

Point of View**SPECIAL ISSUE: The Role of Seed Dispersal in Plant Populations: Perspectives and Advances in a Changing World****Employing plant functional groups to advance seed dispersal ecology and conservation**

Clare Aslan^{1*}, Noelle G. Beckman², Haldre S. Rogers³, Judie Bronstein⁴, Damaris Zurell⁵, Florian Hartig⁶, Katriona Shea⁷, Liba Pejchar⁸, Mike Neubert⁹, John Poulsen¹⁰, Janneke HilleRisLambers¹¹, Maria Miriti¹², Bette Loiselle¹³, Edu Effiom¹⁴, Jenny Zambrano¹⁵, Geno Schupp², Gesine Pufal¹⁶, Jeremy Johnson¹⁷, James M. Bullock¹⁸, Jedediah Brodie¹⁹, Emilio Bruna¹³, Robert Stephen Cantrell²⁰, Robin Decker²¹, Evan Fricke³, Katie Gurski²², Alan Hastings²¹, Oleg Kogan²³, Onja Razafindratsima²⁴, Manette Sandor²⁵, Sebastian Schreiber²¹, Rebecca Snell²⁶, Christopher Strickland²⁷ and Ying Zhou²⁸

¹Landscape Conservation Initiative, Northern Arizona University, Flagstaff, AZ 86011, USA

²Department of Biology, Utah State University, 5305 Old Main Hill, Logan, UT 84322, USA

³Department of Ecology, Evolution, and Organismal Biology, Iowa State University, 251 Bessey Hall, Ames, IA 50011, USA

⁴Department of Ecology and Evolutionary Biology, University of Arizona, 1041 E. Lowell Street, Tucson, AZ 85721, USA

⁵Dynamic Macroecology, Landscape Dynamics, Swiss Federal Research Institute WSL, Zürcherstrasse 111, CH-8903 Birmensdorf, Switzerland

⁶Faculty of Biology and Pre-Clinical Medicine, University of Regensburg, Universitätsstraße 31, D-93053 Regensburg, Germany

⁷Department of Biology, Pennsylvania State University, 208 Mueller Laboratory, University Park, PA 16802, USA

⁸Department of Fish, Wildlife and Conservation Biology, Colorado State University, 1474 Campus Delivery, Fort Collins, CO 80523, USA

⁹Biology Department, Woods Hole Oceanographic Institution, 266 Woods Hole Road, Woods Hole, MA 02543, USA

¹⁰Nicholas School of the Environment, Duke University, 9 Circuit Drive, Durham, NC 27708, USA

¹¹Department of Biology, University of Washington, Seattle, WA 98195, USA

¹²Department of Evolution, Ecology and Organismal Biology, The Ohio State University, 318 W 12th Avenue, Columbus, OH 43210, USA

¹³Department of Wildlife Ecology and Conservation, University of Florida, Gainesville, FL 32611, USA

¹⁴CRS Forestry Commission, Calabar, Nigeria

¹⁵National Socio-Environmental Synthesis Center, 1 Park Place, Annapolis, MD 21401, USA

¹⁶Naturschutz & Landschaftsökologie, Albert-Ludwigs-Universität Freiburg, Tennenbacher Str. 4, D-79106 Freiburg, Germany

¹⁷Department of Geography, Texas A&M University, College Station, TX 77843, USA

¹⁸Centre for Ecology & Hydrology, Wallingford, UK

¹⁹Wildlife Biology Program, University of Montana, 32 Campus Drive, Missoula, MT 59812, USA

²⁰Department of Mathematics, University of Miami, 1365 Memorial Drive, Coral Gables, FL 33146, USA

²¹University of California-Davis, Davis, CA 95616, USA

²²Department of Mathematics, Howard University, Washington, DC 20059, USA

²³Physics Department, California Polytechnic State University, 1 Grand Avenue, San Luis Obispo, CA 93407, USA

²⁴Department of Biology, College of Charleston, Charleston, SC 29424, USA

²⁵Department of Ecology and Evolutionary Biology, University of Connecticut, Storrs, CT 06269, USA

²⁶Environmental and Plant Biology, Ohio University, Athens, OH 45701, USA

²⁷Department of Mathematics, University of Tennessee, Knoxville, TN 37996, USA

²⁸Department of Mathematics, Lafayette College, 730 High Street, Easton, PA 18042, USA

Received: 15 August 2018 **Editorial decision:** 24 January 2019 **Accepted:** 5 February 2019 **Published:** 7 February 2019

Associate Editor: Kim McConkey

*Corresponding author's e-mail address: clare.aslan@nau.edu

© The Author(s) 2019. Published by Oxford University Press on behalf of the Annals of Botany Company.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (<http://creativecommons.org/licenses/by/4.0/>), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

Citation: Aslan C, Beckman NG, Rogers HS, Bronstein J, Zurell D, Hartig F, Shea K, Pejchar L, Neubert M, Poulsen J, HilleRisLambers J, Miriti M, Loiselle B, Effiom E, Zambrano J, Schupp G, Pufal G, Johnson J, Bullock JM, Brodie J, Bruna E, Cantrell RS, Decker R, Fricke E, Gurski K, Hastings A, Kogan O, Razafindratsima O, Sandor M, Schreiber S, Snell R, Strickland C, Zhou Y. 2019. Employing plant functional groups to advance seed dispersal ecology and conservation. *AoB PLANTS* **11**: plz006; doi: 10.1093/aobpla/plz006

Abstract. Seed dispersal enables plants to reach hospitable germination sites and escape natural enemies. Understanding when and how much seed dispersal matters to plant fitness is critical for understanding plant population and community dynamics. At the same time, the complexity of factors that determine if a seed will be successfully dispersed and subsequently develop into a reproductive plant is daunting. Quantifying all factors that may influence seed dispersal effectiveness for any potential seed-vector relationship would require an unrealistically large amount of time, materials and financial resources. On the other hand, being able to make dispersal predictions is critical for predicting whether single species and entire ecosystems will be resilient to global change. Building on current frameworks, we here posit that seed dispersal ecology should adopt plant functional groups as analytical units to reduce this complexity to manageable levels. Functional groups can be used to distinguish, for their constituent species, whether it matters (i) if seeds are dispersed, (ii) into what context they are dispersed and (iii) what vectors disperse them. To avoid overgeneralization, we propose that the utility of these functional groups may be assessed by generating predictions based on the groups and then testing those predictions against species-specific data. We suggest that data collection and analysis can then be guided by robust functional group definitions. Generalizing across similar species in this way could help us to better understand the population and community dynamics of plants and tackle the complexity of seed dispersal as well as its disruption.

Keywords: dependency; directed dispersal; dispersal vectors; generalization; mutualism; seed dispersal effectiveness.

Introduction: Seed Dispersal Is Fundamental to Populations and Communities, Yet Complex

Plants rely on dispersal vectors—for example, animals, wind and water—to move across the landscape. We focus here on the dispersal of seeds, although many of the arguments we make could be generalized to other forms of dispersal. Dispersal occurs when a seed is moved from its origin and deposited elsewhere (Schupp *et al.* 2010). Through dispersal, plants may experience reduced exposure to competition, predation and parasitism (Janzen 1970; Connell 1971; Howe and Miriti 2004); colonize open habitats after disturbance (Wunderle 1997; Puerta-Piñero *et al.* 2013); reach potential suitable microsites in otherwise unsuitable landscapes (Wenny 2001); track climate fluctuations and environmental change (Corlett and Westcott 2013); and contribute to gene flow within and between populations (Bacles *et al.* 2006). As a result of these processes, seed dispersal is a fundamental driver of the diversity, structure, composition and spatial arrangement of plant communities. Seed dispersal ecology thus elucidates mechanisms of species coexistence, implications of species extinctions and impacts of global environmental change.

It is evident that a quantitative understanding of dispersal is key for predicting how environmental changes, and consequent changes in dispersal vectors, will impact plant populations and communities. Operationalizing this goal and moving seed dispersal ecology towards a predictive science, however, requires confronting a wide array of interacting factors and stochastic elements (Robledo-Arnuncio *et al.* 2014). Here, we discuss how a functional group approach may help simplify the complexity of seed dispersal ecology and boost our predictive capacity.

Functional group frameworks, in which species are categorized by ecological functions and the resulting groups treated as analytical units, have helped researchers confront complexity in other ecological sub-disciplines and have been tentatively explored in seed dispersal (e.g. Dennis and Westcott 2006; Brodie *et al.* 2009b; Bastazini *et al.* 2017). However, they have not yet been developed sufficiently to link empirical patterns of seed dispersal with theoretical predictions. In this Viewpoint, we discuss the complexity of seed dispersal and the need to reach generalities about it. We propose that to better understand the importance of seed dispersal in plant populations and communities, it would be useful to identify functional groups that distinguish plant species based on (i) how much it matters if their seeds are dispersed at all, (ii) how much it matters into what ecological context they are dispersed and (iii) how

much it matters by what vector they are dispersed. We list such functional groups and discuss their potential value in achieving general insights. We close by considering key knowledge gaps that this proposed functional group approach may address.

The complexity of seed dispersal

Due to their complexity, seed dispersal processes are difficult to quantify empirically (Fig. 1). Since the quantification of these processes forms the basis for understanding plant population and community dynamics, methods to reduce this complexity are essential. Both biotic and abiotic dispersal vectors can influence which seeds are dispersed, the risks and costs of dispersal, the spatial direction and distance that seeds travel, the probability that seeds will encounter specific microhabitats and the probability of seed aggregation (Howe and Miriti 2004; Côrtes and Uriarte 2013; Morales et al. 2013). For seeds transported by abiotic vectors, wind and water speeds and turbulence determine the distance and direction of seed movement (Katul et al. 2005; Nathan et al. 2011): not only are these factors intrinsically variable, but that variation interacts with the physical structure of the environment and the size and shape of the seed. In biotic dispersal, the set of disperser animals interacting with a seed may dictate its survival, growth and eventual reproduction (García and Martínez 2012). Dispersal vectors vary in their interactions with landscape structure, implying that the mechanism of dispersal may dictate the composition and arrangement of a plant community (Metzger 2000; Albrecht et al. 2012; Effiom et al. 2013; Razafindratsima and Dunham 2016; Chen et al. 2017). We largely focus on biotic seed dispersal because the

behaviours and physiology of biotic dispersers amplify the complexity of seed dispersal. Seed handling, for example, can affect the condition of the seed and change the likelihood of germination and subsequent survival and growth after seed deposition (Ladley and Kelly 1996; Traveset and Verdú 2002; Fricke et al. 2013). Some plant species exhibit extreme specialization in microhabitats and require dispersers to move seeds to these locations (e.g. desert mistletoe requires dispersal to the branches of a very limited range of host trees; Aukema 2004). The preferences and physiology of dispersers may influence the direction and distance of seed dispersal (Beckman and Rogers 2013) (Fig. 1). Stochastic events may include rare, long-distance dispersal events, which are difficult to observe and measure but can be critical for colonization of new geographic regions and provide connectivity among habitat patches across a landscape (Muller-Landau et al. 2003; Jordano et al. 2007; Shea 2007; Auffret et al. 2017). Behavioral aspects of biotic dispersers, such as local aggregation, social organization, mating system, competition and territoriality, can influence both spatial and temporal dispersal of seeds, with potential ramifications for seed aggregation and competition between seeds (reviewed in Karubian and Durães 2009). A given disperser may also disperse seeds of certain shapes or sizes, depending on disperser body or gape sizes (McConkey and Drake 2006; Muñoz et al. 2017). An extensive literature has explored the dispersal syndromes, or seed and fruit traits (e.g. size, shape, colour, chemistry, dormancy) that appear predictive of the primary dispersers of a given plant species, with investigation into the roles of co-evolution, secondary dispersal and specialization (e.g. Vander Wall and Beck 2012;

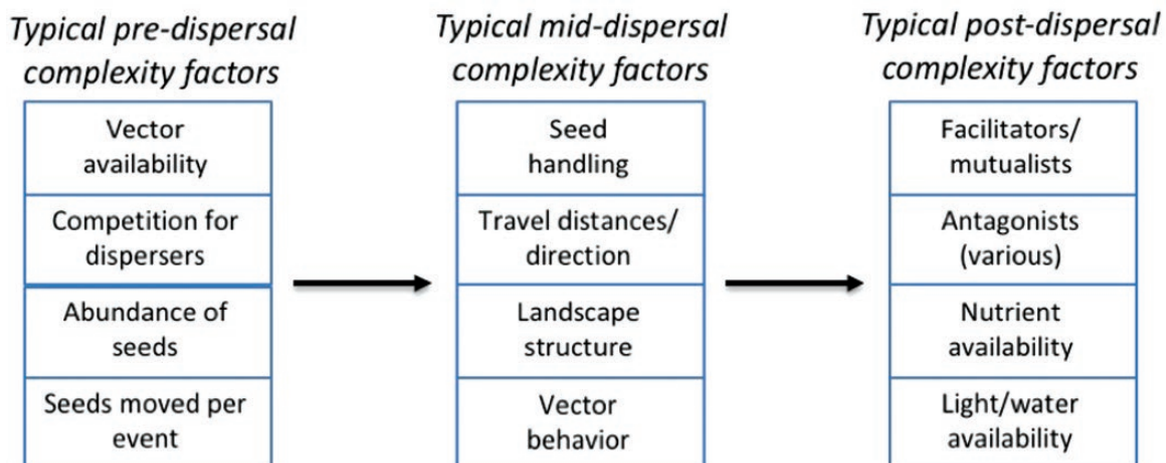


Figure 1. Seed dispersal exemplifies ecological complexity. Survival to adulthood and the fitness of individual adults are influenced by pre-, mid- and post-dispersal variables including the availability of abiotic and biotic vectors; the behaviours, preferences, morphology and physiology of dispersers; the spatio-temporal heterogeneity in seed deposition locations; and the probability of encountering other mutualists, facilitators, predators, pathogens and competitors following dispersal.

Howe 2016). Below, we briefly touch on the importance of dispersal syndromes as a form of functional grouping that categorizes dispersal adaptations. However, our proposed framework focuses instead on functional group delineations that distinguish the importance of dispersal in plant populations and communities.

Where and when a seed is deposited are clearly influenced by many sources of variability (Robledo-Arnuncio et al. 2014). Additionally, the spatial pattern of seed deposition on the landscape can then influence subsequent interspecific interactions (e.g. pollination, mycorrhizal associations, competition, predation, herbivory). Such interactions are important to the fitness of the newly established plant and determine the likelihood of survival and growth, access to limiting resources, the likelihood of mortality due to natural enemies and the probability of successful reproduction (Beckman and Rogers 2013). As a result of these interactions, the resulting plant community may more or less closely reflect the initial template established by seed deposition.

Seeking predictive capacity in light of global change: adapting current frameworks for functional groups

As established above, the plant community in a given location is constrained by the template established by seed deposition, but the post-deposition interactions within the seedscape (i.e. the full environmental context into which the seed is dispersed) determine which subset of those seeds succeeds. Empirically quantifying all relevant pre- and post-dispersal variables is a complex task for even one plant-disperser pair, and impossible for the thousands of species pairs that participate in seed dispersal mutualisms worldwide (Howe and Smallwood 1982; Aslan et al. 2013; Beckman and Rogers 2013). Nevertheless, without an attempt to understand these processes, their variability and the drivers of that variability, our understanding of system dynamics is hamstrung.

The Seed Dispersal Effectiveness (SDE) framework is a comprehensive framework to summarize the full suite of variables affecting the dispersal service provided to any particular plant species by any particular vector (Schupp et al. 2010). The SDE framework summarizes the contribution of each dispersal vector (whether biotic or abiotic) to the production of new adult plants by evaluating variables influencing the *quantity* of seeds dispersed and the *quality* of the seed dispersal event. Quantity metrics within SDE include, for example, the frequency of visits from the disperser to the plant species and the number of seeds dispersed per visit. Quality metrics include, for example, the condition of the deposited seed (which depends, e.g., on an animal's seed-handling behaviour) or a disperser's movement patterns

combined with measures of habitat quality across the landscape (Schupp et al. 2010). The SDE framework examines the complexity of variation within and among seeds, dispersers and other interactors determining the likelihood that a seed grows into a seedling (and, ultimately, reproductive adult), given a specific vector moving that seed to a specific location (e.g. Alvarez-Buylla and Martinez-Ramos 1990; Godinez-Alvarez and Jordano 2007; Escribano-Avila 2014; Rey and Alcántara 2014; Rother et al. 2016).

A full utilization of the SDE framework involves quantifying the effectiveness of dispersal for interacting pairs of seed and disperser species, taking into account pre-, mid- and post-dispersal factors that might affect seed survival and germination and the growth and fecundity of the resulting plant (e.g. Fig. 1). However, parameterization of SDE requires immense investment of empirical resources and includes up to 15 different measurable quantities for a given seed-disperser pair (Schupp et al. 2010). In one study, plant species were dispersed by an average of just over seven different disperser species (Aslan et al. 2013); parameterization of SDE for such a plant would therefore require a minimum of $7 \times 15 = 105$ separately measured parameters—a degree of complexity that would exhaust the resources of most scientific endeavours. Nevertheless, SDE has guided impressive efforts to measure subsets of these parameters, generating important insights. For example, McConkey et al. (2014) measured disperser effectiveness as a combination of the percent of monitored fruit dispersed by each disperser species combined with the distance of dispersal and survival of seedlings at each distance. Nogales et al. (2017) compared the number of seeds dispersed and effect of gut treatment by reptile vs. bird frugivores in the Galápagos. González-Castro et al. (2015) combined the number of seeds dispersed with condition of seeds after dispersal and seedling emergence/survival probabilities to compare SDE for birds and lizards. As these studies illustrate, different dispersers contribute in different ways to the template constraining an eventual plant community. To understand these roles across many more sites and for many more species, we require approaches that build off the SDE framework while simplifying the complexity inherent in biologically diverse systems.

To achieve this goal, we propose using plant functional groups in place of individual species in the SDE framework (Table 1). Functional groups are employed in many fields of ecology and have proven to be useful (e.g. functional group classifications yielded insights into plant species responses to climate change in Africa, Scheiter and Higgins 2009; successional dynamics in a Costa Rican forest, Chazdon et al. 2010; and global vegetation patterns, Sato et al. 2007). By identifying relevant

Table 1. Functional groups relevant to the importance of seed dispersal for seed survival and thereby plant population and community dynamics. We propose that researchers and conservation planners determine whether target plant species belong to functional groups for which dispersal disruption is likely to significantly decrease fitness vs. have only minor effects on fitness. These groups are categorized based on how much it matters whether a seed is dispersed (shown in red; groups 1–3); how much it matters where or when dispersal occurs (shown in blue; groups 4–14); and how much it matters what vector disperses the seed (shown in green; groups 15–17). Applying vulnerability assessments and SDE calculations at the level of these functional groups may enable us to achieve a predictive understanding of seed dispersal ecology in the face of combined global change and complexity. *For species exhibiting a measurable fitness boost from dispersal, seed size may dictate which abiotic or biotic vectors are effective.

Functional group category	Characteristics of functional groups likely significantly affected by seed dispersal disruption	Characteristics of functional groups likely less affected by seed dispersal disruption	Sample references
Groups for which plant fitness is affected by whether seeds are dispersed.	High colonization ability Long-distance dispersal adaptations Density-dependent survival	High competitive ability Local dispersal adaptations Density-independent survival	Coomes and Grubb (2003) Muller-Landau et al. (2003) Rey and Alcántara (2014)
Groups for which plant fitness is affected by where or when dispersal occurs.	Thin/vulnerable seed coats Shade-intolerant Fire-intolerant Self-incompatible Reproduction by seed only Intolerant of low nutrients Low phenotypic plasticity Metapopulation-dependent Negative distance-dependent mortality Inability to seed bank Seasonal dispersal	Thick/hard/spiky seed coats Shade-tolerant Fire-tolerant Self-compatible Reproduces asexually Tolerant of low nutrients High phenotypic plasticity Continuous population distribution No negative distance-dependent mortality Seed banking Low dispersal seasonality	Notman and Gorchov (2001) Alvarez-Clare and Kitajima (2007) Wenny (2001) Bond (1994) Bond (1994) Wenny (2001) Goh et al. (2013) Bohrer et al. (2005) Beckman et al. (2012) Gutterman (2000) Ruggera et al. (2015)
Groups for which plant fitness is affected by the vector of dispersal	Seed size* Intraspecific competitor/ non-facilitator Seed coat with germination inhibitors	Seed size* Intraspecific facilitator No seed coat germination inhibitors	Tamme et al. (2014) Martorell and Freckleton (2014) Traveset and Verdú (2002)

traits, functional group frameworks unite species sharing those traits under a common lens allowing generalization across diverse organisms. Because functional groups by definition describe the ecological functions present in a given site, functional group diversity has in some cases been found to predict whole-ecosystem function almost as well (or better than) species diversity ([Díaz and Cabido 2001](#)). At the same time, functional groups are conceptual constructs and thereby subject to the perspective of ecologists identifying traits they deem important to particular questions. Beginning with SDE allows us to anchor our functional group recommendations in a robust and established guiding

comprehensive framework. Thus, the three broad categories of functional groups described below have been selected to distil the comprehensive SDE framework into straightforward conceptual bins. We acknowledge that other approaches to simplifying matters are possible, perhaps based on different criteria, or on different quantifications of the same criteria. However, as we demonstrate here, considerable insights can be obtained with our approach. It is also important to acknowledge that, compared with a species-level SDE analysis, a functional group-level SDE analysis carries a risk of overgeneralization, whereby meaningful sources of variation are dismissed due to limited understanding.

Previous uses of functional groups in seed dispersal ecology have been narrow in scope (focusing on single systems or a small number of focal functions) but are indicative of the usefulness of this approach. For example, [Dennis and Westcott \(2006\)](#) distilled 26 detailed measurements of seed disperser traits into 10 trait dimensions. They used these dimensions to identify 15 functional groups in a suite of 65 Australian seed disperser species; their mathematical approach could be more broadly applied to reduce complexity in other systems ([Dennis and Westcott 2006](#)). [Rodríguez-Rodríguez et al. \(2017\)](#) categorized plant–animal interactions into typologies and evaluated links between these typologies and plant fitness. [Zamora \(2000\)](#) explored how the consistency of fitness benefits offered by seed dispersal across systems and groups of species permits generalization within functional groups. A key benefit of a functional group approach is that it could provide an understanding of the functions that may be lost when extinctions occur ([Blondel 2003](#); [Bastazini et al. 2017](#)). This conservation-oriented conceptual application was highlighted by [Schleuning et al. \(2014\)](#) in their call for more work examining the linkage between trait-based approaches such as functional group delineation and structural approaches such as network analysis ([Ruggera et al. 2015](#)). Functional groups can be used to predict the role of suites of species in an ecosystem and the response of those species to drivers of global change.

Meaningful Functional Groups in Seed Dispersal Ecology

We define plant functional groups based on traits influencing the importance of seed dispersal for plants ([Table 1](#)). Our proposed functional groups categorize plants based on: (i) how important it is to plant recruitment if seeds are dispersed at all, (ii) how important the location and timing of seed deposition are and (iii) how much vector identity matters. These groups thus define important points in the dispersal process at which seed fate may be influenced, with a focus on the fitness benefits derived from dispersal events. Applying SDE to these functional groups will enable researchers to predict how populations of the species within a group will be affected by total or partial dispersal disruption, changes in phenology or habitat conditions, or entry of non-native species into dispersal networks. A given plant species may display traits that make them likely vulnerable to dispersal disruption based on one functional group category and less vulnerable based on another; in such a case, these categories will help to pinpoint sources of such vulnerability.

Functional groups distinguishing how much it matters if a seed is dispersed at all

Species may be categorized based on the importance of dispersal for their survival and reproduction. Well-established frameworks examining fitness benefits that may be derived from seed dispersal can guide functional group determination in this arena, since functional groups can be defined by traits linked to such fitness benefits. The *escape hypothesis* states that seeds will experience fitness boosts as a result of removal from the neighbourhood of natural enemies, including pathogens, parasites, herbivores and competitors ([Howe and Smallwood 1982](#); [Howe and Miriti 2004](#)). Escape from beneath the canopy of a parent tree reduces the chances of pathogens and herbivores finding a seed ([Janzen 1970](#); [Connell 1971](#)), as well as the chance that a seed will be deposited immediately adjacent to a close relative and thus compete for necessary resources. A logical extension of this hypothesis suggests that species can be assembled into functional groups by traits indicating dependence upon such escape (i.e. susceptible to infection or herbivory; exhibiting negative density-dependence in survival and growth) vs. those less dependent upon escape (i.e. exhibiting thick seed coats or other protections against infection and herbivory; exhibiting low negative density-dependence). As an example, in a study of olive (*Olea europaea*) regeneration in human-altered vs. unaltered landscapes in Spain, proximity to maternal trees was associated with elevated seedling mortality; *O. europaea* thus appears to occupy a functional group characterized by escape dependence and negative density-dependence ([Rey and Alcántara 2014](#)) ([Table 1](#)). Similarly, fungal pathogens led to strong density-dependent mortality in *Pleradenophora longicuspis* in Belize, evidence that functional group categorization based on density-dependence is appropriate for this species ([Bagchi et al. 2010](#)). By contrast, species with low density-dependence, and thus likely to be classified into functional groups with reduced dispersal-dependence, include a suite of common species in a Panamanian rainforest, where density-dependence varies considerably among tree species ([Comita et al. 2010](#)). Species with greater seed mass exhibited reduced negative density-dependence on Barro Colorado Island ([Lebrija-Trejos et al. 2016](#)).

Previous species-specific studies have examined density-dependent damage and mortality in seeds and seedlings encountering abundant herbivores, pathogens and predators in close proximity to parent trees (the Janzen–Connell effect) (e.g. [Petermann et al. 2008](#); [Bagchi et al. 2010](#); [Liu et al. 2012](#)). Study results have been mixed, but largely show increased success

of seeds and seedlings after removal from the parent, with various explanatory mechanisms (e.g. Thomas 1990; Blundell and Peart 1998; Packer and Clay 2000; Petermann et al. 2008; Bagchi et al. 2014). These studies suggest that assigning seeds to functional groups based on whether they exhibit negative density-dependence is important. For those groups that do exhibit such density-dependence (Table 1), loss of dispersers may be expected to affect plant fitness significantly. Assigning plants to functional groups may in some cases be possible through observational studies, generalizing from what we know about similar species, and in other cases may require experimental assessments—which are still far less extensive than a traditional species-specific SDE assessment (Table 1).

Interspecific interactions can affect parameters of matrix population models, enabling their effect on fitness to be examined using elasticity and sensitivity analyses (McGraw and Caswell 1996; Horvitz et al. 1997; Benton and Grant 1999; Mills et al. 1999; Carslake et al. 2009; Jongejans et al. 2011). The effect of seed dispersal failure can be explored via elasticity analyses simulating loss of dispersers and resulting failure to escape from natural enemies or encounter recruitment sites (Howe and Miriti 2004; Brodie et al. 2009a; Rodríguez-Pérez and Traveset 2012; Traveset et al. 2012; Caughlin et al. 2015; Pérez-Méndez et al. 2015). Applying such analyses to functional groups that enable generalization beyond a few carefully measured surrogates to other species within a group might greatly expand the predictive capacity of such analyses across systems.

Other fitness benefits of seed dispersal may arise from *colonization* of unpredictable and newly available germination sites and *directed dispersal* to hospitable microsites located within a non-hospitable matrix (Wenny 2001; Howe and Miriti 2004). Dispersal is likely to matter most to functional groups of species with specialized spatio-temporal germination and growth site requirements or low competitive ability and thus high dependence on vacant establishment sites. Identification of such species may be informed by competition/colonization trade-off theory, which predicts that species exhibit a trade-off between dispersal ability and competitive ability (for example, plant species may trade off the production of a few large, well-provisioned seeds for the production of many smaller seeds) (e.g. Bolker and Pacala 1999; Dalling and Hubbell 2002; but see Coomes and Grubb 2003). Similarly, theoretical ecologists have investigated when long-distance dispersal vs. local dispersal is evolutionarily advantageous, given the fitness advantages of colonizing new sites and the lower probability of finding habitats sharing specific characteristics

at greater distances from one another (Snyder and Chesson 2003; Snyder 2011).

Functional groups distinguishing how much it matters where or when a seed is dispersed

A large fraction of the ‘quality’ element of the SDE framework centres on where and when a seed is dispersed. Fundamentally, this will dictate which abiotic and biotic resources and threats are encountered by the seed and subsequent plant (Schupp et al. 2010; Beckman and Rogers 2013). Abiotic resources may include nutrients, moisture, space and light. Abiotic threats could include drought, nutrient deficiencies, frost and fire. On the biotic side, resources could include mutualists such as soil mycorrhizae, pollinators, seed dispersers and facilitators, and threats could include herbivores, competitors, predators and pathogens. Functional groups that categorize species by whether deposition setting matters to a seed may include (i) groups of plants that are particularly susceptible to abiotic stressors/disturbances or natural enemies (e.g. plants with low competitive ability or thin seed coats) vs. (ii) those tolerant of threats (e.g. shade-tolerant, fire-tolerant, drought-tolerant, etc.). Other relevant functional groups would include species dependent on mutualists or facilitators (Calvo and Horvitz 1990; Onguene and Kuyper 2002; Hoehn et al. 2008; Teste et al. 2009), frost-intolerant species that require nurse plants, species dependent on forest gaps to escape shading and species that require a narrow range of soil nutrient content.

Plant species with plastic phenotypes may be relatively generalized with regard to their interspecific interaction requirements, suggesting that functional groups defined by plasticity may be appropriate. Plasticity may influence dependence upon certain abiotic conditions or interspecific interactions. For example, mycorrhizal associations could provide critical assistance to plant individuals with delicate or small root systems, but individuals with plastic growth (e.g. those able to divert resources towards robust root growth as required) might be less affected by an absence of root symbionts (Valladares et al. 2007; Goh et al. 2013).

Dispersal also matters for plants living in habitats that are temporally or spatially variable. Important functional groups include those with specific habitat requirements that are spatially heterogeneous (e.g. species dependent on metapopulation processes for persistence; Bohrer et al. 2005) vs. general habitat requirements that are widespread and homogeneous. For example, in a human-disturbed, patchy landscape, affinity of dispersers for seedling habitat leads to increased germination of the relic Chinese yew (*Taxus*

chinensis), indicating that directed dispersal matters for this endangered plant species (Li et al. 2016) (Table 1). In another example, seeds of the shrub *Daphne rod-riguezii* dispersed to sites below nurse plants exhibit higher seedling survival (Rodríguez-Pérez and Traveset 2010) (Table 1). Lower location specificity can be found in, for example, shade-tolerant species that exhibit physical defence mechanisms and are thus able to survive and reproduce in conditions of high competition; eight such species were studied in Panama and their physical traits documented (Alvarez-Clare and Kitajima 2007). This indicates that material characteristics can be identified to classify such species into functional groups with less dependence on dispersal (Table 1). Some functional groups may be affected by positive or negative distance- or density-dependent mortality (e.g. if seeds must be dispersed in clumps to germinate and grow; Beckman et al. 2012), or may require rare micro-conditions (Pufal and Garnock-Jones 2010). Temporally, some species can protect themselves against poor dispersal years by living many years as adults or remaining viable in a seed bank for a long time (e.g. Gutterman 2000). In other cases, the timing of dispersal interacts with characteristics that determine habitat quality (e.g. ephemeral environmental conditions or seasonally migratory dispersers) (Ruggera et al. 2015). Timing can matter on the plant side, too: in a study of *Pistacia lentiscus* dispersal in Spain, seed viability was found to vary during the fruiting season, such that dispersers interacting with the species when viability is high were more effective than those handling fruits at other times (González-Varo et al. 2018).

In theory, the functional groups most dependent upon dispersal include species in patchy habitats, those with strong density-/distance-dependent mortality, those lacking the ability to maintain a seed bank and those with specific requirements for the timing and location of the dispersal event (Table 1).

Functional groups enabling us to distinguish how much the identity of the dispersal vector matters

Dispersal syndromes are used to categorize plants by the type of vector known or assumed to best disperse their seeds. Syndromes are the most common functional group classifications used in seed dispersal ecology. Illustrating the potential value of generalization across similar dispersers, Tamme et al. (2014) successfully used plant traits to predict dispersal distances for over 500 species. Dispersal distances could then be related to dispersal syndromes, growth form and other plant traits, such as plant height and seed size (Thomson et al. 2011; Tamme et al. 2014). Previous studies have reported an interaction between seed size and dispersal vector size,

as seed size sets a lower limit on the type and size of dispersal vector that can lift (e.g. wind) or ingest (e.g. animals) the seed (Wheelwright 1985; Ganeshiah and Shaanker 1991; McConkey and Drake 2002). In some cases, dispersal syndromes explain some variation in dispersal distances and can be used to predict dispersal distances (Tamme et al. 2014), but the variation within dispersal syndromes can be very high (Clark et al. 2005; Muller-Landau et al. 2008). Dispersal syndromes tend to be broad categories (e.g. large mammal vs. small mammal vs. wind). Even within these categories, species may be dispersed by a diversity of vectors, and in some cases secondary dispersal is performed by an altogether different class of vector than primary dispersal (Böhning-Gaese et al. 1999; Vander Wall and Beck 2012). Whether the identity of the vector matters to the eventual success of the seed is an important component of understanding the role of dispersal in eventual plant population and community dynamics.

In spite of these successful attempts to achieve general insights, there are certain risks associated with generalizing across vectors (or dispersers). Identifying a dispersal syndrome may suggest that a broad category of vector is the likely disperser, but such categories could include many potential disperser species varying in effectiveness (Jordano et al. 2007; Howe 2016). Thus, dispersal syndromes are not sufficient to predict the effects of losing certain vectors. Nor do dispersal syndromes give us information on the likelihood of being dispersed by a ‘non-standard’ dispersal vector—that is, a vector other than the most common vector or vectors interacting with a particular plant—which might be more influential than ‘standard’ vectors in long-dispersal events (Higgins et al. 2003; Jordano et al. 2007) and therefore exert larger effects on plant populations (Kot et al. 1996; Neubert and Caswell 2000). As an important lesson for dispersal ecology, the concept of syndromes has faced substantial criticism in pollination ecology (Ollerton et al. 2009). Careful empirical study has demonstrated that in most cases both plants and pollinators are much more opportunistic and interact with a much broader suite of partners than morphological pollination syndromes would suggest (Waser et al. 1996; Fenster et al. 2004; Ollerton et al. 2009; Waser et al. 2018). If syndromes are similarly uninformative in dispersal, this carries implications for conservation and management, since incorrect generalization stemming from syndromes could lead to fallacious assumptions about the redundancy of dispersers within interaction networks and, consequently, about restoration and conservation needs (Howe 2016).

Although the use of dispersal syndromes *per se* thus carries a risk of drawing conclusions at too crude a

scale, straightforward functional groups founded on seed morphology and physiology (e.g. determined by seed size and shape) may dictate potential disperser suites and indicate how important different dispersal vectors may be, relative to one another, for a given plant species (Table 1). Identifying plant species at risk from dispersal disruption (e.g. McConkey et al. 2018) may be possible when the importance of vector identity is understood. Vectors may differ in the number of seeds dispersed, the condition of dispersed seeds, dispersal distances and dispersal spatial arrangements. Wind, for example, is most likely to move seeds that are small in mass (Shea 2007; Nathan et al. 2011). Disperser animals with large gape sizes are more likely than small dispersers to disperse greater numbers of larger seeds over longer distances (Cox et al. 1991). Since large dispersers with low reproductive rates are often most threatened by direct human exploitation coupled with low reproductive rates (Farwig and Berens 2012), the plant functional group that includes large-seeded species is of particular interest in seed dispersal conservation. Losses of key large dispersers can threaten plant species and functional group diversity in seed dispersal networks (Donoso et al. 2017). Dispersers with specialized habitat requirements may aggregate seeds by returning frequently to a limited number of sites (Howe 1989). Because different vectors may provide dispersal services in different ways, plant species may experience complementary dispersal services from them, with a greater diversity of vectors maximizing the success of a plant (Levin et al. 2003; Jordano et al. 2007; Bueno et al. 2013; Escribano-Avila et al. 2014; González-Varo et al. 2017). Plant functional groups of interest when determining whether a specific vector is important include groups defined by seed size, seed coat thickness (e.g. groups of species with thick coats requiring substantial gut treatment for germination), presence of germination inhibitors, and intraspecific facilitation or positive density-dependence.

Exemplifying the importance of this functional group delineation, different behaviours of large mammal dispersers resulted in differential contributions to dispersal of the large-seeded *Platymitra macrocarpa* in Thailand, with some species dispersing higher quantities of seeds with poor survival outcomes and others dispersing fewer seeds with greater success per seed (McConkey et al. 2018). In this case, dispersers contributed differentially to the dispersal of the plant but overall plant regeneration was poor, leading researchers to speculate that there may be important dispersers that are missing or rare (McConkey et al. 2018). In that context, the large seed size of the plant suggests that the identity of the dispersal vector in this example is important according

to the functional groups we propose (Table 1) (McConkey et al. 2018). By contrast, SDE of a suite of bird species was studied for two *Miconia* species in Brazil (Santos et al. 2017). Although the birds varied in the quantity of seeds they dispersed, they did not vary in quality of dispersal (Santos et al. 2017). *Miconia* species with their small seeds and large disperser suites (e.g. Levey and Byrne 1993) therefore appear to fall into a proposed functional group for which vector identity is less important (Table 1).

Dispersal vector identity has been shown to affect population growth rates for some but not all of the few vertebrate-dispersed plant species that have been studied (e.g. Godinez-Alvarez and Jordano 2007; Brodie et al. 2009b; Loayza and Knight 2010). However, the importance of different vectors is unknown for most plant species, and that lack of clarity hampers our ability to predict the outcomes of changes in vectors. Predictions are better-developed for ballistic- and wind-mediated dispersal than for animal-mediated dispersal (Skarpaas and Shea 2007; Nathan et al. 2011; Bullock et al. 2012), in large part because of the complexity of animal behaviour and movement and the diffuse nature of most seed dispersal systems, wherein multiple animals disperse any given plant (Shea 2007). Even when detailed information about the role of specific vectors has been obtained for a given plant species, studies are often narrow in spatial and temporal extent and thus context-dependent (i.e. information is specific to a particular time and place, given a particular disturbance history), and the importance of individual vectors may change under different contexts.

Using Functional Groups to Close Our Knowledge Gaps

The use of functional groups defined by dispersal-related traits can reduce the amount of data needed to parameterize models (Mokany et al. 2014). The digital availability of trait data is increasing (e.g. via publicly accessible databases such as TRY; Kattge et al. 2011) but continued empirical research is needed to relate those data to dispersal processes. Even so, certain functional groups can now be defined and used to distinguish species that are relatively more or less strongly dispersal-dependent (Table 1). If a particular plant species belongs to a group for which fitness is strongly linked to dispersal (Table 1, column 2), we can predict that this species is likely to be vulnerable in the face of dispersal disruption, based strictly on functional group.

While functional groups may enable us to generalize across full plant communities, overgeneralization could

cause us to lose sight of meaningful sources of variation. It is thus necessary to test the value of functional group approaches in seed dispersal ecology by generating predictions based on them and then testing those predictions against species-specific data. As functional groups ‘pass’ these tests, our ability to generalize by constructing our models and predictions around functional groups will help us manage the broad variability and context-dependence that can characterize seed dispersal events (Robledo-Arnuncio *et al.* 2014).

Compiling large amounts of data from multiple systems to test functional group approaches can be resource-intensive, although increasingly global databases are becoming available and serving as common data hubs (e.g. COMPADRE Plant Matrix Database and COMADRE Animal Matrix Database; Max Planck Institute for Demographic Research (Germany); available at www.compadre-db.org). One promising approach is to standardize data collection by many research groups working across many systems (e.g. NutNet; Borer *et al.* 2014). If groups collect the same data across systems, those data can be introduced to a common modelling platform to explore patterns that hold across systems. By assembling data sets in this way, the costs are spread across research groups, and data collection can be useful even when sample sizes within a specific system are limited. As an example, standardized data collection across systems has enabled researchers to identify consistent patterns of grassland responses to land use change (Garnier *et al.* 2007). We thus recommend that research teams join forces to collect standardized data exploring varying effectiveness of dispersal by different agents, the role of spatio-temporal dynamics and the influence of interspecific interactions pre-, mid- and post-dispersal. Once data are collected in many systems, it will be necessary to bring them together to make them available for broad analysis. Depositing seed dispersal data into public-access repositories is therefore important. Useful repositories include Dryad (<http://datadryad.org>) and KNB (<http://knb.ecoinformatics.org>).

To grow our understanding of seed dispersal ecology and to predict the likely effects of environmental changes, we advocate a new focus on seed dispersal functional groups. The plant functional groups described here are defined based on plant dependence on seed dispersal for plant population persistence and the likelihood of experiencing meaningful dispersal disruption. Analysis at a functional group level generalizes across species and systems and may enable us to more effectively assess the role of seed dispersal in the shaping of

plant populations and communities. We invite the ecological community to join us in this effort.

Conflict of interest

None declared.

Sources of funding

Ideas for this manuscript initiated during the Seed Dispersal Workshop held in May 2016 at the Socio-Environmental Synthesis Center in Annapolis, MD and supported by the US National Science Foundation Grant DEB-1548194 to N.G.B. and the National Socio-Environmental Synthesis Center under the US National Science Foundation Grant DBI-1052875. D.Z. received funding from the Swiss National Science Foundation (SNF, grant: PZ00P3_168136/1) and from the German Science Foundation (DFG, grant: ZU 361/1- 1).

Contributions by the authors

C.A. led the development of the concepts, writing, and revising of the manuscript with input from N.G.B. and H.S.R. All authors contributed to the development of concepts and are listed in order of contribution and alphabetical order within each level of contribution.

Acknowledgements

Our discussion emerged from a National Science Foundation-funded Seed Dispersal Workshop held at the National Socio-Environmental Synthesis Center in May 2016. All workshop participants are co-authors on this paper and have contributed to the development of these ideas.

Literature Cited

- Albrecht J, Neuschulz EL, Farwig N. 2012. Impact of habitat structure and fruit abundance on avian seed dispersal and fruit predation. *Basic and Applied Ecology* **13**:347–354.
- Alvarez-Buylla ER, Martínez-Ramos M. 1990. Seed bank versus seed rain in the regeneration of a tropical pioneer tree. *Oecologia* **84**:314–325.
- Alvarez-Clare S, Kitajima K. 2007. Physical defence traits enhance seedling survival of neotropical tree species. *Functional Ecology* **21**:1044–1054.
- Aslan CE, Zavaleta ES, Tershy B, Croll D. 2013. Mutualism disruption threatens global plant biodiversity: a systematic review. *PLoS One* **8**:e66993.
- Auffret AG, Rico Y, Bullock JM, Hooftman DA, Pakeman RJ, Soons MB, Suárez-Esteban A, Traveset A, Wagner HH, Cousins SA. 2017. Plant functional connectivity-integrating

- landscape structure and effective dispersal. *Journal of Ecology* **105**:1648–1656.
- Aukema JE. 2004. Distribution and dispersal of desert mistletoe is scale-dependent, hierarchically nested. *Ecography* **27**:137–144.
- Bacles CF, Lowe AJ, Ennos RA. 2006. Effective seed dispersal across a fragmented landscape. *Science* **311**:628.
- Bagchi R, Gallery RE, Gripenberg S, Gurr SJ, Narayan L, Addis CE, Freckleton RP, Lewis OT. 2014. Pathogens and insect herbivores drive rainforest plant diversity and composition. *Nature* **506**:85–88.
- Bagchi R, Swinfield T, Gallery RE, Lewis OT, Gripenberg S, Narayan L, Freckleton RP. 2010. Testing the Janzen-Connell mechanism: pathogens cause overcompensating density dependence in a tropical tree. *Ecology Letters* **13**:1262–1269.
- Bastazini VAG, Debastiani VJ, Azambuja BO, Guimaraes PR, Pillar VD. 2017. Loss of generalist plant species and functional diversity decreases the robustness of a seed dispersal network. *bioRxiv*. doi:10.1101/187179.
- Beckman NG, Neuhauser C, Muller-Landau HC. 2012. The interacting effects of clumped seed dispersal and distance- and density-dependent mortality on seedling recruitment patterns. *Journal of Ecology* **100**:862–873.
- Beckman NG, Rogers HS. 2013. Consequences of seed dispersal for plant recruitment in tropical forests: interactions within the seedscape. *Biotropica* **45**:666–681.
- Benton TG, Grant A. 1999. Elasticity analysis as an important tool in evolutionary and population ecology. *Trends in Ecology & Evolution* **14**:467–471.
- Blondel J. 2003. Guilds or functional groups: does it matter? *Oikos* **100**:223–231.
- Blundell AG, Peart DR. 1998. Distance-dependence in herbivory and foliar condition for juvenile *Shorea* trees in Bornean dipterocarp rain forest. *Oecologia* **117**:151–160.
- Böhning-Gaese K, Gaese BH, Rabemanantsoa SB. 1999. Importance of primary and secondary seed dispersal in the Malagasy tree *Commiphora guillaumini*. *Ecology* **80**:821–832.
- Borner G, Nathan R, Volis S. 2005. Effects of long-distance dispersal for metapopulation survival and genetic structure at ecological time and spatial scales. *Journal of Ecology* **93**:1029–1040.
- Bolker BM, Pacala SW. 1999. Spatial moment equations for plant competition: understanding spatial strategies and the advantages of short dispersal. *The American Naturalist* **153**:575–602.
- Bond WJ. 1994. Do mutualisms matter? Assessing the impact of pollinator and disperser disruption on plant extinction. *Philosophical Transactions of the Royal Society of London B: Biological Sciences* **344**:83–90.
- Borer ET, Harpole WS, Adler PB, Lind EM, Orrock JL, Seabloom EW, Smith MD. 2014. Finding generality in ecology: a model for globally distributed experiments. *Methods in Ecology and Evolution* **5**:65–73.
- Brodie JF, Helmy OE, Brockelman WY, Maron JL. 2009a. Bushmeat poaching reduces the seed dispersal and population growth rate of a mammal-dispersed tree. *Ecological Applications* **19**:854–863.
- Brodie JF, Helmy OE, Brockelman WY, Maron JL. 2009b. Functional differences within a guild of tropical mammalian frugivores. *Ecology* **90**:688–698.
- Bueno RS, Guevara R, Ribeiro MC, Culot L, Bufalo FS, Galetti M. 2013. Functional redundancy and complementarities of seed dispersal by the last neotropical megafrugivores. *PLoS One* **8**:e56252.
- Bullock JM, White SM, Prudhomme C, Tansey C, Perea R, Hooftman DA. 2012. Modelling spread of British wind-dispersed plants under future wind speeds in a changing climate. *Journal of Ecology* **100**:104–115.
- Calvo RN, Horvitz CC. 1990. Pollinator limitation, cost of reproduction, and fitness in plants: a transition-matrix demographic approach. *The American Naturalist* **136**:499–516.
- Carslake D, Townley S, Hodgson DJ. 2009. Patterns and rules for sensitivity and elasticity in population projection matrices. *Ecology* **90**:3258–3267.
- Caughlin TT, Ferguson JM, Lichstein JW, Zuidema PA, Bunyavechewin S, Levey DJ. 2015. Loss of animal seed dispersal increases extinction risk in a tropical tree species due to pervasive negative density dependence across life stages. *Proceedings of the Royal Society of London B: Biological Sciences* **272**:20142095.
- Chazdon RL, Finegan B, Capers RS, Salgado-Negret B, Casanoves F, Boukili V, Norden N. 2010. Composition and dynamics of functional groups of trees during tropical forest succession in north-eastern Costa Rica. *Biotropica* **42**:31–40.
- Chen Q, Tomlinson KW, Cao L, Wang B. 2017. Effects of fragmentation on the seed predation and dispersal by rodents differ among species with different seed size. *Integrative Zoology* **12**:468–476.
- Clark CJ, Poulsen JR, Bolker BM, Connor EF, Parker VT. 2005. Comparative seed shadows of bird-, monkey-, and wind-dispersed trees. *Ecology* **86**:2684–2694.
- Comita LS, Muller-Landau HC, Aguilar S, Hubbell SP. 2010. Asymmetric density dependence shapes species abundances in a tropical tree community. *Science* **329**:330–332.
- Connell JH. 1971. On the role of natural enemies in preventing competitive exclusion in some marine animals and in rain forest trees. Dynamics of numbers in populations. In: Proceedings of the Advanced Study Institute on Dynamics of Numbers in Populations, Oosterbeek, The Netherlands, 7–18 September 1970, 298–312.
- Coomes DA, Grubb PJ. 2003. Colonization, tolerance, competition and seed-size variation within functional groups. *Trends in Ecology & Evolution* **18**:283–291.
- Corlett RT, Westcott DA. 2013. Will plant movements keep up with climate change? *Trends in Ecology & Evolution* **28**:482–488.
- Côrtes MC, Uriarte M. 2013. Integrating frugivory and animal movement: a review of the evidence and implications for scaling seed dispersal. *Biological Reviews* **88**:255–272.
- Cox PA, Elmqvist T, Pierson ED, Rainey WE. 1991. Flying foxes as strong interactors in South Pacific island ecosystems: a conservation hypothesis. *Conservation Biology* **5**:448–454.
- Dalling JW, Hubbell SP. 2002. Seed size, growth rate and gap microsite conditions as determinants of recruitment success for pioneer species. *Journal of Ecology* **90**:557–568.
- Dennis AJ, Westcott DA. 2006. Reducing complexity when studying seed dispersal at community scales: a functional classification of vertebrate seed dispersers in tropical forests. *Oecologia* **149**:620–634.
- Diaz S, Cabido M. 2001. Vive la différence: plant functional diversity matters to ecosystem processes. *Trends in Ecology & Evolution* **16**:646–655.
- Donoso I, Schleuning M, Garcia D, Fründ J. 2017. Defaunation effects on plant recruitment depend on size matching and size trade-offs in seed-dispersal networks. *Proceedings of the Royal Society of London B: Biological Sciences* **284**:20162664.

- Effiom EO, Nunez-Iturri G, Smith HG, Ottosson U, Olsson O. 2013. Bushmeat hunting changes regeneration of African rainforests. *Proceedings of the Royal Society of London B: Biological Sciences* **280**:20130246.
- Escribano-Avila G, Calviño-Cancela M, Pías B, Virgós E, Valladares F, Escudero A. 2014. Diverse guilds provide complementary dispersal services in a woodland expansion process after land abandonment. *Journal of Applied Ecology* **51**:1701–1711.
- Farwig N, Berens DG. 2012. Imagine a world without seed dispersers: a review of threats, consequences and future directions. *Basic and Applied Ecology* **13**:109–115.
- Fenster CB, Armbruster WS, Wilson P, Dudash MR, Thomson JD. 2004. Pollination syndromes and floral specialization. *Annual Review of Ecology, Evolution, and Systematics* **35**:375–403.
- Fricke EC, Simon MJ, Reagan KM, Levey DJ, Riffell JA, Carlo TA, Tewksbury JJ. 2013. When condition trumps location: seed consumption by fruit-eating birds removes pathogens and predator attractants. *Ecology Letters* **16**:1031–1036.
- Ganeshiah KN, Shaanker RU. 1991. Seed size optimization in a wind dispersed tree *Butea monosperma*: a trade-off between seedling establishment and pod dispersal efficiency. *Oikos* **60**:3–6.
- García D, Martínez D. 2012. Species richness matters for the quality of ecosystem services: a test using seed dispersal by frugivorous birds. *Proceedings of the Royal Society of London B: Biological Sciences* **279**:3106–3113.
- Garnier E, Lavorel S, Ansquer P, Castro H, Cruz P, Dolezal J, Eriksson O, Fortunel C, Freitas H, Golodets C, Grigulis K, Jouany C, Kazakou E, Kigel J, Kleyer M, Lehsten V, Leps J, Meier T, Pakeman R, Papadimitriou M, Papanastasis VP, Queded H, Quéfier F, Robson M, Roumet C, Rusch G, Skarpe C, Sternberg M, Theau JP, Thébault A, Vile D, Zarovali MP. 2007. Assessing the effects of land-use change on plant traits, communities and ecosystem functioning in grasslands: a standardized methodology and lessons from an application to 11 European sites. *Annals of Botany* **99**:967–985.
- Godinez-Alvarez H, Jordano P. 2007. An empirical approach to analysing the demographic consequences of seed dispersal by frugivores. In: Dennis AJ, Schupp EW, Green RJ, Westcott DA, eds. *Seed dispersal: theory and its application in a changing world*. Wallingford, UK: CAB International, 391–406.
- Goh CH, Veliz Vallejos DF, Nicotra AB, Mathesius U. 2013. The impact of beneficial plant-associated microbes on plant phenotypic plasticity. *Journal of Chemical Ecology* **39**:826–839.
- González-Castro A, Calviño-Cancela M, Nogales M. 2015. Comparing seed dispersal effectiveness by frugivores at the community level. *Ecology* **96**:808–818.
- González-Varo JP, Arroyo JM, Jordano P. 2018. The timing of frugivore-mediated seed dispersal effectiveness. *Molecular Ecology*. doi:10.1111/mec.14850.
- González-Varo JP, Carvalho CS, Arroyo JM, Jordano P. 2017. Unravelling seed dispersal through fragmented landscapes: frugivore species operate unevenly as mobile links. *Molecular Ecology* **26**:4309–4321.
- Gutterman Y. 2000. Environmental factors and survival strategies of annual plant species in the Negev Desert, Israel. *Plant Species Biology* **15**:113–125.
- Higgins SI, Nathan R, Cain ML. 2003. Are long-distance dispersal events in plants usually caused by nonstandard means of dispersal? *Ecology* **84**:1945–1956.
- Hoehn P, Tschardt T, Tylianakis JM, Steffan-Dewenter I. 2008. Functional group diversity of bee pollinators increases crop yield. *Proceedings of the Royal Society of London B: Biological Sciences* **275**:2283–2291.
- Horvitz C, Schemske DW, Caswell H. 1997. The relative “importance” of life-history stages to population growth: prospective and retrospective analyses. In: Tuljapurkar S, Caswell H, eds. *Structured-population models in marine, terrestrial, and freshwater systems*. New York: Springer, 247–271.
- Howe HF. 1989. Scatter-and clump-dispersal and seedling demography: hypothesis and implications. *Oecologia* **79**:417–426.
- Howe HF. 2016. Making dispersal syndromes and networks useful in tropical conservation and restoration. *Global Ecology and Conservation* **6**:152–178.
- Howe HF, Miriti MN. 2004. When seed dispersal matters. *BioScience* **54**:651–660.
- Howe HF, Smallwood J. 1982. Ecology of seed dispersal. *Annual Review of Ecology and Systematics* **13**:201–228.
- Janzen DH. 1970. Herbivores and the number of tree species in tropical forests. *The American Naturalist* 501–528.
- Jongejans E, Shea K, Skarpaas O, Kelly D, Ellner SP. 2011. Importance of individual and environmental variation for invasive species spread: a spatial integral projection model. *Ecology* **92**:86–97.
- Jordano P, García C, Godoy JA, García-Castaño JL. 2007. Differential contribution of frugivores to complex seed dispersal patterns. *Proceedings of the National Academy of Sciences of the United States of America* **104**:3278–3282.
- Karubian J, Durães R. 2009. Effects of seed disperser social behavior on patterns of seed movement and deposition. *Oecologia Brasiliensis* **13**:45–57.
- Kattge J, Díaz S, Lavorel S, Prentice IC, Leadley P, Bönsch G, Garnier E, Westoby M, Reich PB, Wright IJ, Cornelissen JHC. 2011. TRY – a global database of plant traits. *Global Change Biology* **17**:2905–2935.
- Katul GG, Porporato A, Nathan R, Siqueira M, Soons MB, Poggi D, Horn HS, Levin SA. 2005. Mechanistic analytical models for long-distance seed dispersal by wind. *The American Naturalist* **166**:368–381.
- Kot M, Lewis MA, van den Driessche P. 1996. Dispersal data and the spread of invading organisms. *Ecology* **77**:2027–2042.
- Ladley JJ, Kelly D. 1996. Dispersal, germination and survival of New Zealand mistletoes (Loranthaceae): dependence on birds. *New Zealand Journal of Ecology* **20**:69–79.
- Lebrija-Trejos E, Reich PB, Hernández A, Wright SJ. 2016. Species with greater seed mass are more tolerant of conspecific neighbours: a key driver of early survival and future abundances in a tropical forest. *Ecology Letters* **19**:1071–1080.
- Levey DJ, Byrne MM. 1993. Complex ant-plant interactions: rain-forest ants as secondary dispersers and post-dispersal seed predators. *Ecology* **74**:1802–1812.
- Levin SA, Muller-Landau HC, Nathan R, Chave J. 2003. The ecology and evolution of seed dispersal: a theoretical perspective. *Annual Review of Ecology, Evolution, and Systematics* **34**:575–604.
- Li N, Li XH, An SQ, Lu CH. 2016. Impact of multiple bird partners on the seed dispersal effectiveness of China’s relic trees. *Scientific Reports* **6**:17489.
- Liu X, Liang M, Etienne RS, Wang Y, Staehelin C, Yu S. 2012. Experimental evidence for a phylogenetic Janzen-Connell effect in a subtropical forest. *Ecology Letters* **15**:111–118.

- Loayza AP, Knight T. 2010. Seed dispersal by pulp consumers, not “legitimate” seed dispersers, increases *Guettarda viburnoides* population growth. *Ecology* **91**:2684–2695.
- Martorell C, Freckleton RP. 2014. Testing the roles of competition, facilitation and stochasticity on community structure in a species-rich assemblage. *Journal of Ecology* **102**:74–85.
- McConkey KR, Brockelman WY, Saralamba C. 2014. Mammalian frugivores with different foraging behavior can show similar seed dispersal effectiveness. *Biotropica* **46**:647–651.
- McConkey KR, Drake DR. 2002. Extinct pigeons and declining bat populations: are large seeds still being dispersed in the tropical Pacific? In: Levey DJ, Silva WR, Galetti M, eds. *Seed dispersal and frugivory: ecology, evolution, and conservation*. Wallingford, UK: CABI, 381.
- McConkey KR, Drake DR. 2006. Flying foxes cease to function as seed dispersers long before they become rare. *Ecology* **87**:271–276.
- McConkey KR, Nathalang A, Brockelman WY, Saralamba C, Santon J, Matmoon U, Somnuk R, Srinoppawan K. 2018. Different mega-fauna vary in their seed dispersal effectiveness of the mega-faunal fruit *Platymitra macrocarpa* (Annonaceae). *PLoS One* **13**:e0198960.
- McGraw JB, Caswell H. 1996. Estimation of individual fitness from life-history data. *The American Naturalist* **147**:47–64.
- Metzger JP. 2000. Tree functional group richness and landscape structure in a Brazilian tropical fragmented landscape. *Ecological Applications* **10**:1147–1161.
- Mills LS, Doak DF, Wisdom MJ. 1999. Reliability of conservation actions based on elasticity analysis of matrix models. *Conservation Biology* **13**:815–829.
- Mokany K, Prasad S, Westcott DA. 2014. Loss of frugivore seed dispersal services under climate change. *Nature Communications* **5**:3971.
- Morales JM, García D, Martínez D, Rodríguez-Pérez J, Herrera JM. 2013. Frugivore behavioural details matter for seed dispersal: a multi-species model for Cantabrian thrushes and trees. *PLoS One* **8**:e65216.
- Muller-Landau HC, Levin SA, Keymer JE. 2003. Theoretical perspectives on evolution of long-distance dispersal and the example of specialized pests. *Ecology* **84**:1957–1967.
- Muller-Landau HC, Wright SJ, Calderón O, Condit R, Hubbell SP. 2008. Interspecific variation in primary seed dispersal in a tropical forest. *Journal of Ecology* **96**:653–667.
- Muñoz MC, Schaefer HM, Böhning-Gaese K, Schleuning M. 2017. Importance of animal and plant traits for fruit removal and seedling recruitment in a tropical forest. *Oikos* **126**:823–832.
- Nathan R, Katul GG, Bohrer G, Kuparinen A, Soons MB, Thompson SE, Trakhtenbrot A, Horn HS. 2011. Mechanistic models of seed dispersal by wind. *Theoretical Ecology* **4**:113–132.
- Neubert MG, Caswell H. 2000. Demography and dispersal: calculation and sensitivity analysis of invasion speed for structured populations. *Ecology* **81**:1613–1628.
- Notman E, Gorchov DL. 2001. Variation in post-dispersal seed predation in mature Peruvian lowland tropical forest and fallow agricultural sites 1. *Biotropica* **33**:621–636.
- Nogales M, González-Castro A, Rumeu B, Traveset A, Vargas P, Jaramillo P, Olesen JM, Heleno RH. 2017. Contribution by vertebrates to seed dispersal effectiveness in the Galápagos Islands: a community-wide approach. *Ecology* **98**:2049–2058.
- Ollerton J, Alarcón R, Waser NM, Price MV, Watts S, Cranmer L, Hingston A, Peter CI, Rotenberry J. 2009. A global test of the pollination syndrome hypothesis. *Annals of Botany* **103**:1471–1480.
- Onguene NA, Kuyper TW. 2002. Importance of the ectomycorrhizal network for seedling survival and ectomycorrhiza formation in rain forests of south Cameroon. *Mycorrhiza* **12**:13–17.
- Packer A, Clay K. 2000. Soil pathogens and spatial patterns of seedling mortality in a temperate tree. *Nature* **404**:278–281.
- Pérez-Méndez N, Jordano P, Valido A. 2015. Downsized mutualisms: consequences of seed dispersers’ body-size reduction for early plant recruitment. *Perspectives in Plant Ecology, Evolution and Systematics* **17**:151–159.
- Petermann JS, Fergus AJ, Turnbull LA, Schmid B. 2008. Janzen-Connell effects are widespread and strong enough to maintain diversity in grasslands. *Ecology* **89**:2399–2406.
- Puerta-Piñero C, Muller-Landau HC, Calderón O, Wright SJ. 2013. Seed arrival in tropical forest tree fall gaps. *Ecology* **94**:1552–1562.
- Pufal G, Garnock-Jones P. 2010. Hygrochastic capsule dehiscence supports safe site strategies in New Zealand alpine veronica (Plantaginaceae). *Annals of Botany* **106**:405–412.
- Razafindratsima OH, Dunham AE. 2016. Frugivores bias seed-adult tree associations through nonrandom seed dispersal: a phylogenetic approach. *Ecology* **97**:2094–2102.
- Rey PJ, Alcántara JM. 2014. Effects of habitat alteration on the effectiveness of plant-avian seed dispersal mutualisms: consequences for plant regeneration. *Perspectives in Plant Ecology, Evolution and Systematics* **16**:21–31.
- Robledo-Arnuncio JJ, Klein EK, Muller-Landau HC, Santamaría L. 2014. Space, time and complexity in plant dispersal ecology. *Movement Ecology* **2**:16.
- Rodríguez-Pérez J, Traveset A. 2010. Seed dispersal effectiveness in a plant-lizard interaction and its consequences for plant regeneration after disperser loss. *Plant Ecology* **207**:269–280.
- Rodríguez-Pérez J, Traveset A. 2012. Demographic consequences for a threatened plant after the loss of its only disperser. Habitat suitability buffers limited seed dispersal. *Oikos* **121**:835–847.
- Rodríguez-Rodríguez MC, Jordano P, Valido A. 2017. Functional consequences of plant-animal interactions along the mutualism-antagonism gradient. *Ecology* **98**:1266–1276.
- Rother DC, Pizo MA, Jordano P. 2016. Variation in seed dispersal effectiveness: the redundancy of consequences in diversified tropical frugivore assemblages. *Oikos* **125**:336–342.
- Ruggera RA, Blendinger PG, Gomez MD, Marshak C. 2015. Linking structure and functionality in mutualistic networks: do core frugivores disperse more seeds than peripheral species? *Oikos* **125**:541–555.
- Santos AM, Jacobi CM, Silveira FA. 2017. Frugivory and seed dispersal effectiveness in two *Miconia* (Melastomataceae) species from ferruginous campo rupestre. *Seed Science Research* **27**:65–73.
- Sato H, Itoh A, Kohyama T. 2007. SEIB-DGVM: a new dynamic global vegetation model using a spatially explicit individual-based approach. *Ecological Modelling* **200**:279–307.
- Scheiter S, Higgins SI. 2009. Impacts of climate change on the vegetation of Africa: an adaptive dynamic vegetation modelling approach. *Global Change Biology* **15**:2224–2246.
- Schleuning M, Fründ J, García D. 2014. Predicting ecosystem functions from biodiversity and mutualistic networks: an extension

- of trait-based concepts to plant–animal interactions. *Ecography* **38**:380–392.
- Schupp EW, Jordano P, Gómez JM. 2010. Seed dispersal effectiveness revisited: a conceptual review. *The New Phytologist* **188**:333–353.
- Shea K. 2007. Ecology. How the wood moves. *Science* **315**:1231–1232.
- Skarpaas O, Shea K. 2007. Dispersal patterns, dispersal mechanisms, and invasion wave speeds for invasive thistles. *The American Naturalist* **170**:421–430.
- Snyder RE. 2011. Leaving home ain't easy: non-local seed dispersal is only evolutionarily stable in highly unpredictable environments. *Proceedings of the Royal Society of London B: Biological Sciences* **278**:739–744.
- Snyder RE, Chesson P. 2003. Local dispersal can facilitate coexistence in the presence of permanent spatial heterogeneity. *Ecology Letters* **6**:301–309.
- Tamme R, Götzenberger L, Zobel M, Bullock JM, Hooftman DA, Kaasik A, Pärtel M. 2014. Predicting species' maximum dispersal distances from simple plant traits. *Ecology* **95**:505–513.
- Teste FP, Simard SW, Durall DM, Guy RD, Jones MD, Schoonmaker AL. 2009. Access to mycorrhizal networks and roots of trees: importance for seedling survival and resource transfer. *Ecology* **90**:2808–2822.
- Thomas CD. 1990. Herbivore diets, herbivore colonization, and the escape hypothesis. *Ecology* **71**:610–615.
- Thomson FJ, Moles AT, Auld TD, Kingsford RT. 2011. Seed dispersal distance is more strongly correlated with plant height than with seed mass. *Journal of Ecology* **99**:1299–1307.
- Traveset A, González-Varo JP, Valido A. 2012. Long-term demographic consequences of a seed dispersal disruption. *Proceedings of the Royal Society of London B: Biological Sciences* **279**:3298–3303.
- Traveset A, Verdú M. 2002. A meta-analysis of the effect of gut treatment on seed germination. In: Levey DJ, Silva WR, Galetti M, eds. *Seed dispersal and frugivory: ecology, evolution, and conservation*. Wallingford, UK: CABI, 339.
- Valladares F, Gianoli E, Gómez JM. 2007. Ecological limits to plant phenotypic plasticity. *The New Phytologist* **176**:749–763.
- Vander Wall SB, Beck MJ. 2012. A comparison of frugivory and scatter-hoarding seed-dispersal syndromes. *The Botanical Review* **78**:10–31.
- Waser NM, CaraDonna PJ, Price MV. 2018. Atypical flowers can be as profitable as typical hummingbird flowers. *The American Naturalist* **192**:644–653.
- Waser NM, Chittka L, Price MV, Williams NM, Ollerton J. 1996. Generalization in pollination systems, and why it matters. *Ecology* **77**:1043–1060.
- Wenny DG. 2001. Advantages of seed dispersal: a re-evaluation of directed dispersal. *Evolutionary Ecology Research* **3**:51–74.
- Wheelwright NT. 1985. Fruit-size, gape width, and the diets of fruit-eating birds. *Ecology* **66**:808–818.
- Wunderle JM. 1997. The role of animal seed dispersal in accelerating native forest regeneration on degraded tropical lands. *Forest Ecology and Management* **99**:223–235.
- Zamora R. 2000. Functional equivalence in plant–animal interactions: ecological and evolutionary consequences. *Oikos* **88**:442–447.