation of dispersal. With respect to habitat structure, much relevant work has been from the perspective of anthropogenic impacts of habitat fragmentation and degradation on seed dispersal rather than from the perspective of interspecific variation in seed dispersal.

Beyond limited empirical work, we are further hindered by an even greater lack of theory related to the drivers of intraspecific, especially interindividual variation in seed dispersal. While there have been some theoretical developments around fruit crop size and seed dispersal success (see earlier discussion of [Howe and Estabrook 1977](#); [Carlo et al. 2007](#)), we are aware of no other developed theory that can guide our understanding of the drivers of interindividual variation in dispersal and potential demographic and evolutionary responses to such variation.

Looking forward towards potential research directions, in [Box 2](#) we highlight a selection of outstanding questions concerning intrinsic drivers of intraspecific variation in seed dispersal that we personally believe to be especially informative and intriguing to answer. We present these questions as a starting point to advance our understanding of intraspecific drivers of seed dispersal. One promising approach to answer these questions and disentangle the complexity inherent in intraspecific seed dispersal is a frugivore-centred modelling approach ([Côrtes and Uriarte 2013](#)). This approach advocates parameterizing field data on intrinsic animal factors and behaviour, as well as extrinsic landscape factors, to test and quantify the strength of the variables affecting the spatially explicit deposition of seeds across the landscape ([Côrtes and Uriarte 2013](#)). Mechanistic simulations can be used in a hierarchical manner to test the effect of multiple factors one at a time, to quantify their relative influence on patterns of seed deposition ([Côrtes and Uriarte 2013](#)). Studies using this approach have successfully quantified the impact on seed dispersal of edge-following behaviour in a fragmented landscape ([Levey et al. 2005](#)), fruiting neighbourhoods ([Carlo and Morales 2008](#)) and drivers of reduced LDD ([Uriarte et al. 2011](#)). Although primarily envisioned to study endozoochory, similar methods have been applied to epizoochory ([Will and Tackenberg 2008](#)) and other dispersal modes by considering relevant intrinsic and extrinsic factors (e.g. anemochory, [Nathan et al. 2001](#)). Additionally, a powerful molecular approach that matches individual seeds or seedlings to maternal plants ([Grivet et al. 2009](#)) across dispersal modes is also promising for studying individual variation in seed dispersal and may compliment simulation modelling approaches. Despite the daunting complexity of drivers of intraspecific variation in seed dispersal, the combination of

Box 2. Questions

There is an abundance of questions that remain to be answered. Here we highlight a selection of outstanding questions concerning intrinsic drivers of intraspecific variation in seed dispersal that we personally believe to be especially informative and intriguing to answer.

General

- How strong and widespread are the major drivers of intraspecific variation in seed dispersal? Do their relative strengths differ across dispersal modes, and to what extent do drivers operate independently versus interactively?
- What is the relative contribution of intraindividual versus interindividual variation in traits to variation in seed dispersal patterns and SDE?
- How variable is the extent of intraindividual variation in dispersal traits and dispersal patterns, both within and among populations? Do the answers to these questions depend on dispersal mode?

Plant–Animal Dispersal Mutualisms

- When and to what extent do animal seed dispersers respond to intraindividual versus interindividual variation in fruit or seed traits? To what extent do seed dispersers respond to interindividual mean versus variance in fruit or seed traits?
- How does interindividual variation in plant traits interact with interindividual variation in seed disperser traits to affect patterns of seed dispersal and SDE?

quantitative approaches and tools available provide ample starting points to answer the questions we pose in **Box 2** and improve our understanding of this important aspect of seed dispersal.

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Contributions by the Authors

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Conflict of Interest

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