# Functional Ecology

Volume 32 • Number 12 • December 2018

ISSN 1365-2435 (Online)



# Editors: Charles Fox, Ken Thompson, Alan Knapp, Lara Ferry and Enrico Rezende

- Structural traits dictate abiotic stress amelioration by intertidal oysters
- Change in drivers of mangrove crown displacement along a salinity stress gradient
- Trait evolution in tropical rubber trees is related to dry season intensity
- Frugivore biodiversity and complementarity in interaction networks enhance landscape-scale seed dispersal function
- Effects of host colony size and hygiene behaviours on social spider kleptoparasite loads along an elevation andioat
- Plasticity in leaf litter traits partly mitigates the impact of thinning on forest floor carbon cycling



#### Editors

Charles Fox Department of Entomology, University of Kentucky, USA Ken Thompson Department of Animal and Plant Sciences, University of Sheffield, UK Alan Knapp Department of Biology, Colorado State University, USA Lara Ferry Arizona State University, USA Enrico Rezende Department of Ecology, P. Universidad Católica de Chile, Chile Associate Editors J. Alahuhta Oulu, Finland D. Allen Oklahoma, USA F. Angelier Villiers en Bois, France . Baltzer Sackville, Canada S. Barribeau Liverpool, UK K. Barton Hawai'i, USA J. Bell Rothamsted Research, UK A. Bennett Dundee, UK A. Biere Wageningen, The Netherlands N. Boogert Fife, UK M. J. I. Briones Vigo, Spain A. K. Brody Vermont, USA

D. R. Campbell California, USA E. Carrington Washington, USA S. Clusella-Trullas Stellenbosch, South Africa

J. Cooke Milton Keynes, UK D. Costantini Antwerp, Belgium

D. Crocker California, USA G. Davidowitz Arizona, USA

S.E. Diamond Cleveland, OH, USA

E. Dorrepaal Sweden R. El-Sabaawi Victoria, Canada

K. Field Sheffield, UK

R. Gallery Arizona, USA I. Galvan Seville, Spain

D. Garcia Asturias, Spain

D. Goddy Seville, Spain J. Goldbogen California, USA D. Grémillet Montpellier, France J. Grindstaff Oklahoma, USA M. Hart British Columbia, Canada

D. Hawley Virginia, USA

R. Heleno Coimbra, Portugal T. E. Higham Riverside, USA

L. Holeski Arizona, USA

W. A. Hopkins Virginia, USA J. Husak St. Thomas, USA

C. Isaksson Lund, Sweden

T. H. Jones Cardiff, UK S. Killen Aberdeen, UK

J. Koricheva Royal Holloway, UK G. Kudo Hokkaido, Japan

M. Larjavaara Helsinki, Finland K. P. Lee Seoul, Republic of Korea J-F. Lemaître Lyon, France

S. Leroux Newfoundland, Canada

D. Levesque Maine, USA L. Llorens Girona, Spain

H. MacMillan Ontario, Canada J. Manson Alberta, Canada

A. J. Manzaneda Jaén, Spain

D. Marshall Monash, Australia K. Marshall Ohio, USA C. McArthur Sydney, Australia

C. Meynard Virginia, USA C. W. Miller Gainesville, FL, USA

K. Mooney California, USA

E. Morriën, Amsterdam, The Netherlands H. Muller-Landeau Washington DC, USA

S. Niu Beijing, China J. Ohlberger Washington, USA R. Oliveira Campinas, Brazil

R. Ostertag Hilo, HI, USA

J. Overgaard Aarhus, Denmark C.E.T. Paine Stirling, UK

A. B. Pedersen Edinburgh, UK

M. Pfrender Notre Dame, USA S. Portugal Royal Holloway, UK

S. Power Sydney, Australia N. Priest Bath, UK

S. Rasmann Neuchatel, Switzerland

D. N. Reznick Riverside, USA S. Russo Nebraska, USA

A. Sala Missoula, USA

B. Sandercock Trondheim, Norway

E. J. Sayer Lancaster, UK

M. Schleuning Frankfurt, Germany

C. Seymour Claremont, South Africa A. Siepielski Arkansas, USA

K.W. Sockman North Carolina, USA

C. J. Stevens Lancaster, UK M. Tjoelker Sydney, Australia

J. Tobias London, UK

Y. Vindenes Oslo, Norway C. Violle Montpellier, France

F. Wang Guangzhou, PR China C. White Melbourne, Australia

C. Williams California, USA

T. D. Williams Simon Fraser University, Canada

Managing Editor Emilie Aimé (e-mail: managingeditor@functionalecology.org)

Senior Assistant Editor Jennifer Meyer (e-mail: admin@functionalecology.org)

Production Editor: Marion Laws (e-mail: fec@wiley.com)

Editorial policy: Functional Ecology publishes high-impact papers that enable a mechanistic understanding of ecological pattern and process from the organismic to the ecosystem scale. Because of the multifaceted nature of this challenge, papers can be based on a wide range of approaches. Thus, manuscripts may vary from physiological, genetics, life-history, and behavioural perspectives for organismal studies to community and biogeochemical studies when the goal is to understand ecosystem and larger scale ecological phenomena. The Journal publishes the following

Standard Papers - experimental, comparative or theoretical papers.

Reviews - syntheses of topics of broad ecological interest.

Perspectives - short articles presenting new ideas (without data) intended to stimulate scientific debate.

Special Features - a collection of manuscripts, typically Reviews or Perspectives, on a single theme.

Manuscripts should be in English and should be submitted online at http:// mc.manuscriptcentral.com/fe-besjournals.

**Guidelines for Authors** can be found on the journal website at www. functionalecology.org A pdf file of each paper will be provided free of charge and printed copies may be ordered when proofs are returned. A scale of charges will be sent with the proofs

Publisher: Functional Ecology is published by John Wiley & Sons Ltd, 9600 Garsington Road, Oxford OX4 2DQ, UK. Tel: +44 1865 776868; Fax: +44 1865 714591. Functional Ecology is covered by Cambridge Scientific Abstracts, BIOBASE/ Current Awareness in Biological Sciences (CABS), Current Contents and Science Citation Index.

Journal Customer Services: For ordering information, claims and any enquiry concerning your journal subscription, please go to www.wileycustomerhelp.com/ask or contact your nearest office. Europe, Middle East and Africa: E-mail: cs-journals@wiley.com Tel: +1781 388 8599 or 1 800 835 6770 (Toll free in USA and Canada); Asia Pacific: E-mail: cs-journals@wiley.com Tel: +65 6511 8000. Japan: For Japaness ppeaking under the spilles in the spi support, Email: cs-japan@wiley.com. Visit our Online Customer Get-Help available in 7 languages at www.wileycustomerhelp.com /ask

Information for subscribers: Functional Ecology is published in 12 issues per year. The institutional (online only) subscription prices for 2019 are: £1246 (UK), £1580 (Europe), \$2301 (the Americas) and \$2684 (Rest of World). Prices are exclusive of tax. Asia-Pacific GST, Canadian GST and European VAT will be applied at the appropriate rates. For more information on current tax rates, please go to www. wileyonlinelibrary.com/tax-vat. The price includes online access to the current content and all online back sales to 1 January 2015, where available. For other pricing options, including access information and terms and conditions, please visit www.wileyonlinelibrary.com/library-info/products/price-lists

Terms of use can be found here: https://onlinelibrary.wiley.com/terms-and-conditions

View this journal online at wileyonlinelibrary.com/journal/FEC

Visit wileyonlinelibrary.com to search the articles and register for table of contents Visit wileyonine of a founding member of the UN-backed HINARI, AGORA, and OARE initiatives. They are now collectively known as Research4Life, making online scientific content available free or at nominal cost to researchers in developing countries. Please visit Wiley's Content Access – Corporate Citizenship site: http:// www.wiley.com/WileyCDA/Section/id-390082.html

**Back issues:** Single issues from current and recent volumes are available at the current single issue price from cs-journals@wiley.com. Earlier issues may be obtained from Periodicals Service Company, 351 Fairview Avenue – Ste 300, Hudson, NY 12534, USA. Tel: +1 518 822-9300, Fax: +1 518 822-9305, Email: psc@periodicals.com

Copyright and copying (in any format): Functional Ecology © 2018 British Ecological Society. All rights reserved. No part of this publication may be reproduced, stored Society. All rights reserved. No part of this publication may be reproduced, stored or transmitted in any form or by any means without the prior permission in writing from the copyright holder. Authorization to copy items for internal and personal use is granted by the copyright holder for libraries and other users registered with their local Reproduction Rights Organisation (RRO), e.g. Copyright Clearance Center (CCC), 222 Rosewood Drive, Danvers, MA 01923, USA (www.copyright. com), provided the appropriate fee is paid directly to the RRO. This consent does not extend to other kinds of copying such as copying for general distribution, for advertising and promotional purposes, for creating new collective works or for create. Requising the first of a consent does not extend to other kinds of copying constant consent does not extend to other kinds of copying such as copying for general distribution, for advertising and promotional purposes, for creating new collective works or for the prior constant of the supervision of the supervision for advertising and promotional purposes. For creating new collective works or for the supervisions for such results and the supervision for advertising the prior for such results. resale. Permissions for such reuse can be obtained using the RightsLink 'Request Permissions' link on Wiley Online Library. Special requests should be addressed to: permissions@wilev.com

Wiley's Corporate Citizenship initiative seeks to address the environmental, social, economic, and ethical challenges faced in our business and which are important to our diverse stakeholder groups. Since launching the initiative, we have focused on sharing our content with those in need, enhancing community philanthropy, reducing our carbon impact, creating global guidelines and best practices for paper use, establishing a vendor code of ethics, and engaging our colleagues and other stakeholders in our efforts. Follow our progress at www.wiley.com/go/citizenship

Periodical ID Statement: FUNCTIONAL ECOLOGY, (ISSN 0269-8463) is published

**Online Open:** *Functional Ecology* accepts articles for Open Access publication. Please see www.functionalecology.org and select Author Guidelines for further information about OnlineOpen.

Disclaimer: The Publisher, British Ecological Society and Editors cannot be held responsible for errors or any consequences arising from the use of information contained in this journal; the views and opinions expressed do not necessarily reflect those of the Publisher, British Ecological Society and Editors, neither does the publication of advertisements constitute any endorsement by the Publisher, British Ecological Society and Editors of the products advertised.

Cover image: In the temperate forest of the Cantabrian Range (N Spain), small frugivores like blackcap Sylvia atricapilla mostly feed on plants of small fruits like elder Sambucus nigra. This size-matching increases the complementarity in frugivory networks, enhancing the function of seed dispersal at the landscape credit: Rolf Nussbaumer/Alamy. See paper by Garcia et al., pp. 2742-2752.

DOI: 10.1111/1365-2435.13213

# **RESEARCH ARTICLE**

# Functional Ecology

# Frugivore biodiversity and complementarity in interaction networks enhance landscape-scale seed dispersal function

Daniel García<sup>1</sup> | Isabel Donoso<sup>1,2</sup> | Javier Rodríguez-Pérez<sup>1,3</sup>

<sup>1</sup>Depto. Biología de Organismos y Sistemas, Universidad de Oviedo, and Unidad Mixta de Investigación en Biodiversidad (CSIC-Uo-PA), Oviedo, Spain

<sup>2</sup>Senckenberg Biodiversity and Climate Research Centre, Frankfurt (Main), Germany

<sup>3</sup>Centro de Investigação em Biodiversidade e Recursos Genéticos, Universidade de Évora, Évora, Portugal

## Correspondence

Daniel García Email: danielgarcia@uniovi.es

## **Funding information**

Ministerio de Economía y Competitividad, Grant/Award Number: CGL2011-28430, CGL2015-68963-C2-2-R and FPI-BES2012-052863

Handling Editor: Ruben Heleno

# Abstract

- 1. Animal biodiversity matters for the provision of ecosystem functions derived from trophic activity. However, the mechanisms underlying this pattern remain elusive since animal abundance and diversity, which are the components commonly used for representing biodiversity, provide poor information about ecological complementarity in species assemblages. An approach based on species interaction networks may overcome this constraint.
- 2. Here, we relate frugivore biodiversity and frugivore-plant network structure with landscape-scale seed dispersal function. We sampled, for two years, and at fourteen plots with variable assemblages of frugivores and plants in the Cantabrian Range (N Spain), data on the abundance and diversity of frugivorous birds, the consumption of fleshy fruits of woody plants and the landscape-scale patterns of avian seed deposition. As a measure of interaction complementarity in seed dispersal networks, we estimated the degree to which frugivore and plant species specialize in their interacting partners.
- 3. Specialization varied strongly across the seed dispersal networks of the different plots, being higher in networks harbouring smaller bird species that dispersed mostly small-fruited plants, and also in networks with late-ripening, dominant fruiting species dispersed mostly by wintering birds.
- 4. Bird abundance markedly affected seed deposition. Plots harbouring more birds received a higher density of dispersed seeds and showed higher probabilities of seed arrival and seed deposition in open microhabitats. Bird diversity also had a positive effect on the density of dispersed seed and, to a lesser extent, seed arrival probability. Independently of frugivore abundance and diversity, the density of dispersed seeds increased in plots where seed dispersal networks showed a higher degree of specialization.
- 5. This study considers the structure of interaction networks to re-address the relationship between biodiversity and ecosystem functionality, evidencing that specialization in frugivore-plant networks drives the large-scale process of seed dispersal. These results encourage the consideration of interaction complementarity as an underlying mechanism linking animal biodiversity and trophic-related functions.

#### KEYWORDS

avian seed dispersal, bird abundance, bird diversity, fleshy fruits, frugivorous birds, plantanimal interactions, specialization

# 1 | INTRODUCTION

The link between animal biodiversity and the ecosystem functions derived from animals' trophic activity is now widely accepted (Cardinale et al., 2006; Reiss, Bridle, Montoya, & Woodward, 2009). Species-rich and more complex animal assemblages result in stronger and more stable functions than those shrunk by ecological filtering or impoverished by defaunation (Duffy, 2003; Worm et al., 2006). These patterns involve a variety of trophic functions, from plant pollination by flower visitors (e.g., Fründ, Dormann, Holzschuh, & Tscharntke, 2013) to organic matter recycling by detritivores (e.g., Jonsson & Malmqvist, 2000). Although the positive effects of animal biodiversity on ecosystem functions are common, their underpinning mechanisms remain elusive (Reiss et al., 2009; Thompson et al., 2012). Among these mechanisms is niche complementarity through resource partitioning, whereby various species in a given assemblage have complementary manners of exploiting a set of trophic resources (Cardinale, 2011). Both theoretical modelling (e.g., Poisot, Mouquet, & Gravel, 2013) and experimental small-scale studies (e.g., Finke & Snyder, 2008) have evidenced the effects of resource partitioning on trophic functions. However, it is unclear whether these effects are prevalent in real-world ecosystems, as niche complementarity may be contingent on interspecific interactions (e.g., competition; Albrecht et al., 2013; Fründ et al., 2013) and on environmental filtering (Cardinale, 2011).

Using networks of interspecific interactions has been highlighted as a powerful tool for understanding the effects of animal biodiversity on trophic functions (Thompson et al., 2012). Ecological networks, like those representing animal-plant relationships (e.g., pollinators or seed dispersers and plants), assess the actual contribution of individual species to ecosystem functions (pollination and seed dispersal; Schleuning, Fründ, & García, 2015). More importantly, some measures of network complexity, like the degree of specialization which measures how species specialize in terms of their interacting partners, represent resource partitioning and functional complementarity (Blüthgen & Klein, 2010). Despite these advantages, the effects of interaction complementarity on the magnitude of ecosystem functions in the wild remain unclear (e.g., Macfadyen et al., 2009; Theodorou et al., 2017; but see Peralta, Frost, Rand, Didham, & Tylianakis, 2014).

The trophic interactions between frugivorous animals and fleshyfruited plants are suitable systems for evaluating the relevance of biodiversity in ecosystem functioning (Schleuning et al., 2015). By consuming fruits, frugivores shape plant communities through both antagonistic effects (e.g., when preying on pulp and seeds) and mutualistic ones (e.g., when acting as legitimate seed dispersers by regurgitating or defecating intact seeds; Jordano, 2014; Traveset, Heleno, & Nogales, 2014). Alongside plants, frugivores build ecological networks that vary greatly in specialization along environmental gradients at different scales (from landscape to latitudinal), depending mostly on changes in the abundance, richness and composition of animal and fruit resource assemblages (e.g., Schleuning et al., 2012; Albrecht et al., 2013; Vollstädt et al., 2018). Moreover, the abundance and the diversity of frugivores acting as legitimate seed dispersers have been identified as biodiversity components driving fine-scale patterns of seed dispersal (García & Martínez, 2012). Nevertheless, it is unclear whether these effects eventually scale up to landscapes at regional extents. More importantly, it is not known whether complementarity in frugivore-plant networks shapes variability in seed dispersal function.

Here, we focus on the assemblages of frugivorous birds and fleshy-fruited trees in the Cantabrian Range to assess the link between frugivore biodiversity and seed dispersal function, explicitly taking into account the structure of seed dispersal networks. By implementing a large-scale sampling of bird abundance and diversity, their frugivore activity and their derived seed deposition, we seek to: (a) estimate the degree of complementarity within seed dispersal networks and its local ecological determinants, (b) evaluate the relationship between frugivore abundance and diversity and seed dispersal function, in terms of quantitative and qualitative largescale patterns of community-wide seed deposition, and (c) assess the importance of interaction complementarity in the provision of the seed dispersal function relative to that of frugivore biodiversity. As a general prediction, we expect positive effects of both frugivore biodiversity and network complementarity on seed dispersal.

## 2 | MATERIALS AND METHODS

## 2.1 | Study design

This study was conducted in mid-elevation woodland pastures of the Cantabrian Range in northern Spain (for a comprehensive description of study system, design and all methodologies, see Supporting Information Appendix S1). The woodlands contain variable-sized patches of primary and secondary forest, embedded in an extensive matrix of stony meadows and heathland. The secondary forest is typically dominated by fleshy-fruited trees and shrubs with ripening periods from late summer to early winter. The main frugivores are passerine birds (García, 2016), with most species (e.g., thrushes) performing as legitimate seed dispersers, although some are pulpeaters and seed predators with a negligible contribution to the seed rain (Simmons et al., 2018). As the goal of this study was to relate the trophic activity of frugivorous birds with seed deposition, we used a



**FIGURE 1** Representation of study design. (a) Plot including 25 × 25 m grid cells, forest (pale blue) and bramble (brown) cover, bird count points (1–9 numbered red circles) and fruit consumption observation points (yellow dots). (b) Detail of individual trees (green circles: holly; red circles: hawthorn; black dots: other tree species). (c) Detail of stations for sampling dispersed seeds (different figures for different devices in pictures; green squares: open quadrats; red circles: pots in trees; yellow triangles: trays under bramble)

conservative approach and focused exclusively on bird species considered as legitimate seed dispersers, discarding observations not only of non-frugivorous species, but also of antagonistic frugivores. Classification of bird species was based on previous observations of fruit handling and faecal content (e.g., Martínez, García, & Obeso, 2008; see also Simmons et al., 2018).

In August 2012, we delimited fourteen 2.25-ha plots (150 × 150 m) in two sites (Sierra de Peña Mayor and Bandujo-Puertos de Marabio) in Asturias Province, Spain (Figure 1a; Supporting Information Table S1; Figure S1). All plots had similar vegetation types (forest stands embedded in a matrix of pastures and heathland), geomorphology (slope ≤25%, limestone substrate, altitude 990-1,250 m asl) and anthropic management (extensive livestock raising), but were chosen to incorporate wide variability in forest availability (3%-69% of forest cover; Supporting Information Table S1; Figure S1). We assumed that differences in forest cover would lead to high variability in the abundance and richness of fleshy fruits and frugivorous birds (as previously found in the same system; García & Martínez, 2012). Sampling was conducted across two consecutive fruit production and seed dispersal yearly events, from September to March in both 2012-2013 and 2013-2014 (sampling years, hereafter).

## 2.2 | Bird abundance and richness

Point-count bird censuses were carried out consecutively at nine regularly distributed points within each plot. To facilitate bird recording from these points, each plot was subdivided into 36 cells measuring  $25 \times 25$  m, each point being in the centre of a set of four adjacent cells (Figure 1a). Censuses were performed from 09.00 to 15.00, avoiding days of heavy rain and wind. In each census, all birds heard or seen within the set of four cells over a 5-min period were registered. Bird records were summed across points in each plot (total sampling effort of 45 min per plot per census), and nine census rounds were performed per sampling year and plot (1-2 censuses/ month from September to February). For frugivore species (legitimate seed dispersers only), we estimated absolute and relative abundances per year per plot. Frugivore richness was estimated as the number of species detected across all censuses, per year per plot. Frugivore diversity was estimated from reciprocal Simpson index ( $D = 1/\Sigma p_i^2$ ; where  $p_i$  is the relative abundance of species *i*; Loreau & Kinne, 2010).

## 2.3 | Fruit consumption

Bird fruit consumption was recorded in 17 rounds of 1-hr-per-plot observations throughout the period, though independent, of bird censuses (September-February): 8 rounds in 2012-2013 and 9 in 2013-2014. In each round, a given observer visited 3-4 vantage points (Figure 1a), chosen to ensure that the full extent of the plot was covered (i.e., including the nine points for bird censuses) as well as to focus on the different fruiting species present. Observers recorded every fruit consumption event (i.e., an individual bird consuming fruits) and every feeding bout (i.e., a single bird swallowing a single fruit) detected during the observation round.

## 2.4 | Fruit resource abundance

The production of fleshy fruits by woody plants was surveyed at the beginning of the fruiting season, providing an estimate of the yearly, community-wide, fruit resource base available for frugivorous birds. Depending on the different ripening peaks of the various species, we monitored fruit abundance in early September (early-season species) or mid-October (remaining species; Supporting Information Appendix S1) mapping all fruiting trees and shrubs in each plot (Figure 1b). For each individual tree or shrub, we visually estimated the number of fruits using a semi-logarithmic scale later extrapolated to a natural value of crop size (Supporting Information Appendix S1). For each plant species in each plot, we estimated production, both absolute (number of ripe fruits) and relative (number of fruits of the species/total number of fruits of all species). For each fleshy-fruited species, we measured fruit and seed traits from a sample of 25 ripe fruits (five from each of five individuals) collected in 2012-2013 (Supporting Information Appendix S1).

## 2.5 | Seed dispersal function

Avian seed dispersal was studied by identifying and counting the seeds deposited by frugivores, after regurgitation or defecation, during autumn and winter. This was based on data from sampling stations distributed in a grid scheme (108 stations per plot; Figure 1c) across three types of microhabitat (tree cover, fleshyfruited shrub cover and open area not covered by trees or shrubs) each involving a different type of device for seed collection, adapted to the physiognomy of the vegetation and the danger of trampling by ungulates: hanging plastic pots for tree cover, plastic trays under shrub cover and flag-labelled quadrats on the ground for open area (Figure 1c). Thus, number of stations per microhabitat differed in relation to relative cover of each microhabitat in each plot. All stations were set up in August 2012. Seed traps were checked for seed collection in February-March of 2013 and 2014, and open quadrats in late November and late February of 2012-2013 and 2013-2014. Collected seeds were identified (species are easily identifiable from external morphology) and counted in the laboratory, after oven-drying for one week at 70°C. The density of seeds (seeds/m<sup>2</sup>) deposited by birds at each sampling station was estimated taking into account the various surfaces of the different devices (0.07, 0.08 and 0.10 m<sup>2</sup> for, respectively, pots, trays and open quadrats).

Three different metrics (components) were used to represent community-wide seed dispersal function. To represent seed rain in quantitative terms, we estimated *seed density* (average density of dispersed seeds per sampling station per plot) and *seed arrival rate* (proportion of sampling stations per plot receiving dispersed seeds). As a qualitative metric, we used *seed arrival rate in open* (proportion of open-microhabitat sampling stations per plot receiving bird-dispersed seeds), which represents the potential for tree recolonization of deforested land (García & Martínez, 2012).

## 2.6 | Data analysis

## 2.6.1 | Building seed dispersal networks

Seed dispersal networks were based on frugivore-plant interaction matrices built from fruit consumption data. Although restrictive in terms of obtaining large samples of observed interactions (it requires large, often logistically unviable, efforts), this methodology was chosen because of its species unbiased character (García, 2016).

First, we pooled, for each plot, fruit consumption data from the different rounds and both sampling years, in order to reach suitable sampling effort (17 sampling rounds per site) and interaction sample size (number of consumption events and number of fruits consumed per plot ranging from 19 to 153 and 45 to 520, respectively) to ensure accurate estimates of the global metrics of interaction networks. To evaluate the sampling completeness of interaction networks, we constructed plot-level accumulation curves of the number of both bird and plant interacting species, and of the paired links between birds and plants, in relation to the number of sampling rounds and the number of fruit consumption events sampled; these completeness analyses suggested that our sampling effort was adequate to detect the richness of bird and plant interacting species and of bird-plant paired interactions (Supporting Information Appendix S2; Figures S2.1-6).

Second, we created plot-based matrices representing frugivore-plant interactions in terms of the number of seeds of each plant species dispersed by each bird species. For this, we estimated the number of dispersed seeds, for each plant species, by multiplying the cumulative number of fruits consumed by each bird species by the average number of seeds per fruit. The total number of interactions per plot, estimated from the total number of dispersed seeds, averaged 370.6 (±58.6 *SE*; min-max 92–786).

# 2.6.2 | Complementarity in seed dispersal networks and its ecological determinants

We quantified interaction complementarity by means of *complementary specialization* (sensu Blüthgen, 2010; Blüthgen & Klein, 2011), a measure of the degree of specialization among interacting species. In our case, it represents the degree to which frugivorous birds specialize in fruiting plant species as trophic resources as well as that to which plants specialize in bird species as their seed dispersal agents. It is thus a direct measure of segregation of the functional niche (and an inverse measure of functional niche overlap; Blüthgen, 2010). We used the index  $H_2'$  (standardized two-dimensional Shannon entropy; Blüthgen, Menzel & Blüthgen, 2006), based on the deviation of a species' realized number of interactions from that expected from each species' total number of interactions, which ranges from 0 (maximum generalization) to 1 (perfect specialization).

Compared to other network metrics,  $H_2'$  is demonstrably robust to biases induced by differences in sampling effort and network size (Blüthgen, Menzel, & Blüthgen, 2006; Fründ, McCann, & Williams, 2016). Specifically, one study of bird-plant pollination networks evidenced already stable H2' values already after 10 hr of sampling effort (Vizentin-Bugoni et al., 2016; see also Costa, da Silva, Ramos, & Heleno, 2016). Nevertheless, we standardized the raw values of specialization by estimating  $\Delta$ -transformed  $H_2'$ , based on null models ( $\Delta H_2' = H_2'$  observed –  $H_2'$  null models mean; Patefield model with 1,000 replicates; Dormann, Fründ, Blüthgen, & Gruber, 2009), as null-model standardization corrects potential overestimations in local specialization patterns due to small observation numbers (Fründ et al., 2016). Raw and standardized network metrics were estimated with the *Bipartite* package (Dormann et al., 2009), in the R 3.01.2 environment (R Development Core Team, 2014).

As local ecological determinants of complementarity in seed dispersal networks, we considered the abundance, richness and composition of frugivorous birds and fruiting plants, along with their matching traits (bird body mass and fruit size; Supporting Information Appendix S3), features known as intrinsic drivers of mutualistic networks (Dormann, Fründ, & Schaefer, 2017). The composition of both bird and fruiting plant communities was assessed using principal component analysis (PCA, stats package in R) of the relative abundance (fruit production in plants) of the different species across plots. Local composition values were estimated from the scores of the three first rotated factors in PCA. Bird body mass was used to estimate a plot-level, community-wide measure of bird size, by means of the community-weighted mean (CWM =  $\sum p_i \cdot d_i$ ; where  $p_i$  is the relative abundance of the bird species *i* and  $d_i$  is its body mass; Lavorel et al., 2008). A similar procedure, based on fruit diameter and relative fruit production, was applied to estimate CWM fruit size. Additionally, as potential determinants of complementarity, we also included a consumer/resource ratio, estimated from a plotlevel quotient of abundance of birds: abundance of fruits (assuming that low ratios may relax competition among frugivores leading to increased specialization; Albrecht et al., 2013), and forest cover (as a major environmental gradient; Chama, Berens, Downs, & Farwig, 2013). Relationships between standardized specialization and determinants were tested using Pearson's correlation tests.

# 2.6.3 | Frugivore biodiversity, interaction complementarity and seed dispersal

Our first goal was to explain the large-scale spatio-temporal variability of the different components of seed dispersal as a function of the abundance and diversity of frugivorous birds. For this, we considered the data from the various plots in each sampling year. We used generalized linear mixed models (GLMMs) with the components of seed dispersal as different response variables (log- or arcsin-sqrt-transformed when needed), considering Gaussian error distributions and identity links. All models incorporated, as fixed effects, both abundance and diversity of birds, standardized before inclusion. Simpson index was preferred to bird richness to represent diversity because of the significant collinearity between abundance and richness (Pearson's correlation coefficient, r = 0.426, p = 0.024, n = 28), but not between abundance and diversity (r = 0.293, p = 0.130, n = 28). Simpson index thus accounted for both evenness

and richness (correlation diversity—richness; r = 0.515, p = 0.005, n = 28) and represented an "effective number of species" in the community (the number of species expected in a situation of equal species abundances; Loreau & Kinne, 2010). Plot identity (nested within site) was included in all models as a random effect, whereas year (2012–2013; 2013–2014) and site (Sierra de Peña Mayor, Bandujo-Puertos de Marabio) were included as fixed effects (due to low number of levels). Year and site factors were excluded from final models when proven initially non-significant (p > 0.05). Degree of variance explained by the final complete model and by fixed effects was estimated from conditional and marginal  $R^2$  values, respectively (Nakagawa & Schielzeth, 2013).

The second step sought to test the simultaneous effects of interaction complementarity and frugivore biodiversity on the provision of seed dispersal function. Given that network metrics were based on interaction matrices that pooled data from both sampling years, we recalculated the local values of frugivore biodiversity and seed dispersal function per plot also from data pooled across years. We used generalized linear models (GLMs) to check for effects of abundance and diversity of frugivorous birds and of standardized specialization (predictors) on each component of seed dispersal (response variables). Predictors were not correlated between themselves (|r| < 0.40, p > 0.15, n = 14; for all paired combinations). We built models with different combinations of predictors, setting the number of predictors per model at two in order to avoid model overparametrization. Inferences about the relative effects of the different predictors were based on effect size and significance (predictors were standardized before inclusion), and Akaike's information criterion (with correction for small sample sizes, AIC<sub>c</sub>) value was used to identify the most informative model (i.e., that with the lowest AIC, value; Quinn & Keough, 2002). All response variables met normality requirements (after log- or arcsin-sqrt-transformed, when needed) so models considered Gaussian error distributions and identity links. GLMM and GLM analyses were performed with Ime4 package in R.

## 3 | RESULTS

#### 3.1 | General overview

Bird censuses provided 17,732 observations, of which 68.58% corresponded to ten species of legitimate seed dispersers, blackbird *Turdus merula*, redwing *T. iliacus* and European robin *Erithacus rubecula* being the most frequent and abundant (Supporting Information Appendix S3; Table S3). Ten fruiting woody plant species were recorded (Table S3), the most frequent and abundant being holly *llex aquifolium* and hawthorn *Crataegus momogyna* (Supporting Information Table S3). Due to differences in species occurrence and abundances, the composition of bird and fruiting plant assemblages varied markedly across plots (Supporting Information Appendix S4; Table S4, Figure S4).

Ten bird species and seven fleshy-fruited plants were detected in fruit consumption observations. Plot-based interaction matrices varied in size, with the number of species of frugivorous birds, fruiting



**FIGURE 2** Examples of local seed dispersal networks with increasing levels of interaction complementarity (values of standardized degree of specialization  $\Delta H_2$ ' are shown). Bipartite graphs show the proportion of seeds (left column) dispersed by frugivorous birds (right column), and the proportion of seeds of each plant consumed by each bird species (grey links). Species codes are based on abbreviated scientific names (Supporting Information Appendix S3; Table S3). Bird species in right panel are scaled to highlight size differences (artwork: Daniel García)

plants and both birds and plants ranging, respectively, 4–7, 2–5 and 6–11 (Appendix S5; Figure S5). The proportion of observed links between birds and plants, from all potential paired bird–plant combinations per plot, averaged 0.56 (±0.04 *SE*; min–max: 0.35–0.88; Figure S5). In terms of seed dispersal (i.e., number of dispersed seeds estimated from plot-based interaction matrices), holly and haw-thorn–mostly by blackbird and redwing–accounted for 67.11% of all observed interactions.

We recorded seed deposition of fleshy-fruited plants in 52.38% of sampling stations (n = 1512 stations/year), mean density being 222.46 (±11.47 SE) seeds/m<sup>2</sup> per station per year. Eight plant species were recorded in the sampling of the seed rain, which was heavily dominated by holly, hawthorn and bramble *Rubus fruticosus/ulmifo-lius* seeds (Appendix S3; Table S3).

## 3.2 | Complementarity in seed dispersal networks

The degree of specialization in seed dispersal networks varied markedly across plots, with raw  $H_2'$  values ranging from 0.164 to 0.965 (Figure 2; Supporting Information Appendix S5; Figure S5). Poorly specialized networks showed species interacting with many and equally frequent partners, on both the frugivore and the plant side (Figure 2). Conversely, highly specialized networks showed frequent segregation, especially on the bird side, as species seldom overlapped in terms of the plants with which they interacted (Figure 2).  $H_2'$  raw values significantly differed from null model means in all plots (t < -29.66, p < 0.001, in all cases) and were positively correlated with  $\Delta H_2'$  standardized values (r = 0.997, p < 0.001, n = 14).  $H_2'$ was independent of the number of interacting species of birds or plants per plot (|r| < 0.442, p > 0.113, n = 14), the proportion of realized links (r = 0.300, p = 0.296, n = 14) and the number of interactions (i.e., dispersed seeds; r = -0.076, p = 0.794, n = 14).

Concerning the ecological determinants of interaction complementarity, standardized specialization degree was unaffected by local abundance or richness of either frugivorous birds (|r| < 0.128, p > 0.666, n = 14; in both cases) or fruiting plants (|r| < 0.368, p > 0.196, n = 14; in both cases). Specialization was higher in plots with larger relative abundances of blackcap Sylvia atricapilla, chiffchaff Phylloscopus collybita/ibericus and European jay Garrulus glandarius (as represented by the bird PCA3 factor; r = 0.712; p = 0.004, n = 14; Figure 3; Supporting Information Appendix S4; Figure S4), as well as in plots with smaller CWM bird body size (r = -0.552; p = 0.041, n = 14; Figure 3), but it was unaffected by other bird composition trends (PCA1 and PCA2 factors: |r| < 0.194, p > 0.507, n = 14, in both cases). Specialization was also higher in plots with fruit crops dominated by holly (fruiting plants PCA1: r = 0.535; p = 0.048, n = 14; Figure 3; Supporting Information Figure S4), but was unaffected by other fruit composition trends (fruiting plants PCA2 and PCA3) or CWM fruit size (|r| < 0.368, p > 0.196, n = 14, in all cases). Neither consumer/resource ratio nor forest cover affected specialization (|r| < 0.448, p > 0.10, n = 14, in both cases).

# 3.3 | Frugivore biodiversity, interaction complementarity and seed dispersal

When analysed by years using GLMMs, both abundance and diversity of frugivorous birds had an independent positive effect on seed dispersal function, although their respective effects were contingent on the specific seed dispersal component. The density of dispersed seeds was higher in plots with higher values for abundance and diversity of frugivores (Table 1; Figure 4). Seed arrival rate followed a similar pattern, although in this case bird abundance showed a stronger effect than that of frugivore diversity (Table 1; Figure 4). Rate of seed arrival in open microhabitats was exclusively affected by abundance of frugivorous



**FIGURE 3** Responses of interaction complementarity (standardized degree of specialization  $\Delta H_2'$ ) to the composition of the communities of frugivorous birds and fruiting plants (axes of principal component analysis) and frugivore traits (community-weighted mean of body mass). Dots represent different plots (data from both years pooled)

**TABLE 1** Generalized linear mixed models for the effects of abundance and diversity of frugivorous birds (fixed effects) on different components of seed dispersal, with site and year included as fixed effects (removed from final models when p > 0.05), and plot identity (nested within site) included as a random effect. Values of marginal and conditional (in brackets)  $R^2$  are also shown

Estimate	SE	t-Value	p-Value
0.135	0.025	5.325	<0.0001
0.137	0.027	4.951	<0.0001
0.426	0.038	11.02	<0.0001
Variance	SD		
0.009	0.099		
Estimate	SE	t-Value	p-Value
0.105	0.018	5.820	<0.0001
0.073	0.019	3.712	0.0035
0.076	0.027	2.84	0.015
Variance	SD		
0.005	0.075		
Estimate	SE	t-Value	p-Value
0.064	0.022	2.860	0.0135
-0.022	0.021	-1.015	0.5360
Variance	SD		
0.009	0.094		
	Estimate 0.135 0.137 0.426 Variance 0.009 Estimate 0.105 0.073 0.076 Variance 0.005 Estimate 0.064 -0.022 Variance 0.009	EstimateSE0.1350.0250.1370.0270.4260.038VarianceSD0.0090.099Conos0.0190.1050.0180.0730.0190.0760.027VarianceSD0.0050.027VarianceSD0.0050.027VarianceSD0.0050.021VarianceSE0.0640.0220.0210.021VarianceSD0.0090.094	EstimateSEt-Value0.1350.0255.3250.1370.0274.9510.4260.03811.02VarianceSD0.0090.099EstimateSEt-Value0.1050.0185.8200.0730.0193.7120.0760.0272.84VarianceSD0.0050.075EstimateSEt-Value0.0050.0212.8600.0640.021-1.015VarianceSD0.064SD2.8600.0220.021-1.015VarianceSD0.0090.094

birds, with a greater proportion of sampling stations in the open receiving seeds in those plots hosting higher numbers of birds (Table 1).

When applied to the data of the pooled sampling years, GLMs evidenced that interaction complementarity positively affected seed dispersal function, even after accounting for the effects of abundance and/or diversity of frugivorous birds (Table 2; Figure 5). Namely, seed density was best explained by the combined positive effects of frugivore abundance and specialization (Table 2; Supporting Information Appendix S6; Table S6). Concerning seed arrival rate and seed arrival in the open, GLMs for pooled data suggested effects of frugivore abundance and diversity similar to those evidenced by previous yearly models (Table 2; Supporting Information Table S6).

# 4 | DISCUSSION

This study addresses the link between frugivore biodiversity and the ecosystem function of seed dispersal, by considering the structure of



**FIGURE 4** Effects of abundance (cumulative number of individuals per plot) and diversity (Simpson Index) of frugivorous birds on the density of dispersed seeds (No. of dispersed seeds per square metre) and the probability of seed arrival (proportion of sampling stations receiving dispersed seeds) for different plots and years

**TABLE 2** Generalized linear models for the effects of frugivore biodiversity and interaction complementarity on seed dispersal. For each seed dispersal component, the combination of predictors with the lowest AIC<sub>c</sub> value is shown. Model and null deviance values are also shown

Seed density (log <sub>10</sub> )						
	AIC <sub>c</sub>	Model Dev.	Null Dev.			
	-13.54	0.619	0.767			
Predictors	Estimate	SE	t-Value	p-Value		
Abundance of birds	0.174	0.032	5.37	0.0002		
Specialization ( $\Delta H_2'$ )	0.110	0.032	3.42	0.0057		
Seed arrival rate						
	AIC <sub>c</sub>	Model Dev.	Null Dev.			
	-20.46	0.377	0.468			
Predictors	Estimate	SE	t-Value	p-Value		
Abundance of birds	0.128	0.027	4.66	0.0007		
Diversity of birds	0.072	0.028	2.61	0.0240		
Seed arrival rate in open (arcsin sqrt)						
	AIC <sub>c</sub>	Model Dev.	Null Dev.			
	-12.46	0.110	0.270			
Predictors	Estimate	SE	t-Value	p-Value		
Abundance of birds	0.074	0.033	2.25	0.0457		
Specialization ( $\Delta H_2'$ )	-0.059	0.034	-1.75	0.1100		



**FIGURE 5** Effects of abundance (cumulative number of individuals per plot) of frugivorous birds and interaction complementarity (standardized degree of specialization  $\Delta H_2$ ') on the density of dispersed seeds. Dots represent different plots (data from both years pooled)

interaction networks. Although in the long term, by driving plant community dynamics (e.g., Martínez & García, 2017), seed dispersal affects frugivore and fruit assemblages and hence their interaction networks (i.e., the seed dispersal loop; Wang & Smith, 2002), here we focus on the assumed causal relationship between frugivore-plant community attributes and their short-term ecological consequence, seed deposition. As such, we applied an observational approach that incorporates a set of local landscapes harbouring independent gradients of frugivore abundance and diversity, and also matches these community attributes with the fine-scale processes of fruit consumption and seed deposition. We evidence positive responses of seed dispersal process to increases in abundance and diversity of frugivores across landscapes. We also found strong differences in interaction complementarity in frugivore-plant networks, mostly determined by the composition of frugivore and plant assemblages and frugivore size. Importantly, interaction complementarity positively affected the local magnitude of seed dispersal, independent of frugivore abundance and diversity.

## 4.1 | Complementary in seed dispersal networks

Our results reveal large-scale variability in interaction complementarity, estimated from specialization, as shown by other studies over regional extents (e.g., Chama et al., 2013). Local specialization values differed significantly from those expected by the random association of interacting frugivores and plants, suggesting that interaction complementarity resulted from ecological determinants (Dormann et al., 2017). In fact, specialization varied mostly as a response to local differences in bird composition and traits: more specialized networks occurring where common and small-sized bird species, such as blackcap and European robin, feed mostly on small-fruit plants like bramble and elder *Sambucus nigra* (Figure 2). These specialized interactions may derive from morphological matching, that is small birds avoiding large fruits due to gape width constraints (González-Castro, Yang, Nogales, & Carlo, 2015). Some kind of phenological coupling (González-Castro et al., 2015) may also occur, as bramble and elder are typically late summer ripening species, a time when blackcap is still abundant in Cantabrian uplands (Santos et al., 2014). Conversely, the occurrence of abundant and medium-to-large size birds-like blackbird and song thrush T. philomelos-with diversified fruit diets (thanks to their larger gape width relative to fruit diameter [González-Castro et al., 2015]) would lead to highly generalized networks (Figure 2). Thus, specialization in networks would mostly reflect frugivore trophic complementarity (less overlap in the trophic niche determined by the quantity and the species of fruits consumed; Blüthgen & Klein, 2011). Indeed, in our case, network complementarity was negatively correlated with niche overlap among bird species (r = -0.81, p < 0.001, n = 14; estimated from bipartite). Some variability in specialization was also attributable to the composition of fruiting plants, that is the presence of holly, a crop-dominant but late-ripening species, leading to interactions almost monopolized, at least in some plots, by wintering frugivores like redwing (Figure 2; see also Martínez et al., 2008).

# 4.2 | Frugivore biodiversity, interaction complementarity and seed dispersal

Our yearly assessment showed the positive, independent and consistent relationships of both abundance and diversity of frugivorous birds with different components of the seed dispersal function. Namely, those landscapes harbouring more frugivores from a greater number of species received a denser and wider seed rain of fleshyfruited woody plants. We thus evidence how the patterns already seen at finer scales in the study system (García & Martínez, 2012) also scale up at the regional extent and prevailed across years.

We found, on the one hand, that the effects of frugivore abundance on seed dispersal were stronger than those of frugivore diversity, suggesting that this ecosystem function is more dependent on the number of individual animals providing it than on the qualitative functional differences between animals (see also, for pollination by insects, Winfree, Fox, Williams, Reilly, & Cariveau, 2015). On the other hand, frugivore biodiversity effects varied across components of seed dispersal, which may be partly associated with the occurrence of complex interactions between frugivore abundance and diversity (e.g., stronger effects of diversity at small abundances, Rumeu et al., 2017), though the present data do not facilitate testing this (but see Rodríguez-Pérez, García, Martínez, & Morales, 2017). Even so, our results showed positive effects of diversity irrespective of abundance, probably related to sampling effects (the incorporation of functionally dominant frugivores, Schleuning et al., 2015), spatial complementarity (with a greater diversity meaning a higher variety of spatial behaviours after fruit consumption; Morales, García, Martínez, Rodríguez-Pérez, & Herrera, 2013) or even interspecific interactions (with a greater diversity leading to facilitation in, e.g., fruit resource tracking; Donoso, García, Martínez, Tylianakis, & Stouffer, 2017).

Notably, by incorporating the effects of interaction complementarity the present study goes beyond previous evaluations of the role of frugivore biodiversity on seed dispersal. In line with our expectation, we found that specialization also drove seed dispersal function and, when controlling for the effects of frugivore abundance, it explained the regional variability in the density of dispersed seeds better than frugivore diversity. We would suggest that trophic (diet) complementarity between frugivorous birds underpins these effects. Different birds specializing in different plants would facilitate more effective access to available plant resources, increasing the average number of seeds per deposition point (for such a case with pollinators, see Fontaine, Dajoz, Meriguet, & Loreau, 2006). This seems particularly likely to occur when specialization leads to the inclusion of big-crop plant species (like bramble, Supporting Information Table S2). Moreover, increased seed density is also expected when frugivores differ in their spatial patterns of foraging, because they feed selectively on fruiting species that occur in different sectors of the landscape (e.g., bramble shrubs mostly occurs at forest fringes in the study plots, Figure 1a).

# 5 | CONCLUDING REMARKS

Our study re-addresses the relationship between biodiversity and ecosystem functions by explicitly considering the structure of trophic networks. We show, under unmanipulated conditions and at the large scale, a positive link between the degree of network-wide specialization and the magnitude of a function derived from the animal-plant mutualism (for small-scale, microcosm experiments with pollination, see Fontaine et al., 2006; Fründ et al., 2013). These results thus reinforce previous observational findings concerning other types of food webs (e.g., host-parasitoids, Peralta et al., 2014), encouraging the consideration of interaction complementarity as an underlying mechanism of the causal link between animal biodiversity and trophic-related ecosystem functions and services.

## ACKNOWLEDGEMENTS

We thank Carlos Guardado, Juan Rodríguez, Marga García, Lucía García, Diego Méndez, David Luna, René Pérez, Daniel Martínez, Jorge Álvarez and José A. Molina for technical support and Ronnie Lendrum for her review of the English. The Associate Editor and two anonymous referees provided helpful suggestions for improving the manuscript. The research was funded by Spanish MinECo/FEDER grants CGL2011-28430 and CGL2015-68963-C2-2-R, and FPI-BES2012-052863 grant to I.D. Fieldwork was conducted with the permission of the Government of Asturias.

## AUTHORS' CONTRIBUTIONS

D.G. conceived and designed the study; D.G., J.R.-P. and I.D. collected the data; D.G. and I.D. analysed the data; and D.G. wrote the manuscript. All authors revised the manuscript and approved the final version.

## DATA ACCESSIBILITY

Original data associated with this article are deposited in the Dryad Digital Repository https://doi.org/10.5061/dryad.h748c88 (García, Donoso, & Rodríguez-Pérez, 2018).

## ORCID

Daniel García D http://orcid.org/0000-0002-7334-7836 Isabel Donoso D http://orcid.org/0000-0002-0287-9026 Javier Rodríguez-Pérez D http://orcid.org/0000-0002-1292-9090

## REFERENCES

- Albrecht, J., Berens, D. G., Blüthgen, N., Jaroszewicz, B., Selva, N., & Farwig, N. (2013). Logging and forest edges reduce redundancy in plant-frugivore networks in an old-growth European forest. *Journal* of Ecology, 101, 990–999. https://doi.org/10.1111/1365-2745.12105
- Blüthgen, N. (2010). Why network analysis is often disconnected from community ecology: A critique and an ecologist's guide. *Basic and Applied Ecology*, 11, 185–195. https://doi.org/10.1016/j.baae.2010.01.001
- Blüthgen, N., & Klein, A. M. (2011). Functional complementarity and specialisation: The role of biodiversity in plant-pollinator interactions. *Basic and Applied Ecology*, 12, 282–291. https://doi.org/10.1016/j. baae.2010.11.001
- Blüthgen, N., Menzel, F., & Blüthgen, N. (2006). Measuring specialization in species interaction networks. BMC Ecology, 6, 9.
- Cardinale, B. J. (2011). Biodiversity improves water quality through niche partitioning. Nature, 472, 86–89. https://doi.org/10.1038/nature09904
- Cardinale, B. J., Srivastava, D. S., Duffy, J. E., Wright, J. P., Downing, A. L., Sankaran, M., & Jouseau, C. (2006). Effects of biodiversity on the functioning of trophic groups and ecosystems. *Nature*, 443, 989–992. https://doi.org/10.1038/nature05202
- Chama, L., Berens, D. G., Downs, C. T., & Farwig, N. (2013). Habitat characteristics of forest fragments determine specialisation of plant-frugivore networks in a mosaic forest landscape. *PloS ONE*, *8*, e54956. https://doi.org/10.1371/journal.pone.0054956
- Costa, J. M., da Silva, L. P., Ramos, J. A., & Heleno, R. H. (2016). Sampling completeness in seed dispersal networks: When enough is enough. *Basic and Applied Ecology*, 17, 155–164. https://doi.org/10.1016/j. baae.2015.09.008
- Donoso, I., García, D., Martínez, D., Tylianakis, J. M., & Stouffer, D. B. (2017). Complementary effects of species abundances and ecological neighborhood on the occurrence of fruit-frugivore interactions. *Frontiers in Ecology and Evolution*, 5, 133. https://doi.org/10.3389/ fevo.2017.00133
- Dormann, C. F., Fründ, J., Blüthgen, N., & Gruber, B. (2009). Indices, graphs and null models: Analyzing bipartite ecological networks. *The Open Ecology Journal*, 2, 7–24. https://doi.org/10.2174/1874213000902010007
- Dormann, C. F., Fründ, J., & Schaefer, H. M. (2017). Identifying causes of patterns in ecological networks: Opportunities and limitations. *Annual Review of Ecology, Evolution, and Systematics*, 48, 559–584. https://doi.org/10.1146/annurev-ecolsys-110316-022928
- Duffy, J. E. (2003). Biodiversity loss, trophic skew and ecosystem functioning. *Ecology Letters*, *6*, 680–687. https://doi. org/10.1046/j.1461-0248.2003.00494.x
- Finke, D. L., & Snyder, W. E. (2008). Niche partitioning increases resource exploitation by diverse communities. *Science*, 321, 1488–1490. https://doi.org/10.1126/science.1160854
- Fontaine, C., Dajoz, I., Meriguet, J., & Loreau, M. (2006). Functional diversity of plant-pollinator interaction webs enhances the persistence of plant communities. *PLoS Biology*, *4*, e1.
- Fründ, J., Dormann, C. F., Holzschuh, A., & Tscharntke, T. (2013). Bee diversity effects on pollination depend on functional complementarity and niche shifts. *Ecology*, 94, 2042–2054. https://doi.org/10.1890/12-1620.1
- Fründ, J., McCann, K. S., & Williams, N. M. (2016). Sampling bias is a challenge for quantifying specialization and network structure: Lessons from a quantitative niche model. *Oikos*, 125, 502–513. https://doi.org/10.1111/oik.02256

- García, D. (2016). Birds in ecological networks: Insights from birdplant mutualistic interactions. Ardeola, 63, 151–180. https://doi. org/10.13157/arla.63.1.2016.rp7
- García, D., Donoso, I., & Rodríguez-Pérez, J. (2018). Data from: Frugivore biodiversity and complementarity in interaction networks enhance landscape-scale seed dispersal function. *Dryad Digital Repository*, https://doi.org/10.5061/dryad.h748c88.
- 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 B: Biological Sciences*, 279, 3106–3013.
- González-Castro, A., Yang, S., Nogales, M., & Carlo, T. A. (2015). Relative importance of phenotypic trait matching and species' abundances in determining plant-avian seed dispersal interactions in a small insular community. AoB Plants, 7, plv017. https://doi.org/10.1093/aobpla/plv017
- Jonsson, M., & Malmqvist, B. (2000). Ecosystem process rate increases with animal species richness: Evidence from leaf-eating, aquatic insects. Oikos, 89, 519–523. https://doi.org/10.1034/j.1600-0706.2000.890311.x
- Jordano, P. (2014). Fruits and frugivores. In R. S. Gallagher (Ed.), Seeds: The ecology of regeneration in plant communities (3rd ed., pp. 18–62). Wallingford, UK: CAB International.
- Lavorel, S., Grigulis, K., McIntyre, S., Williams, N. S., Garden, D., Dorrough, J., ... Bonis, A. (2008). Assessing functional diversity in the field-methodology matters!. *Functional Ecology*, 22, 134–147.
- Loreau, M., & Kinne, O. (2010). *The challenges of biodiversity science* (Vol. 17). Oldendorf/Luhe, Germany: International Ecology Institute.
- Macfadyen, S., Gibson, R., Polaszek, A., Morris, R. J., Craze, P. G., Planqué, R., ... Memmott, J. (2009). Do differences in food web structure between organic and conventional farms affect the ecosystem service of pest control?. *Ecology Letters*, 12, 229–238. https://doi. org/10.1111/j.1461-0248.2008.01279.x
- Martínez, D., & García, D. (2017). Role of avian seed dispersers in tree recruitment in woodland pastures. *Ecosystems*, 20, 616–629. https:// doi.org/10.1007/s10021-016-0043-6
- Martínez, I., García, D., & Obeso, J. R. (2008). Differential seed dispersal patterns generated by a common assemblage of vertebrate frugivores in three fleshy-fruited trees. *Écoscience*, 15, 189–199. https:// doi.org/10.2980/15-2-3096
- Morales, J. M., García, D., Martínez, D., Rodríguez-Pérez, J., & Herrera, J. M. (2013). Frugivore behavioural details matter for seed dispersal: A multispecies model for Cantabrian thrushes and trees. *PLoS ONE*, 6, e65216.
- Nakagawa, S., & Schielzeth, H. (2013). A general and simple method for obtaining R2 from generalized linear mixed-effects models. *Methods* in Ecology and Evolution, 4, 133–142.
- Peralta, G., Frost, C. M., Rand, T. A., Didham, R. K., & Tylianakis, J. M. (2014). Complementarity and redundancy of interactions enhance attack rates and spatial stability in host-parasitoid food webs. *Ecology*, 95, 1888–1896. https://doi.org/10.1890/13-1569.1
- Poisot, T., Mouquet, N., & Gravel, D. (2013). Trophic complementarity drives the biodiversity–ecosystem functioning relationship in food webs. *Ecology Letters*, 16, 853–861. https://doi.org/10.1111/ele.12118
- Quinn, G. P., & Keough, M. J. (2002). Experimental design and data analysis for biologists. Cambridge, UK: Cambridge University Press.
- R Core Team (2014). R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing. Retrieved from https://www.R-project.org/
- Reiss, J., Bridle, J. R., Montoya, J. M., & Woodward, G. (2009). Emerging horizons in biodiversity and ecosystem functioning research. *Trends in Ecology* & Evolution, 24, 505–514. https://doi.org/10.1016/j.tree.2009.03.018
- Rodríguez-Pérez, J., García, D., Martínez, D., & Morales, J. M. (2017). Seed dispersal by changing frugivore assemblages: A mechanistic test of global change effects. Oikos, 126, 672–681. https://doi.org/10.1111/ oik.03176
- Rumeu, B., Devoto, M., Traveset, A., Olesen, J. M., Vargas, P., Nogales, M., & Heleno, R. (2017). Predicting the consequences of disperser extinction: Richness matters the most when abundance is low. *Functional Ecology*, 31, 1910–1920. https://doi.org/10.1111/1365-2435.12897

- Santos, T., Carbonell, R., Galarza, A., Perez-Tris, J., Ramirez, A., & Tellería, J. L. (2014). The importance of northern Spanish farmland for wintering migratory passerines: A quantitative assessment. *Bird Conservation International*, 24, 1–16. https://doi.org/10.1017/ S0959270913000191
- Schleuning, M., Fründ, J., & García, D. (2015). Predicting ecosystem functions from biodiversity and mutualistic networks: An extension of trait-based concepts to plant-animal interactions. *Ecography*, 38, 380–392. https://doi.org/10.1111/ecog.00983
- Schleuning, M., Fründ, J., Klein, A. M., Abrahamczyk, S., Alarcón, R., Albrecht, M., ... Dalsgaard, B. (2012). Specialization of mutualistic interaction networks decreases toward tropical latitudes. *Current Biology*, 22, 1925–1931. https://doi.org/10.1016/j.cub.2012.08.015
- Simmons, B. I., Sutherland, W. J., Dicks, L. V., Albrecht, J., Farwig, N., García, D., ... González-Varo, J. P. (2018). Moving from frugivory to seed dispersal: Incorporating the functional outcomes of interactions in plant-frugivore networks. *Journal of Animal Ecology*, 87, 995–1007. https://doi.org/10.1111/1365-2656.12831
- Theodorou, P., Albig, K., Radzevičiūtė, R., Settele, J., Schweiger, O., Murray, T. E., & Paxton, R. J. (2017). The structure of flower visitor networks in relation to pollination across an agricultural to urban gradient. *Functional Ecology*, 31, 838–847. https://doi.org/10.1111/1365-2435.12803
- Thompson, R. M., Brose, U., Dunne, J. A., Hall, R. O. Jr, Hladyz, S., Kitching, R. L., ... Tylianakis, J. M. (2012). Food webs: Reconciling the structure and function of biodiversity. *Trends in Ecology & Evolution*, 27, 689–697. https://doi.org/10.1016/j.tree.2012.08.005
- Traveset, A., Heleno, R., & Nogales, M. (2014). The ecology of seed dispersal. In R. S. Gallagher (Ed.), Seeds: The ecology of regeneration in plant communities (3rd ed., pp. 62–93). Wallingford, UK: CAB International.
- Vizentin-Bugoni, J., Maruyama, P. K., Debastiani, V. J., Duarte, L. D. S., Dalsgaard, B., & Sazima, M. (2016). Influences of sampling effort on detected patterns and structuring processes of a Neotropical plant-hummingbird network. *Journal of Animal Ecology*, 85, 262–272. https://doi.org/10.1111/1365-2656.12459
- Vollstädt, M. G. R., Ferger, S. W., Hemp, A., Howell, K. M., Böhning-Gaese, K., & Schleuning, M. (2018). Seed-dispersal networks respond differently to resource effects in open and forest habitats. *Oikos*, 127, 847–854. https://doi.org/10.1111/oik.04703
- Wang, B. C., & Smith, T. B. (2002). Closing the seed dispersal loop. Trends in Ecology & Evolution, 17, 379–386. https://doi.org/10.1016/ S0169-5347(02)02541-7
- Winfree, R., Fox, J., Williams, N. M., Reilly, J. R., & Cariveau, D. P. (2015). Abundance of common species, not species richness, drives delivery of a real-world ecosystem service. *Ecology Letters*, 18, 626–635. https://doi.org/10.1111/ele.12424
- Worm, B., Barbier, E. B., Beaumont, N., Duffy, J. E., Folke, C., Halpern, B. S., ... Sala, E. (2006). Impacts of biodiversity loss on ocean ecosystem services. *Science*, 314, 787–790. https://doi.org/10.1126/science.1132294

#### SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: García D, Donoso I, Rodríguez-Pérez J. Frugivore biodiversity and complementarity in interaction networks enhance landscape-scale seed dispersal function. *Funct Ecol.* 2018;32:2742-2752. <u>https://doi.org/10.1111/1365-</u> 2435.13213