

Limited potential for bird migration to disperse plants to cooler latitudes

<https://doi.org/10.1038/s41586-021-03665-2>

Received: 15 July 2020

Accepted: 21 May 2021

Published online: 23 June 2021

 Check for updates

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Climate change is forcing the redistribution of life on Earth at an unprecedented velocity^{1,2}. Migratory birds are thought to help plants to track climate change through long-distance seed dispersal^{3,4}. However, seeds may be consistently dispersed towards cooler or warmer latitudes depending on whether the fruiting period of a plant species coincides with northward or southward migrations. Here we assess the potential of plant communities to keep pace with climate change through long-distance seed dispersal by migratory birds. To do so, we combine phenological and migration information with data on 949 seed-dispersal interactions between 46 bird and 81 plant species from 13 woodland communities across Europe. Most of the plant species (86%) in these communities are dispersed by birds migrating south, whereas only 35% are dispersed by birds migrating north; the latter subset is phylogenetically clustered in lineages that have fruiting periods that overlap with the spring migration. Moreover, the majority of this critical dispersal service northwards is provided by only a few Palearctic migrant species. The potential of migratory birds to assist a small, non-random sample of plants to track climate change latitudinally is expected to strongly influence the formation of novel plant communities, and thus affect their ecosystem functions and community assembly at higher trophic levels.

Anthropogenic climate change is forcing the redistribution of life on Earth at an unprecedented rate^{1,2}. The distribution of organisms is constrained by the climatic conditions that they can tolerate (known as their climatic envelope)⁵. Driven by global warming, climatic envelopes are shifting towards higher (cooler) latitudes^{6,7}. The mean global velocity at which organisms need to shift their distributional range to retain the same temperatures has been estimated at 4.2 km per decade, although estimates exceed 100 km per decade in some regions^{2,8}. A crucial question is whether species and locally adapted genotypes will be able to move sufficiently fast to track a rapidly changing climate, which depends on their dispersal capacities^{5,7,9,10}. Tackling this issue is key for understanding and predicting the effects of climate change on biological communities and the ecosystem functions that they mediate, including those that affect human welfare and even climate itself (via vegetation shifts)¹¹.

Plants are the cornerstone of terrestrial ecosystems, but there is a major knowledge gap regarding their dispersal abilities and latitudinal range shifts under current rates of global warming^{5–7,9–11}. Dispersal beyond range edges is necessary for plant species to colonize novel areas that become suitable owing to climate change^{6,12}, whereas dispersal within

species ranges allows the immigration of genotypes from warm-adapted populations to cooler areas that are becoming warmer¹³. However, plants are sessile and the dispersal of their seeds (the process that allows new individuals to recruit far away) generally occurs within 1 km of source plants^{14–17}. Although local dispersal is crucial for plant recruitment, it is clearly insufficient to track current climate change, particularly in plants with generation times of several years to decades^{12,18}. Therefore, long-distance seed dispersal is required; however, we need a better mechanistic understanding of these less frequent—yet highly relevant—seed-dispersal events¹⁹. Migratory birds have recently been identified as possible suppliers of these dispersal events^{3,4}, because these migrants can transport viable seeds over tens or even hundreds of kilometres in short time periods^{3,20,21}. The most notable evidence comes from the Canary Islands in the Atlantic Ocean, where about 1.2% of birds caught in migration by Eleonora's falcons (*Falco eleonorae*) were found to carry seeds in their guts from the mainland, over 170 km away²⁰. Further evidence comes from island colonization by fleshy-fruited plants^{22,23}, mechanistic models parameterized with empirical data of migratory movements and gut retention times of ingested seeds²⁴, and large-scale patterns of

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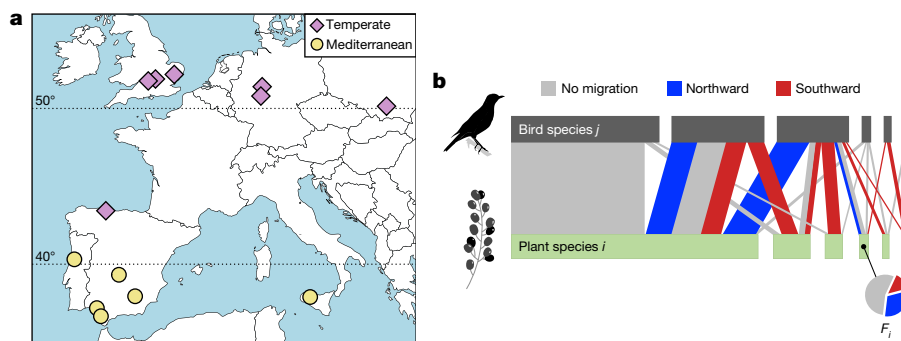


Fig. 1 | Location of the 13 European seed-dispersal networks we studied, and network with bird–plant interactions in relation to bird migration. a, Study sites in Portugal, Spain, UK, Germany, Italy and Poland. Symbols denote the biome of the locations (Mediterranean or temperate). **b**, Hypothetical seed-dispersal network illustrating how the weight of each pairwise interaction ij can be partitioned in relation to the migratory state of the bird using the

plant genetic structure along migratory routes²⁵ (Supplementary Discussion 1). Although such dispersal events seem rare, they are numerically compensated for by the fact that billions of birds migrate every year worldwide through seasonal and directional displacements that are highly predictable in space and time²⁶.

In the Northern Hemisphere, birds typically migrate towards the Equator in autumn (postnuptial migration) and towards the North Pole in spring (prenuptial migration)²⁶. Thus, plants could be consistently dispersed towards warmer or cooler latitudes depending on whether their fruiting period overlaps with southward or northward bird migrations, respectively (Extended Data Fig. 1). The relationship between migration directionality, plant phenology and dispersal potential towards cooler latitudes has, to our knowledge, been overlooked to date, despite this being crucial to predict the ability of plants to track climate change^{5,6,11}. In this Article, we provide an assessment of the potential of European plant communities to keep pace with climate change through long-distance seed dispersal towards cooler latitudes. We combined data on fruiting phenology and bird migration with information on pairwise interactions between frugivorous birds and fleshy-fruited plants from 13 woodland communities distributed across Europe (Fig. 1a). We focused on fleshy-fruited plants because many of their seed dispersers are migratory birds that far outnumber (both numerically and functionally) resident frugivores in European forests and woodlands^{27,28}. Moreover, fleshy-fruited plants are an important component of woody floras that account for a mean of 35% of species in temperate forests and 44% in Mediterranean woodlands²⁹.

We used data on seed-dispersal networks (that is, local communities of interacting bird and plant species) with links that describe the presence and intensity of pairwise interactions³⁰; in this case, the quantity of seeds of each plant species that is dispersed by each bird species (Fig. 1b). Importantly for the purpose of this study, all networks were sampled all year-round, covering the entire fruiting phenology of all plant species and the entire migration periods of all migratory birds (Extended Data Table 1). The 13 study networks were distributed across the Mediterranean ($n = 6$) and temperate ($n = 7$) biomes of Europe (Fig. 1a), and included a total of 949 interactions (range = 24–204 per network) between 46 bird species (range = 8–21) and 81 plant species (range = 8–29) (Extended Data Tables 1, 2). Most plant species were woody (89%), and the remainder were herbs (Extended Data Table 2).

In each network, we partitioned each bird–plant interaction according to the migratory state of the bird: migrating southwards, migrating northwards and non-migrating (Fig. 1b). To do so, we collated and combined information on the fruiting phenology of the plants and the phenology of bird migrations (Extended Data Fig. 1) (Methods). Then, for each plant species i , we calculated the fraction of its total interaction weight (F_i) that corresponds to interactions with each of the migratory states of the birds

phenological overlaps between the seed-dispersal period of plant species and periods of no migration, northward migration and southward migration of bird species (Extended Data Fig. 1). F_i are the interaction frequency values out of the total interaction weight of each plant species i that was with birds migrating southwards or northwards or with non-migrating birds.

(Fig. 1b, Extended Data Fig. 2). These calculations considered whether bird populations were full migrants or partial migrants, in which a fraction of the population migrates and the rest stays as residents²⁶ (Methods).

Our analysis tested whether the proportion of plant species interacting with migrating birds (prevalence), the frequency of such interactions and the number of bird species dispersing each plant species were significantly associated with the migration direction (southward or northward) (Methods). We used generalized linear mixed models to account for the non-normal error distributions and repeated measures per network and plant species. We also tested for differences between Mediterranean and temperate biomes in potential for plants to be dispersed towards cooler latitudes, because these biomes differ in fruiting seasonality: Mediterranean woodlands are characterized by longer fruiting seasons²⁹, which increase the probability of phenological overlap with the northward spring migration. We found that the majority (86%) of plant species across European communities are dispersed by birds migrating southwards, whereas only about one third (35%) are dispersed by northward-migrating birds (direction: $P < 0.001$) (Fig. 2a, Extended Data Table 3). This trend was consistent across biomes, but was less pronounced in Mediterranean (80% and 42% for southward and northward migrations, respectively) than in temperate communities (89% and 29% for southward and northward migrations, respectively) (interaction of biome \times direction, $P = 0.008$) (Fig. 2a, Extended Data Table 3). The sums of these percentages are greater than 100% because interactions with southward- and northward-migrating birds are not mutually exclusive (Fig. 1b), and many plant species are dispersed during both migrations (Extended Data Fig. 2, Supplementary Methods). The interaction frequency between plants and migrating birds was also much higher during the southward (36%) than during the northward (11%) migration (direction, $P < 0.001$) (Fig. 2b, Extended Data Table 3). This trend was also consistent across biomes, but was more pronounced in Mediterranean communities (40% and 9% for southward and northward migrations, respectively) than in temperate communities (32% and 13% for southward and northward migrations, respectively) (interaction of direction \times biome, $P = 0.011$) (Fig. 2b). Finally, plants were dispersed by more bird species migrating southwards than northwards (estimated mean = 2.9 and 2.3 species per plant, respectively; direction, $P = 0.017$), a small but consistent difference across biomes (Fig. 2c, Extended Data Table 3). Importantly, these results were not an artefact of analysing networks that were sampled with different methods (Supplementary Discussion 2). Our findings are congruent with general patterns in fruiting seasonality and bird migrations, as the fruiting peak in temperate and Mediterranean plant communities occurs between late summer and early winter²⁹, when migratory birds move southwards²⁶.

We further tested whether closely related plant species tend to have similar seed-dispersal interactions with birds migrating southwards or

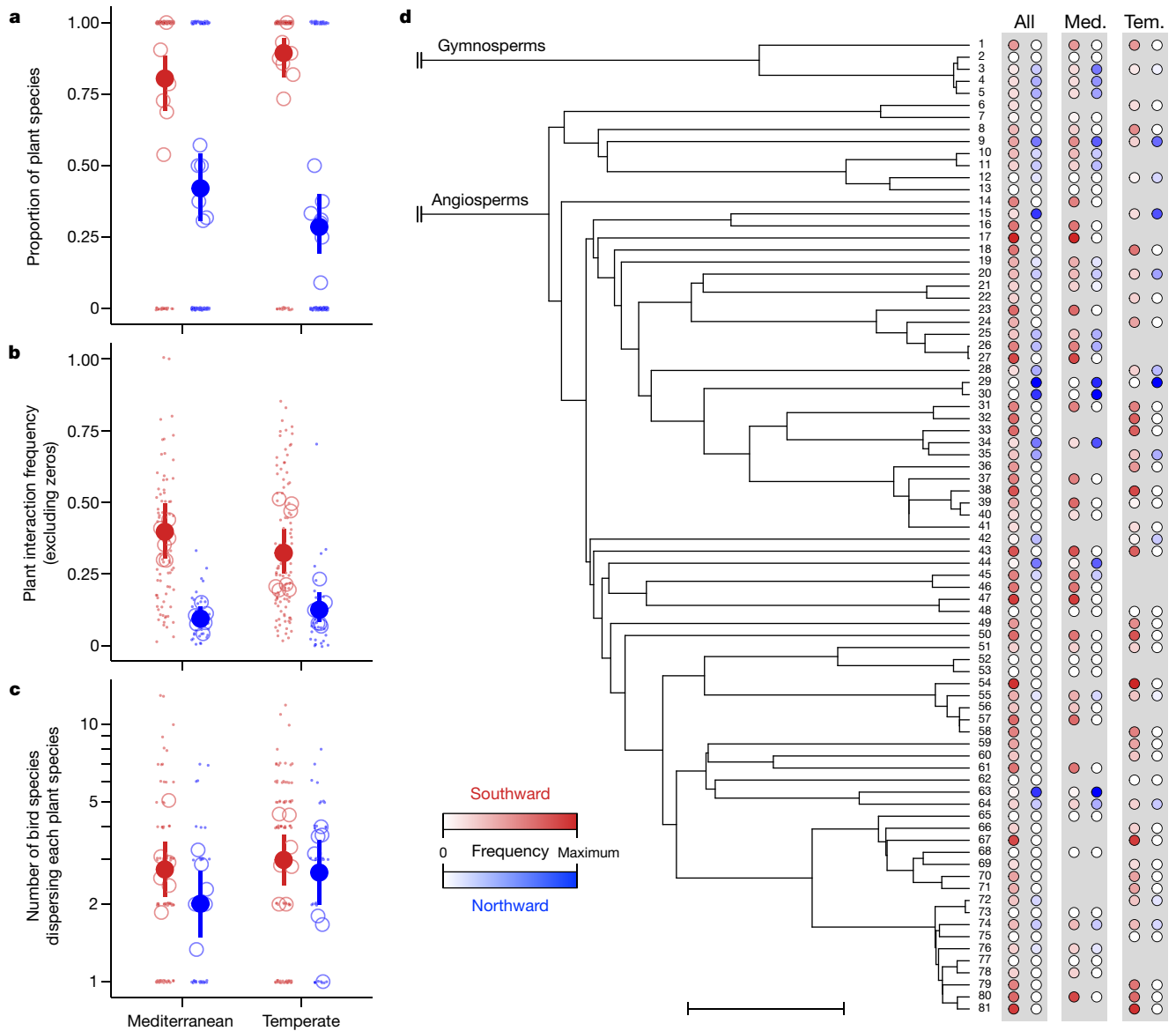


Fig. 2 | Seed-dispersal interactions of plants with migratory birds in relation to southward and northward migration and Mediterranean or temperate biome. a–c, Large dots and bars denote means \pm 95% confidence intervals estimated by generalized linear mixed models predicting the proportion of plant species interacting with birds during migration ($n = 434$ observations from 13 networks across plant species and directions) (**a**), the frequency of interactions with migrating birds when these occurred (zeros excluded) ($n = 260$ observations), out of the total seed-dispersal interactions (**b**), and the number of bird species dispersing each plant species ($n = 260$ observations) (**c**). Circles denote mean values for each seed-dispersal network;

tiny dots denote plant-level data. **d**, Dated phylogeny of the fleshy-fruited plants in the studied networks, with the root at 325 million years ago (shown in Extended Data Fig. 3). Scale bar, 50 million years. Numbers at the tips indicate species' codes (see species names in Extended Data Fig. 3). Coloured circles at the right of the tips indicate species-level means in interaction frequency (F_i) with birds migrating southwards or northwards (red and blue colour gradients, respectively) calculated across all networks (all) and, separately, for Mediterranean (Med.) and temperate (tem.) networks (maximum frequency means in all, Mediterranean and temperate, respectively: southward = 0.80, 0.80 and 0.76; northward = 0.30, 0.23 and 0.34).

northwards. The rationale is that the fruiting period of plants shows a phylogenetic signal³¹ and can thus be related to the phenological overlap with seasonal migrations (Extended Data Fig. 1b). We calculated the mean interaction frequency with birds migrating in either direction at the plant-species level, both across all networks ($n = 81$ species) and separately across Mediterranean ($n = 53$ species) and temperate ($n = 45$ species) networks (Fig. 2d). We found no phylogenetic signal for interaction frequency with southward migrants (all Pagel's $\lambda \ll 0.01$, $P = 1.0$). Conversely, we detected a strong phylogenetic signal for interaction frequency with northward migrants, both when considering all networks ($\lambda = 0.944$, $P = 1.2 \times 10^{-6}$) and when considering plants from Mediterranean ($\lambda = 0.895$, $P = 0.025$) or temperate ($\lambda = 0.999$, $P = 0.001$)

networks separately. We also detected significant phylogenetic signal for phenological variables describing the fruiting period of the study plants (specifically, the start and end dates, and the period length) (Extended Data Fig. 3). Accordingly, plant lineages that are frequently dispersed by northward-migrating birds are characterized by long fruiting periods (for example, *Juniperus* spp.; labels 3–5 in Fig. 2d) or late fruiting periods that extend until the spring of the next calendar year (for example, *Hedera* spp.; labels 29–30 in Fig. 2d). By contrast, plant lineages that are not dispersed by northward migrants are characterized by short fruiting periods between summer and early winter (for example, *Arum* spp. and *Prunus* spp.; labels 6–7 and 65–71 in Fig. 2d, respectively). Our results indicate that the potential of plants to track

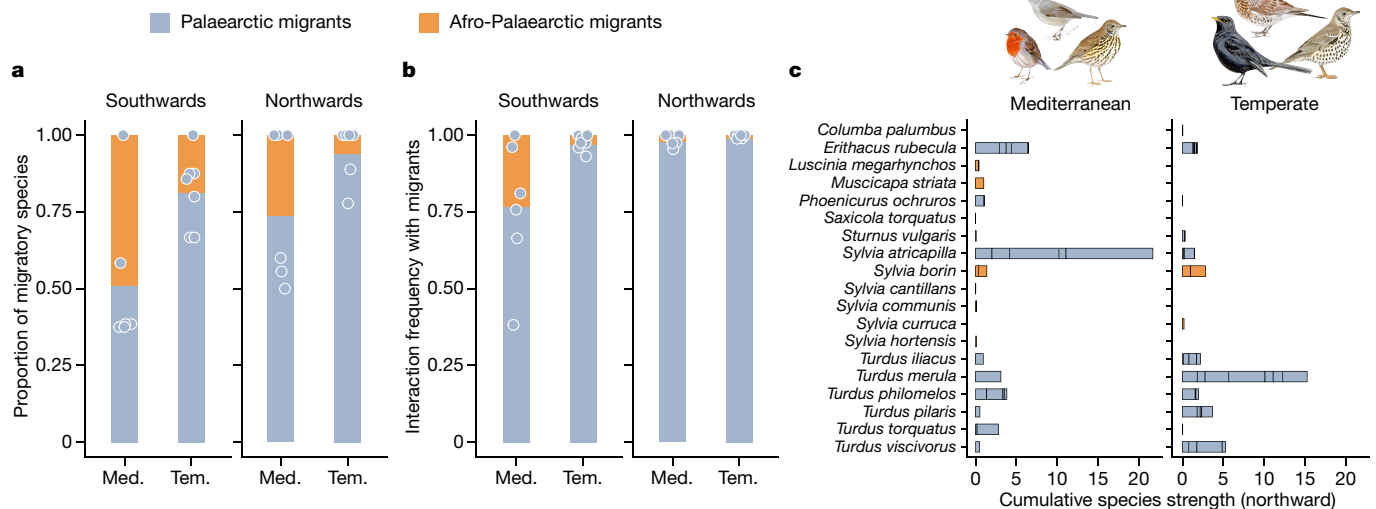


Fig. 3 | Relevance of Palaearctic and Afro-Palaearctic migratory birds dispersing seeds during their southward and northward migration in Mediterranean and temperate communities. **a**, Mean proportion of Palaearctic (grey) and Afro-Palaearctic (orange) species (wintering ranges in Europe and Africa north of the Sahara, and in sub-Saharan Africa, respectively) dispersing seeds while migrating, in relation to migration direction (southwards (left) or northwards (right)) and biome (Mediterranean or temperate). **b**, Mean relative contributions of Palaearctic and Afro-Palaearctic migrants to network-level interaction weight with migratory birds during their southward (left) and northward (right) migrations in Mediterranean and temperate communities. Circles in **a**, **b** denote network-level observations for

Palaearctic migrants. **c**, Relevance of bird species for seed dispersal towards cooler latitudes across Mediterranean (left) and temperate (right) fleshy-fruited plant communities, measured as the cumulative strength of bird species in subnetworks of interactions between plants and birds migrating northwards (blue links in Fig. 1b). Species strength quantifies the relevance of a bird species across the whole plant community; high cumulative values are found in bird species with high strength values in several subnetworks within each biome. Bird drawings represent the three most relevant species in each biome (from left to right, *E. rubecula*, *S. atricapilla*, *T. philomelos*, *T. merula*, *T. pilaris* and *T. viscivorus*). Illustrations by Juan M. Varela.

climate change is clustered in particular lineages, which suggests that the novel communities that may emerge in northern latitudes in the long term will incorporate non-random subsets of the evolutionary tree of southern floras. This phylogenetic filtering might have unanticipated consequences for ecosystem functions³² and community composition at higher trophic levels^{33,34}.

European migratory birds can be classified into two distinct groups according to their migratory strategy: Palaearctic migrants winter in southern Europe and northern Africa, whereas Afro-Palaearctic migrants winter in sub-Saharan Africa²⁶. In addition, both groups differ in their population trends; in contrast to Palaearctic migrants, Afro-Palaearctic migrants are experiencing major population declines across Europe for reasons that remain poorly understood^{35,36}. More bird species per community dispersed seeds during the southward than during the northward migration (mean = 9.5 and 5.9, respectively), a difference that was consistent across biomes (Extended Data Fig. 4). However, most of these species were Palaearctic migrants (Fig. 3a), particularly during the northward migration (mean = 87%; direction, $P = 0.005$) and in temperate communities (mean = 89%; biome, $P = 0.003$) (Extended Data Table 4). Furthermore, we found that both types of migrant differed in their functional relevance as seed dispersers, measured in terms of their interaction frequency (the proportion of the total interaction weight per network with all birds in migration (Methods)). Palaearctic migrants accounted for almost all interaction frequency (Fig. 3b), particularly during the northward migration (mean = 98%; direction, $P < 0.001$) (Extended Data Table 4). Palaearctic migrants were also significantly more relevant in temperate (mean = 98%) than in Mediterranean communities (mean = 87%; biome, $P < 0.001$), in which Afro-Palaearctic birds had a more important role—particularly when migrating southwards (interaction of direction \times biome, $P = 0.008$) (Extended Data Table 4). These results reveal that almost all seed-dispersal services towards cooler latitudes in Mediterranean (98%) and temperate (99%) communities across Europe are provided by a handful (about 5) of Palaearctic migrant species per locality. This stronger role of Palaearctic migrants can be explained by their occurrence in Europe

during the winter (when invertebrates are scant and fruits abundant) and their earlier spring migration northwards³⁷.

Finally, we assessed the identity and importance of different bird species contributing to seed dispersal during their northward migration. For this, we obtained bird species strength, which measures the sum of plant dependencies (relative interaction frequencies) on each bird species and therefore quantifies the relevance of a bird species for community-wide seed dispersal towards cooler latitudes (Methods). To do so, we used subnetworks of the original networks that included only interactions during northward migrations (blue links in Fig. 1b). Then, we obtained the cumulative species strength per biome as the sum of species strengths across Mediterranean and temperate subnetworks (Fig. 3c). Results showed that a few Palaearctic migrants are disproportionately important during their northward migration. Across Mediterranean communities, the blackcap (*Sylvia atricapilla*) was by far the most important species, followed by the European robin (*Erithacus rubecula*) and the song thrush (*Turdus philomelos*) (Fig. 3c); these three species accounted for 73% of cumulative strength (blackcap, 49%; robin, 15%; and song thrush, 9%). Across temperate communities, the blackbird (*Turdus merula*) was prominently the most relevant species, followed by the mistle thrush (*Turdus viscivorus*) and the fieldfare (*Turdus pilaris*) (Fig. 3c); these thrushes accounted for 69% of cumulative strength (blackbird, 44%; mistle thrush, 15%; and fieldfare, 10%). Thus, bird species did not have similar relevance in both biomes (Extended Data Fig. 5). Notably, the key bird species for plant dispersal towards cooler latitudes are—in general—common and abundant birds, highlighting their importance for the functioning and dynamics of ecological communities³⁸. However, their functional role in providing long-distance seed dispersal towards cooler latitudes could be at risk because bird migrations are already being disrupted by climate change^{39,40}. Moreover, some of these species are severely hunted (both legally and illegally), particularly in the Mediterranean region. In fact, *S. atricapilla* and *T. philomelos* are in the top five of the most

illegally hunted birds in the Mediterranean Basin (estimated over 1 million individuals of each species killed per year⁴¹).

Available evidence shows that seed-dispersal distances by resident animals are typically insufficient for plants to track current climate change; however, it suggests that migratory birds can supply the long-distance dispersal services required (Supplementary Discussion 1). Given that our approach is based on mainstream migratory movements, our results provide a general template of the potential for directional, long-distance seed dispersal. A further step to accurately estimate dispersal distances and directionality requires detailed movement data of migratory birds, which are necessary for the development of mechanistic seed-dispersal models²⁴. Such data are expected to come during this decade, as we are witnessing a revolution in next-generation GPS tags that will enable the tracking of small frugivorous birds with unprecedented spatiotemporal resolution⁴².

The Earth is warming rapidly and is expected to continue to do so in the near future^{2,8}. Our study reveals that only about a third of fleshy-fruited plant species across European biomes will benefit from directed long-distance dispersal by migratory birds towards northern latitudes to track favourable conditions. These few 'winners' are phylogenetically clustered in plant lineages characterized by either long or late fruiting periods, and are mostly dispersed by a few common bird species with a relevance that is biome-specific. Our findings are expected to be broadly generalizable to other regions in the Northern Hemisphere (North America and Asia), where the fruiting period of most fleshy-fruited plants occurs in autumn^{43,44}, when most birds move southwards, and where bird migration is a much more obvious phenomenon than in the Southern Hemisphere²⁶. The extent to which our findings are generalizable to other plant–bird systems, such as aquatic plants dispersed internally or externally by waterbirds^{21,45,46}, deserves further research. Understanding large-scale dispersal is necessary to develop conservation practices aimed at halting and mitigating biodiversity loss driven by climate change¹. Our study suggests that migratory birds are only helping a phylogenetically clustered minority of plant species to disperse towards cooler latitudes, while they are dispersing most species towards increasingly drier and hotter regions. This divergent dispersal is expected to strongly influence the formation of novel communities in the future. Finally, our results provide a baseline to assess whether climate-driven phenological shifts will exacerbate or improve this situation.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41586-021-03665-2>.

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Methods

No statistical methods were used to predetermine sample size. Randomization and blinding do not apply to our study, as we did not conduct experiments.

Study seed-dispersal networks

We refer to interacting communities of frugivorous birds that disperse the seeds of fruiting plants as networks, which are interaction matrices in which each row i represents a plant species and each column j represents a bird species. Elements in the matrices (w_{ij}) denote whether pairwise plant–bird interactions were observed ($w_{ij} > 0$) or not ($w_{ij} = 0$) and—if so—their value account for interaction weight.

Our study includes 13 seed-dispersal networks evenly distributed between the Mediterranean ($n = 6$) and the temperate ($n = 7$) biomes of Europe^{47,48} (Fig. 1a and Extended Data Table 1). All these networks are quantitative (interactions are weighted) and were sampled in natural forests and woodlands, most of them in lowland habitats (<600 metres above sea level) in which agricultural fields surround the remnant vegetation. The single exception is the ‘Nava Correhuelas’ network, which is located on a well-preserved Mediterranean mountain at an elevation of 1,600 metres. Eight of the 13 networks include new data obtained by the authors, and five were compiled from previous studies^{27,49–51}. Seven of the eight new networks were sampled within the EU project ‘MobileLinks’, through field sampling of bird-dispersed seeds and subsequent disperser identification by means of DNA-barcoding analysis⁵² (as described in ‘Methods for the new network data’). The other six networks were obtained either through focal plant observations of birds feeding on fruits or through dietary analysis of birds captured in mist nets (Extended Data Table 1). In networks obtained through focal observations, we focused exclusively on pairwise interactions in which the bird behaves as a legitimate seed disperser (swallowing the fruit and defecating or regurgitating viable seeds), discarding pulp-pecking and seed-predation interactions⁵³.

Importantly, the study networks were sampled all year-round (for 1–6 years; mean = 2 years), a prerequisite to cover the entire fruiting periods of all local fleshy-fruited species, as well as the pre-nuptial and post-nuptial migration periods of all migratory birds; the single exception was the network from ref.²⁷, which was sampled during nine months (August to early May), covering most of the year and both migrations (Extended Data Table 1). We thus avoided using other European networks^{54–56} that were sampled during short temporal periods (5–6 months). The study networks included a total of 949 interactions between frugivorous birds and fleshy-fruited plants (median = 52 per network, range = 24–204). Some interactions occurred in more than one network, resulting in 563 unique pairwise interactions between 46 bird species (median = 14 per network, range = 8–21) and 81 plant species (median = 15 per network, range = 8–29). The number of bird species, plant species or interactions did not differ significantly between Mediterranean and temperate networks ($P \geq 0.20$ in one-way analyses of variance (ANOVAs) $\log_{10}[n]$ -biome). Bird and plant species included 16 and 28 families, respectively (Extended Data Table 2). The plants included trees and shrubs (79% of species), herbs (11%) and woody vines (10%); thus, most plant species (89%) were woody.

Because we were interested in the seed-dispersal function, we expressed the interaction weights (w_{ij}) of all networks as the number of seeds of each plant species i (or the seed-rain density as seeds per m²) dispersed by each bird species j . These weights were directly obtained in networks that sampled bird-dispersed seeds, either in seed traps for subsequent DNA-barcoding analysis or in droppings from birds captured in mist nets (as described in ‘Methods for the new network data’). Yet, in networks based on feeding observations, interaction weights were originally expressed as number of bird visits to focal plants⁵³. We then converted number of visits into number of seeds through the following two steps. First, we converted visits into fruits consumed using

the parameters of a linear mixed model ($R^2_{\text{GLMM}(m)} = 0.924$) fitted to data from two European networks^{56,57} for which the number of both visits and fruits consumed were recorded for each pairwise interaction (Supplementary Methods). In a second step, we converted fruits consumed into seeds dispersed by multiplying the former by the average number of seeds per fruit of each plant species, which was obtained from the literature^{50,58–60} and from data generated by the authors. In cases in which the product did not result in an integer, values were rounded to the nearest integer.

All networks were combined into a single data table for subsequent incorporation of data on seed-dispersal phenology and bird-migration periods (as described in ‘Seed-dispersal phenology’ and ‘Migrant types and phenology of bird migrations’), with columns for network identity, network biome, network country and bioclimatic zone, plant and bird species, and interaction weight. Hereafter, we refer to ‘seed-dispersal period’ rather than to ‘fruiting period’ because a part of our phenological data was based on the presence of seeds dropped by birds in seed traps or during mist netting (as described in ‘Methods for the new network data’).

Methods for the new network data

Authorship of the eight unpublished networks is shown in the ‘Author contributions’ section.

MobileLinks networks. Community-wide seed dispersal by frugivorous birds was sampled within the EU project ‘MobileLinks’ (H2020-MSCA-IF-2014-656572) in seven European landscapes (plots of 1–4 km²) located in Spain, UK, Germany, Italy and Poland (Extended Data Table 1). Six of these plots were sampled for one year (2016–2017) and one plot for two complete years (2013–2015). In all plots, seed traps were placed beneath tree and shrub canopies (natural perches), and under electricity pylons (anthropogenic perches) used by birds, to quantify the magnitude of bird-mediated seed rain in the landscape⁶¹. Seed traps were 0.22-m² plastic trays covered with wire mesh to prevent postdispersal seed removal. Between 40 and 77 seed traps (mean = 46.3) were monitored in each study plot. Sampling surveys, in which the number of bird-dispersed seeds per trap was recorded, were conducted fortnightly; seeds were visually identified (as described below in this section). The route used to survey the seed traps was also used as a 1-m wide single fixed belt transect (range 2,630–9,110 m length, mean 4,410 m) to search for bird-dispersed seeds and quantify seed rain in canopy-free open interspaces, where bird-mediated seed rain is less likely⁶¹. Individual seeds or droppings with seeds were sampled for DNA barcoding analysis into 1.5- or 2.0-ml sterile tubes that were labelled and stored in a freezer at –20 °C until DNA extraction. Because DNA barcoding identification generally fails (PCR failure) in 5–10% of samples^{28,52,61}, some bird-dispersed seeds visually detected outside the transects were also sampled for DNA barcoding analysis aiming at increasing sample sizes, particularly for locally rare plant species. Conversely, only a subsample of the seeds was generally sampled when seed traps received many seeds of particular plant species; for example, 40–50% of the hyper-abundant *Pistacia lentiscus* seeds in Garrapilos during its fruiting peak²⁸.

We used DNA barcoding analysis (mitochondrial COI (cytochrome c oxidase subunit I)) to identify the bird species responsible for the seed-dispersal events, as DNA of animal origin can be extracted from the surface of defecated or regurgitated seeds^{28,52,61}. Detailed laboratory protocols for DNA extraction, PCR, sequencing and species identification can be found in Supplementary Methods. Resulting sequences were identified at the species level based on best sequence matches in the ‘BARCODE OF LIFE DATA’ identification system (BOLD⁶²) (www.boldsystems.org), typically at a 98–100% similarity (Supplementary Fig. 1). We successfully identified the disperser species of 2,991 samples (that is, 2,991 sequences; 123–1,753 per network) including 3,014 interaction events between a bird–plant species pair, and containing

4,812 seeds (144–2,193 per network); overall 3,234 samples containing 5,181 seeds were analysed, with an identification success of 92.5% (PCR failure occurred in 7.5% of samples). All barcoding sequences obtained in the present study are publicly available in the data file 'MOBILELINKS_DNA_barcoding_data.csv' deposited at the DRYAD repository (<https://doi.org/10.5061/dryad.15dv41nx3>).

After DNA extraction for bird DNA barcoding, we visually identified seed species according to their morphology. To do so, we compared the seeds against a personal reference collection (owned by J.P.G.-V.) and pictures from a guide of seeds of European fleshy-fruited species that includes plants from the Mediterranean and temperate biomes⁵⁸. The exception were 11 samples for which initial identification was not possible and for which we conducted DNA-barcoding analysis using chloroplast *MaturaseK* gene (*matK*)⁶³ (detailed laboratory protocols are provided in Supplementary Methods). Seed species from these 11 samples were identified through the following three steps: (1) we obtained a short list of species from the best sequence matches in BLAST⁶⁴; (2) we used such short list to identify candidate fleshy-fruited plant species that were present around the study sites; and (3) we used the final reduced list of candidate plant species to identify seed species visually according to seed morphology. Thus, this DNA-barcoding analysis served us to short list and guide visual identification (further details are provided in Supplementary Methods). All seed samples are stored by J.P.G.-V. at the Laboratory of Botany in the University of Cádiz, and plant sequences are publicly available in the data file 'MOBILELINKS_DNA_barcoding_data.csv' deposited at the DRYAD repository (<https://doi.org/10.5061/dryad.15dv41nx3>).

We used DNA barcoding identifications to calculate the relative contribution (f_{ijk}) of each bird species j to the seed-rain density of plant species i beneath perch type k as $f_{ijk} = n_{\text{DNA-}ijk} / n_{\text{DNA-}ik}$, in which n_{DNA} is the number of DNA-barcoded seeds. We then estimated the seed rain of each plant species dispersed by each bird species beneath different perch types as $sr_{ijk} = sr_{ik} \times f_{ijk}$, in which sr_{ik} is average seed rain (seeds per m²) of plant species i measured in seed traps located beneath perch type k (similar to procedures in refs.^{28,65}). Finally, we calculated the site-level (network-level) seed-rain density of each plant species dispersed by each bird species (sr_{ij}) as the weighted mean of sr_{ijk} values across perch types, using the number of seed traps per perch type as weighting factor. The result (sr_{ij}) was the weight (w_{ij}) of pairwise interactions in these networks, expressed as seeds per m².

Vale Soeiro network. Community-wide seed dispersal by frugivorous birds was sampled in a plot of natural woodland of about 0.5 km² located in central Portugal. A total of 168 m of mist nets of different lengths (nine mist nets of 15 m, two of 12 m and one of 9 m) were operated fortnightly during 5 h after dawn for 6 years (2012–2018). Nets were visited every 30 min and captured birds were individually placed in ringing bags for up to 30 min until they ejected droppings. Out of a total of 4,462 bird captures, 1,330 produced droppings with seeds ($n = 3,398$ seeds). Defecated or regurgitated seeds were later extracted and identified by comparison with a reference collection. A quantitative seed-dispersal network was built in which interaction weights (w_{ij}) represented the total number of seeds of each plant species i dispersed by each bird species j .

Seed-dispersal phenology

Seed-dispersal phenology is the period in which plants bear ripe fruits and disperse their seeds. We obtained bioclimate-level data on seed-dispersal phenology of the plant species in the study networks; the bioclimatic zone of the study networks is shown in Extended Data Table 1. We targeted on bioclimate-level phenology because the dispersal period of a given plant species may differ between bioclimatic zones⁶⁶. We used distinct data sources: published studies^{27,50,55,59,67–77}, from which information was extracted from figures, tables and text; and our own data associated with the new eight networks that we

obtained through fortnightly sampling surveys. Published studies included data on entire fleshy-fruited plant communities (for example, ref.⁶⁷), in some cases associated to the published networks we compiled (for example, refs.^{50,59,72}), as well as data on specific taxa (for example, refs.^{68,70}). In some cases, we also used personal observations for the phenology of particular plant species at specific bioclimatic zones. From each data source, we obtained the 'start' and the 'end' of the seed-dispersal period ($D_{\text{start}} - D_{\text{end}}$) of each plant species. We used a monthly scale (0–12) in which exact values represent the transition between months (for example, 0 = end of December–beginning of January; 1 = end of January–beginning of February; and so on) and half values represent the midpoint within months (for example, 1.5 = mid-February). For instance, a fruiting period from mid-June to late September was expressed as $D_{\text{start}} = 5.5$ and $D_{\text{end}} = 9$. We added 12 to D_{end} whenever it belonged to the next calendar year; hence, a dispersal period from mid-November to late March was expressed as $D_{\text{start}} = 10.5$ and $D_{\text{end}} = 15$ (3 + 12) (Extended Data Fig. 1). Most phenological data were obtained at a 0.5-month accuracy (about 2 weeks), although in a few data sources the information was found at a 0.25-month accuracy. We obtained data for 143 of the 150 unique 'plant species–bioclimate' combinations (95%) from 288 original data entries as, in many cases, we obtained data from several sources or years for the same plant species at a particular bioclimate (Extended Data Fig. 6). When the same data source included information from different fruiting seasons (for example, refs.^{72,76}), we averaged D_{start} and D_{end} values of each plant species across seasons. Then, we obtained unique D_{start} and D_{end} values for each plant species–bioclimate combination by averaging across data sources. With this procedure, we aimed at conservatively obtaining the most representative and generalizable seed-dispersal period of each plant species within each bioclimate. We also obtained the minimum D_{start} and the maximum D_{end} recorded (that is, the longest fruiting period per plant species–bioclimate combination) to perform a complementary analysis using a less conservative approach (Supplementary Discussion 2). For the seven remaining plant species–bioclimate combinations lacking specific information, we used data from the closest bioclimate. For example, we used phenological data of *Rhamnus lycioides* from thermo-Mediterranean bioclimate for one meso-Mediterranean network. Phenological data were finally incorporated into the network data according to plant species and network bioclimate.

Migrant types and phenology of bird migrations

We used published information^{59,78–85} and our own data (periodic bird censuses and mist netting captures) to classify the bird species in each study network as 'resident' (non-migrant), 'Palearctic migrant' (birds that breed in Europe and winter in southern Europe and Africa north of the Sahara) and 'Afro-Palearctic migrant' (birds that breed in Europe and winter in sub-Saharan Africa)²⁶. Palearctic migrants are often partial migrants (that is, only a fraction of their populations migrates while the other fraction behaves as resident)²⁶. For this reason, we characterized the proportion of migrants (P_{migrants}) in the bird populations of the study networks by means of a semiquantitative variable: 0, non-migrant population; 0.1, only a minor fraction migrates; 0.25, a larger fraction migrates but non-migrants prevail; 0.5, roughly half of the population migrates; 0.75, migrants prevail; 0.9, only a minor fraction does not migrate; 1: the whole population migrates (for this variable, we also used published information^{79,80,82–91} and our own data). Hence, Palearctic migrants showed P_{migrants} values ranging from 0.1 to 1. We also classified fully migrant populations ($P_{\text{migrants}} = 1$) as 'wintering', 'summer-breeding' or 'transient' depending, respectively, on whether birds occur locally during the winter, the breeding season or short periods while migrating (stopover site)²⁶.

We obtained country-level phenological data for the pre-nuptial (northward) and post-nuptial (southward) migrations of the bird species in the study networks (countries in Extended Data Table 1). In this

case, we targeted on country-level phenology to capture geographical variation in the timing of migrations, as this information was obtained mainly from bird migration atlases of the study networks' countries: Spain and Portugal (Iberia)^{79,80}, Italy^{82,83}, UK⁸⁴, Germany⁸⁶ and Poland⁸⁷. We also obtained data from specific references from Poland⁸⁸⁻⁹¹ and websites from recognized ornithological organizations in the case of Spain (www.seo.org/listado-aves-2)⁸¹ and the UK (www.birdtrack.net)⁹². We gathered phenological data for the 119 unique 'bird species-country' combinations. From each data source, we obtained the 'start' and the 'end' of both the northward (N , prenuptial) and southward (S , postnuptial) migration periods ($N_{\text{start}}-N_{\text{end}}$ and $S_{\text{start}}-S_{\text{end}}$, respectively) from figures, tables and text. All phenological data were obtained at a 0.5-month accuracy (about 2 weeks). Again, we used a monthly scale in which exact values represent the transition between months and half values represent the midpoint within months (as described in 'Seed-dispersal phenology'). Only in 3 cases (2.5% of the 119 bird species-country combinations) for which we did not obtain some of the four migration dates at the country level ($N_{\text{start}}-N_{\text{end}}$ and $S_{\text{start}}-S_{\text{end}}$), we used migration phenology available for the Western Palearctic region⁸⁵ or at a continental coarse scale (www.eurobirdportal.org). Phenological data were finally incorporated into the network data according to bird species and network country.

Directional migration in seed-dispersal interactions

Phenological overlap during migrations. For each plant-bird interaction in each network, we calculated the phenological overlap between the seed-dispersal period of the plant and the northward and southward migration periods of the bird ($O_{\text{north-}ij}$ and $O_{\text{south-}ij}$, respectively; O units are months). We calculated these overlaps as the difference between the minimum 'end' and the maximum 'start' of both periods; $O_{\text{north-}ij} = \min(D_{\text{end-}i}, N_{\text{end-}j}) - \max(D_{\text{start-}i}, N_{\text{start-}j})$, and $O_{\text{south-}ij} = \min(D_{\text{end-}i}, S_{\text{end-}j}) - \max(D_{\text{start-}i}, S_{\text{start-}j})$. Before these calculations, we added 12 to the dates of the northward migration (spring) of bird species j whenever the seed-dispersal period of plant species i extended to the next calendar year (if $D_{\text{end-}i} > 12$). This solved, for instance, the fact that a period of northward migration $N_{\text{start-}j}-N_{\text{end-}j} = 2-4$ does not overlap mathematically with a seed dispersal period $D_{\text{start-}i}-D_{\text{end-}i} = 10-16$, despite there being a true phenological overlap ($N_{\text{start-}j}-N_{\text{end-}j} = 2-4 = 14-16$). Negative and 'NA' values obtained were converted into zeros (no overlap) as they represented, respectively, the lack of either phenological overlap or migration (resident birds). We provide a graphical representation for the overlap calculations of this section in Extended Data Fig. 1b and Supplementary Fig. 3.

Total phenological overlap. Apart from the phenological overlap during migrations, we also calculated the total phenological overlap ($O_{\text{total-}ij}$) as the whole period during which a bird species coincides locally with the seed-dispersal period of each plant species in the study networks. When bird populations were fully or partially resident ($P_{\text{migrants}} < 1$), the bird species occurs locally all year round and, thus, $O_{\text{total-}ij}$ was equal to the length of the seed-dispersal period ($O_{\text{total-}ij} = D_{\text{end-}i} - D_{\text{start-}i}$). When bird populations were transient (only occur locally during migration), $O_{\text{total-}ij}$ was equal to the sum of phenological overlap during northward and southward migrations ($O_{\text{total-}ij} = O_{\text{north-}ij} + O_{\text{south-}ij}$). In the case of wintering migrants, their occurrence in the local communities spans from their arrival at the beginning of the southward migration (S_{start}) to the end of their departure at the end of the northward migration (N_{end}); thus, for wintering migrants: $O_{\text{total-}ij} = \min(D_{\text{end-}i}, N_{\text{end-}j}) - \max(D_{\text{start-}i}, S_{\text{start-}j})$. In the case of summer migrants, their presence in local communities spans from their arrival at beginning of the northward migration (N_{start}) to their complete departure at the end of the southward migration (S_{end}); thus, for summer migrants: $O_{\text{total-}ij} = \min(D_{\text{end-}i}, S_{\text{end-}j}) - \max(D_{\text{start-}i}, N_{\text{start-}j})$. Whenever $D_{\text{end-}i}$ extended to the next calendar year ($D_{\text{end-}i} > 12$), we added 12 to the migration dates to calculate the actual $O_{\text{total-}ij}$ (Supplementary Fig. 3).

Frequency of seed-dispersal interactions during migrations. We then used the phenological overlaps during migration ($O_{\text{north-}ij}$ and $O_{\text{south-}ij}$) and the total phenological overlap ($O_{\text{total-}ij}$) to calculate, for each plant-bird interaction ij , the frequency of seed-dispersal interactions in which the bird is migrating northwards as $F_{\text{north-}ij} = P_{\text{migrants-}j} \times O_{\text{north-}ij} / O_{\text{total-}ij}$, and southwards as $F_{\text{south-}ij} = P_{\text{migrants-}j} \times O_{\text{south-}ij} / O_{\text{total-}ij}$. The calculation was the fraction of the total phenological overlap accounted for by each migration period and weighted by the proportion of migrants in the bird population ($P_{\text{migrants-}j}$). For instance, if $O_{\text{north-}ij} = 3$ and $O_{\text{total-}ij} = 6$, then $F_{\text{north-}ij} = 0.5$ if the whole bird population migrates ($P_{\text{migrants-}j} = 1$; $F_{\text{north-}ij} = 1 \times 3/6$), but $F_{\text{north-}ij} = 0.05$ if only a minor fraction of the bird population migrates ($P_{\text{migrants-}j} = 0.1$; $F_{\text{north-}ij} = 0.10 \times 3/6$). For fully resident populations, $O_{\text{north-}ij}$, $O_{\text{south-}ij}$ and $P_{\text{migrants-}j}$ equal 0, and thus $F_{\text{north-}ij}$ and $F_{\text{south-}ij}$ too. We calculated the frequency of seed-dispersal interactions in which the bird is non-migrating as $F_{\text{non-}ij} = 1 - (F_{\text{north-}ij} + F_{\text{south-}ij})$. Through this approach, we made the assumption that interaction frequency is uniformly distributed throughout $O_{\text{total-}ij}$. We consider it to be a conservative assumption because the magnitude of seed dispersal by frugivorous birds throughout the fruiting season can be roughly constant (our assumption), unimodal symmetric, unimodal skewed or even multimodal, depending on the plant species (for example, refs. 27,50,71) and the local context (for example, ref. 74).

Finally, for each plant species i in each study network, we calculated the frequency of seed-dispersal interactions during which the birds are migrating south ($F_{\text{south-}i}$), north ($F_{\text{north-}i}$) or are not migrating ($F_{\text{non-}i}$), as the weighted means of F_{ij} across j bird species. Weighting was done by the interaction weight w_{ij} of each pairwise interaction. Hence, F_i values represent the fraction of the total interaction weight of plant species dispersed by birds with distinct migratory states ($F_{\text{south-}i} + F_{\text{north-}i} + F_{\text{non-}i} = 1$) (Extended Data Fig. 2).

Statistical analyses

Four out of the 81 plant species (*Crataegus monogyna*, *Hedera hibernica*, *Rosa canina* and *Rubus fruticosus*) actually represented operational taxonomic units in some networks owing to the local occurrence of congeneric species with seeds that did not allow for unambiguous species-level identification (*Crataegus laevigata*, *Hedera maderensis*, other *Rosa* and other *Rubus* species, respectively). In these cases, we used the name of the most common species to match the species name across networks, which allowed us to use plant species as random factor in mixed models and match a unique tip label in the plant phylogeny.

All generalized linear mixed models (GLMMs) described below were fitted using the R package glmmTMB (v.0.2.3)⁹³ and the significance of fixed effects (P values of type II Wald χ^2 tests) was computed using the Anova function of the R package car (v.2.1-6)⁹⁴.

Seed-dispersal interactions. We fitted GLMMs to test whether the migration direction (southward or northward), the biome (Mediterranean or temperate) and the interaction between these two fixed factors were significantly associated with (1) the proportion of plant species (prevalence) interacting with birds during migration ($n = 434$ observations), (2) the frequency of seed-dispersal interactions with birds during migration (whenever these interactions occurred; non-zero F_i) out of the total interaction weight, and (3) the number of bird species dispersing each plant species during migration (whenever interactions during migration occurred; non-zero values); $n = 260$ observations in (2) and (3). Importantly, the prevalence and frequency of interactions with migrants were not interrelated in both migrations (Supplementary Methods). All models included network identity and plant species nested within network as random factors (random intercepts) to account for the repeated measures per network (different plant species) and per plant species within networks (same plant interacting with birds migrating southwards and northwards). Prevalence among plant species was modelled as a Bernoulli-distributed variable with

logit link function (1: $F_i > 0$; 0: $F_i = 0$). Frequency ($F_i > 0$) was modelled as a mixed-effects beta regression with logit link function in which the dispersion parameter ϕ of the beta distribution was allowed to vary in response to the interactive effects of direction and biome ($\Delta\text{AIC} = -25$ relative to a model with fixed ϕ)⁹⁵. For modelling purposes, we transformed F_i values as follows: $F'_i = (F_i(n-1) + 0.5)/n$, in which n is the total number of observations⁹⁵. This transformation compresses the closed interval ($0 \leq y \leq 1$) within the open interval ($0 < y < 1$) because the values modelled by beta distribution are defined on the latter⁹⁵ (range of $F_i > 0$: 0.0009–1; range of F'_i : 0.0020–0.9988); the estimated means and 95% confidence intervals reported in the Article (Fig. 2b) were previously back-transformed ($F_i = (F'_i n - 0.5)/(n - 1)$). The number of migrating bird species that dispersed each plant species was modelled as a Poisson-distributed variable with log link function.

Phylogenetic signal in plants. We tested for the presence of phylogenetic signal in the plant species means across networks regarding their interaction frequency (F_i values, including zeros) with birds migrating southwards and northwards, for all plant species ($n = 81$) and separately for species in Mediterranean and temperate networks ($n = 53$ and 45, respectively). We calculated plant species means across networks because many plant species participated in several networks (mean = 2.7, range = 1–12), either from the same or different biomes. For example, *Cornus sanguinea* participated in six temperate networks, *Myrtus communis* in four Mediterranean networks, and *C. monogyna* in 12 networks from both biomes. We extracted information about the phylogenetic relatedness of the plants present in the study networks from a dated phylogeny of seed plants (Spermatophyta)⁹⁶ with a backbone based on a previous publication⁹⁷. The tree was prepared by dropping tips other than the 81 plant species of interest using the R package *ape* (v.5.3)⁹⁸; the resulting tree contained one polytomy, which was resolved randomly using the function *multi2di*. Phylogenetic signal was assessed through Pagel's λ (ref.⁹⁹), a statistic that varies between 0 (phylogenetic independence) and 1 (species' traits covary in direct proportion to their shared evolutionary history under a Brownian motion model of quantitative trait evolution)¹⁰⁰. Intermediate values of λ indicate that traits have evolved according to a process in which the effect of phylogeny is weaker than in the Brownian model¹⁰⁰. Pagel's λ seems strongly robust to polytomies and suboptimal branch-length information¹⁰¹. Significant phylogenetic signal ($\lambda > 0$) is calculated through a likelihood ratio test comparing the likelihood of the model fitted to the data (observed λ) to that of a model in which λ was fixed to 0¹⁰⁰. These analyses were performed using the R package *phytools* (v.0.6-99)¹⁰². As a complementary analysis, we also tested for phylogenetic signal in seed-dispersal phenology as the frequency of interactions with migrant birds is ultimately related to fruiting phenology (Extended Data Fig. 3).

Migratory birds. We fitted GLMMs to test whether the migration direction, the biome and the interaction between these two fixed factors were significantly associated with the number of migratory bird species in the study networks dispersing plants during migration, and with the proportion of Palaeartic and Afro-Palaeartic species. The species richness was modelled as a Poisson-distributed variable with log link function and the proportion of Palaeartic species as a Bernoulli-distributed variable with logit link function (1: Palaeartic; 0: Afro-Palaeartic; the proportion of both migrant types are fully interdependent). Network identity was included as a random factor (random intercepts) to account for the repeated measures within networks ($n = 26$ observations in each case, that is, 'network–direction' combinations). We also assessed whether the relevance of Palaeartic and Afro-Palaeartic migrants varied between migrations and biomes. To do so, we calculated the frequency of interactions with Palaeartic and Afro-Palaeartic birds on migration in each network out of the total interaction weight with all migrant birds during the southward

and northward migrations (for example, $f_{\text{Palaeartic}} = W_{\text{Palaeartic}}/W_{\text{allmigrants}}$, in which $W_{\text{allmigrants}}$ is the total interaction weight with all migrating birds per network, thus, $W_{\text{allmigrants}} = W_{\text{Palaeartic}} + W_{\text{Afro-Palaeartic}}$). For this analysis, we used only data from Palaeartic migrants ($f_{\text{Palaeartic}}$, $n = 26$ observations, that is, network–direction combinations) because frequencies from both migrant types are fully interdependent ($f_{\text{Palaeartic}} + f_{\text{Afro-Palaeartic}} = 1$). We fitted a GLMM to test whether the migration direction, the biome and their interaction were significantly associated with the interaction frequency with Palaeartic migrants. This model was as a mixed-effects beta regression with logit link function⁹⁵, in which the dispersion parameter ϕ of the beta distribution was allowed to vary in response to the additive effects of direction and biome ($\Delta\text{AIC} = -29$ relative to a model with fixed ϕ). For modelling purposes, we transformed values for beta regression as explained in 'Seed-dispersal interactions' (range of $f_{\text{Palaeartic}}$: 0.3818–1.0; range of $f'_{\text{Palaeartic}}$: 0.3841–0.9904); the estimated means reported in the article (Fig. 3b) were also back-transformed as explained in 'Seed-dispersal interactions'. Network identity was included as random factor (random intercepts) to account for the repeated measures within networks.

Finally, we used 'species strength', a species-level network metric¹⁰³, to identify the most relevant bird species dispersing seeds during each migration. Species strength is the sum of plant dependencies (relative interaction frequencies) on each bird species, therefore, it quantifies the relevance of a bird species across all the fleshy-fruited plant community¹⁰⁴. We calculated species strength of migratory birds ($n = 24$ species) using the R package *bipartite* (v.2.13)¹⁰³ in subnetworks of the original networks that only included seed-dispersal interactions either during southward or northward migration (subnetworks of red or blue links in Fig. 1b, respectively), in which strength quantifies the relevance of a bird species as a seed disperser during each migration. We then obtained the cumulative species strength (sum across sub-networks) per direction and biome combinations (Mediterranean–southward, Mediterranean–northward, temperate–southward and temperate–northward). This way, very high cumulative values can only be found in migratory bird species with high strength values in several networks per biome. We used nonparametric Kendall's rank correlations to test whether, in each biome, the cumulative species strength across southward and northward subnetworks were correlated, which would indicate that bird species generally display a proportional role in both migrations (Extended Data Fig. 5). Besides, we used Kendall's rank correlations to test whether, for each migration, the cumulative species strength across Mediterranean and temperate subnetworks were correlated, which would indicate that bird species generally display a proportional role in both biomes (Extended Data Fig. 5).

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data availability

All data used in the analyses are available through the Dryad Digital Repository (<https://doi.org/10.5061/dryad.15dv41nx3>). The dated phylogeny of seed plants (Spermatophyta) used to obtain our phylogenetic tree is available through GitHub (https://github.com/FePhyFoFum/big_seed_plant_trees/releases). Data on bird body weight used for size classification (Supplementary Fig. 2) were obtained from EltonTraits 1.0 available through Figshare (<https://doi.org/10.6084/m9.figshare.c.3306933>).

Code availability

The R scripts used to generate all results and figures are available through the Dryad Digital Repository (<https://doi.org/10.5061/dryad.15dv41nx3>).

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Acknowledgements The ‘Molecular Ecology Laboratory’ (LEM-EBD-CSIC; ISO9001:2015 and ISO14001:2015 certifications) and the ‘Research Unit of Biodiversity’ (UO-CSIC-PA) provided logistical support for molecular analyses. We thank L. Viesca and E. Cires for laboratory assistance, and J. M. Varela for the bird illustrations. Barcoding data were obtained within an Individual Fellowship from the Marie Skłodowska-Curie Actions (H2020-MSCA-IF-2014-656572: MobileLinks) and supported by a GRUPIN grant from the Regional Government of Asturias (IDI/2018/000151). ‘King Jaume I’ awarded to A.T. supported data collation during two postdoctoral contracts. J.P.G.-V. is supported by a Spanish ‘Ramón y Cajal’ fellowship (RYC-2017-22095) and a grant from the Spanish MICINN (PID2019-104922GA-I00/AEI/10.13039/501100011033). B.R. is supported by a Spanish ‘Juan de la Cierva Incorporación’ fellowship (IJC1-2017-33475). R.H.H. is funded by the Portuguese Foundation for Science and Technology (UID/BIA/04004/2020). B.I.S. is supported by a Royal Commission for the Exhibition of 1851 Research Fellowship. W.J.S. is funded by Arcadia.

Author contributions J.P.G.-V. conceived the study. J.P.G.-V., J.A., J.M.A., R.S.B., T.B., G.E.-Á., N.F., D.G., J.C.I., P.J., P.K., W.J.S. and E.V. obtained data of seven new seed-dispersal networks within the EU project ‘MobileLinks’. J.M.A. and J.C.I. also conducted the molecular analyses for these networks. L.P.d.S. and R.H.H. provided data of one unpublished network. P.J. provided data of two published networks. B.R., J.P.G.-V. and A.T. gathered data on fruiting phenology and bird migrations; J.P.G.-V., B.R., J.A. and B.I.S. analysed the data; J.P.G.-V. wrote the first manuscript draft, and all authors worked on the final version.

Competing interests The authors declare no competing interests.

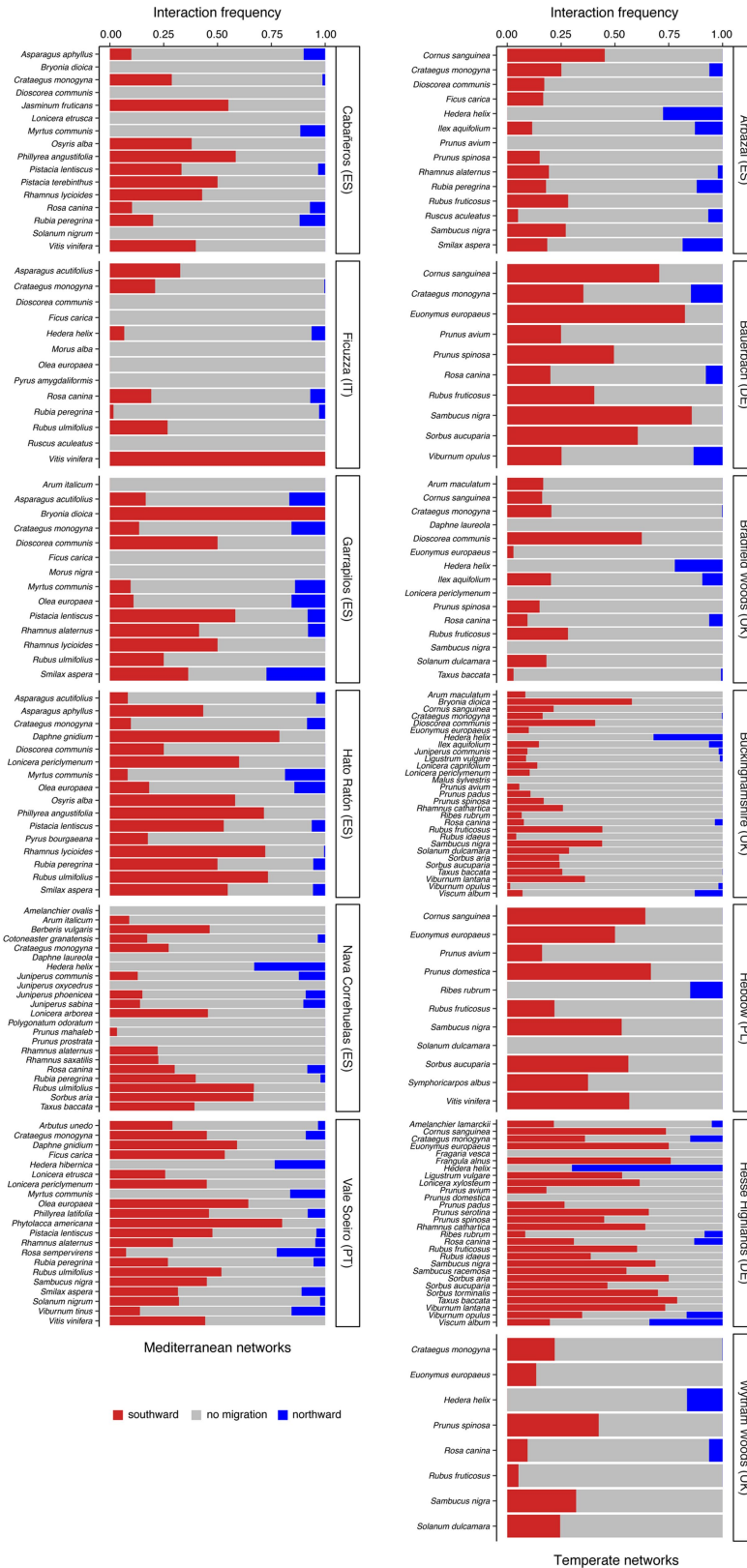
Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41586-021-03665-2>.

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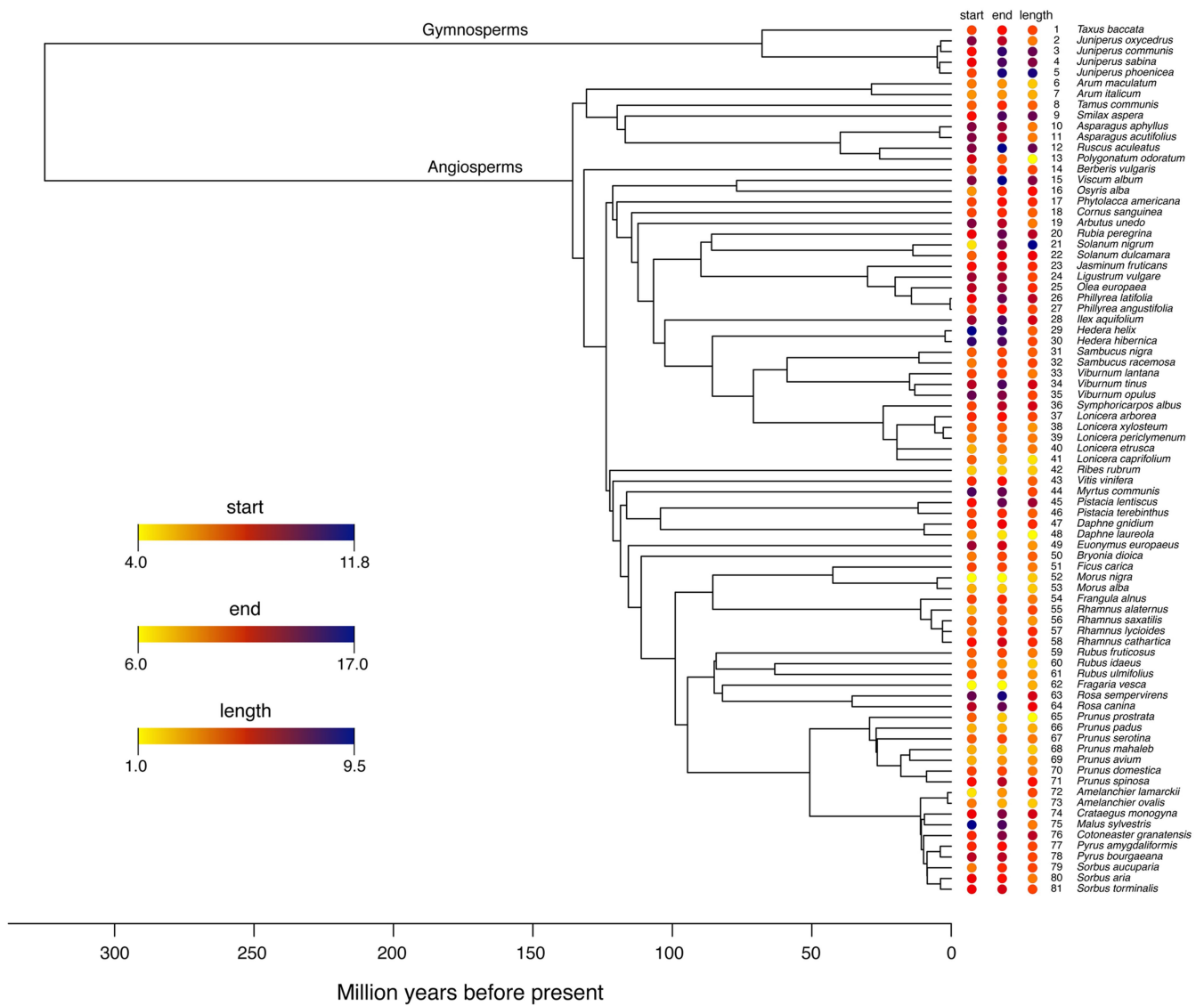
Peer review information Nature thanks Christian Both, Barnabas Daru, David Inouye, Duarte Viana and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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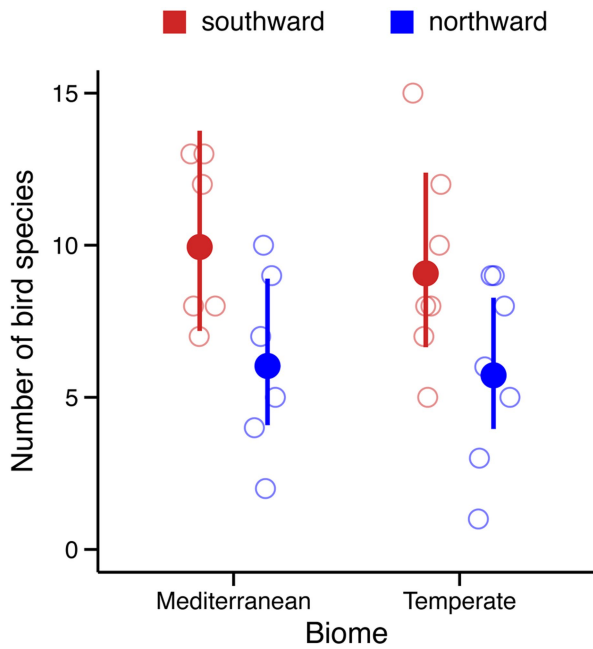
Extended Data Fig. 2 | Estimated interaction frequencies of plant species within each study network with birds migrating northwards, southwards or not migrating. Blue, interactions during northward migration; red, interactions during southward migration; grey, non-migration interactions.

Each panel represents a seed-dispersal network. The left column of panels includes Mediterranean networks, whereas the right column includes temperate networks. DE, Germany; ES, Spain; IT, Italy; PL, Poland; PT, Portugal; UK, United Kingdom.

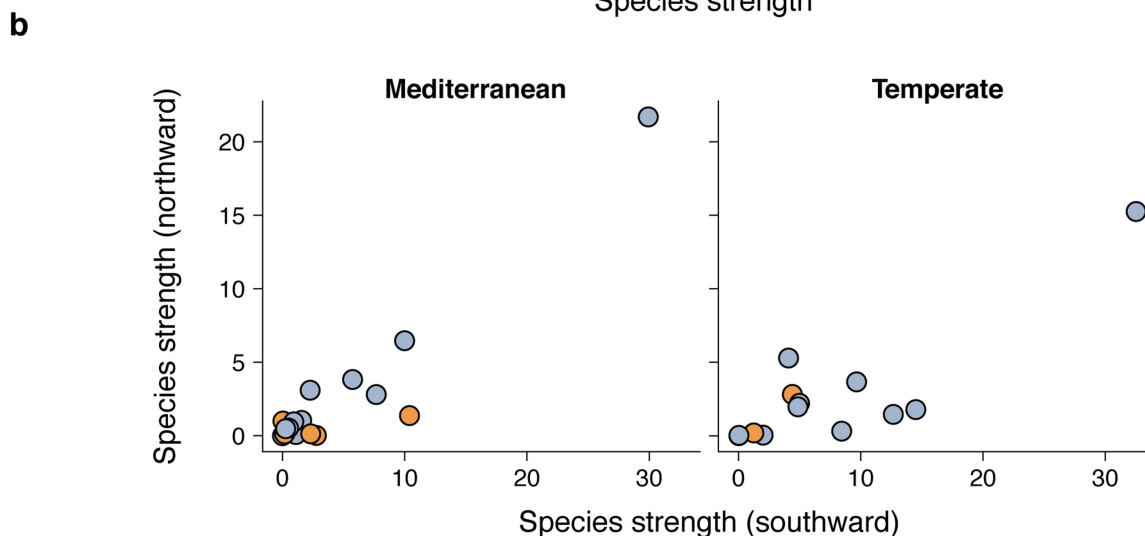
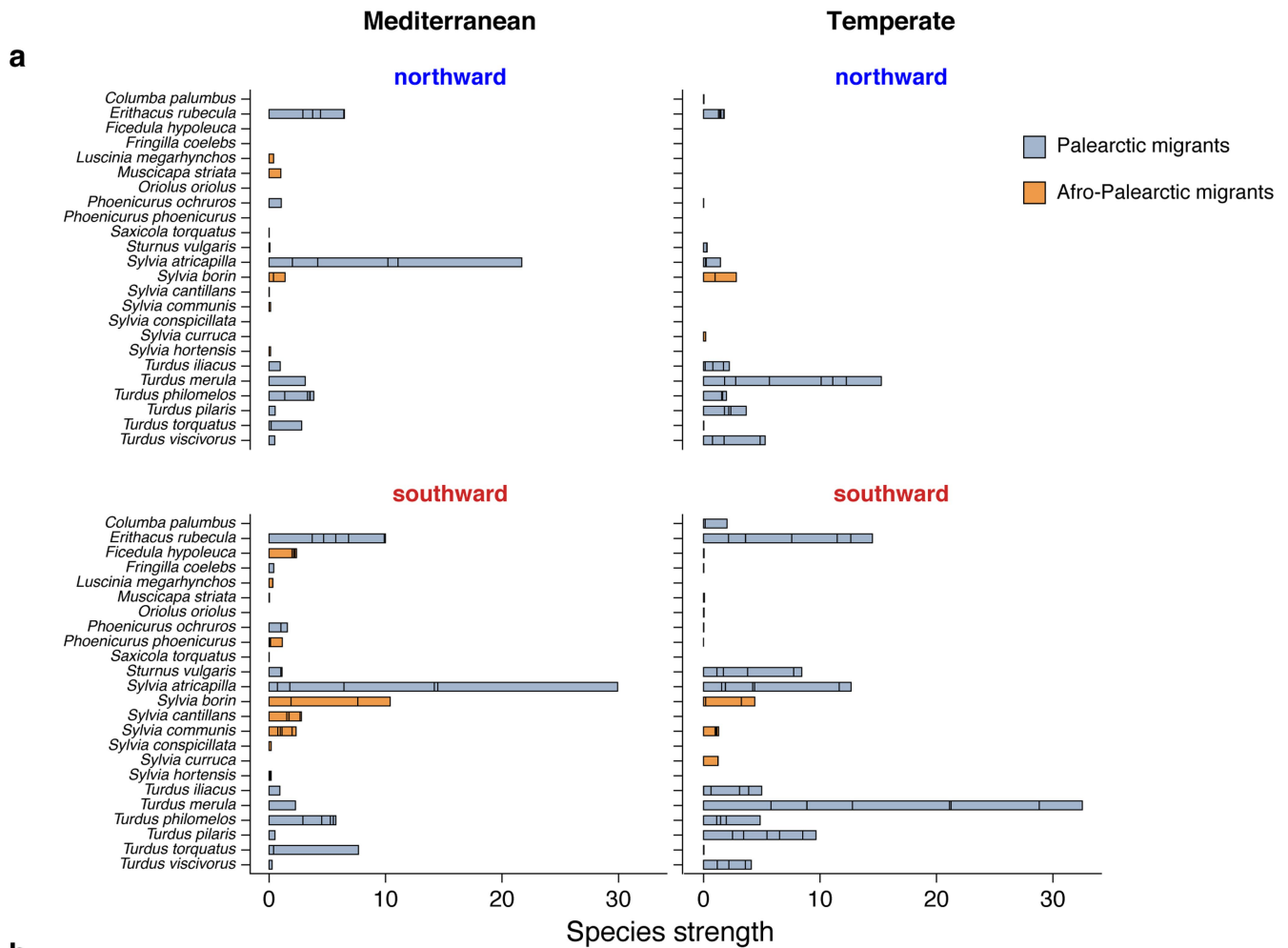


Extended Data Fig. 3 | Variables of the seed-dispersal phenology across the phylogenetic tree of plants. Phylogenetic signal was tested in plant-species means across networks in start and end dates (D_{start} and D_{end}), as well as in length ($D_{length} = D_{end} - D_{start}$) of the seed-dispersal period ($n = 81$ plant species) by means of Pagels' λ , as described in 'Phylogenetic signal in plants' in 'Statistical analyses' (Methods). The three phenological variables showed significant phylogenetic signal ($D_{start}, \lambda = 0.800, P = 0.0103$; $D_{end}, \lambda = 0.781, P = 0.0015$; and $D_{length}, \lambda = 0.419, P = 0.0343$). To test for phylogenetic signal, we previously calculated the species-level means for D_{start} , D_{end} and D_{length} across bioclimates (Extended Data Fig. 6). For this reason, we assessed the amount of variance in these phenological variables that is accounted for by bioclimate, as compared

to that accounted for by species through linear-mixed models (LMMs) that included 'bioclimate' as fixed factor and 'plant species' as random factor to account for the repeated measures per species. Bioclimate accounted for only a minor fraction of variance (1–3%) in D_{start} , D_{end} and D_{length} , as shown by the marginal R^2 values (variance explained by fixed effects; $R^2_{LMM(f)} = 0.028, 0.01$ and 0.023 , respectively). By contrast, the high conditional R^2 values (variance explained by both fixed and random effects; $R^2_{LMM(c)} = 0.780, 0.845$ and 0.643 , respectively) indicated that plant species accounted for most variance in the three phenological variables. LMMs were fitted with the R package package lme4 (v.1.1-19)¹⁰⁵.

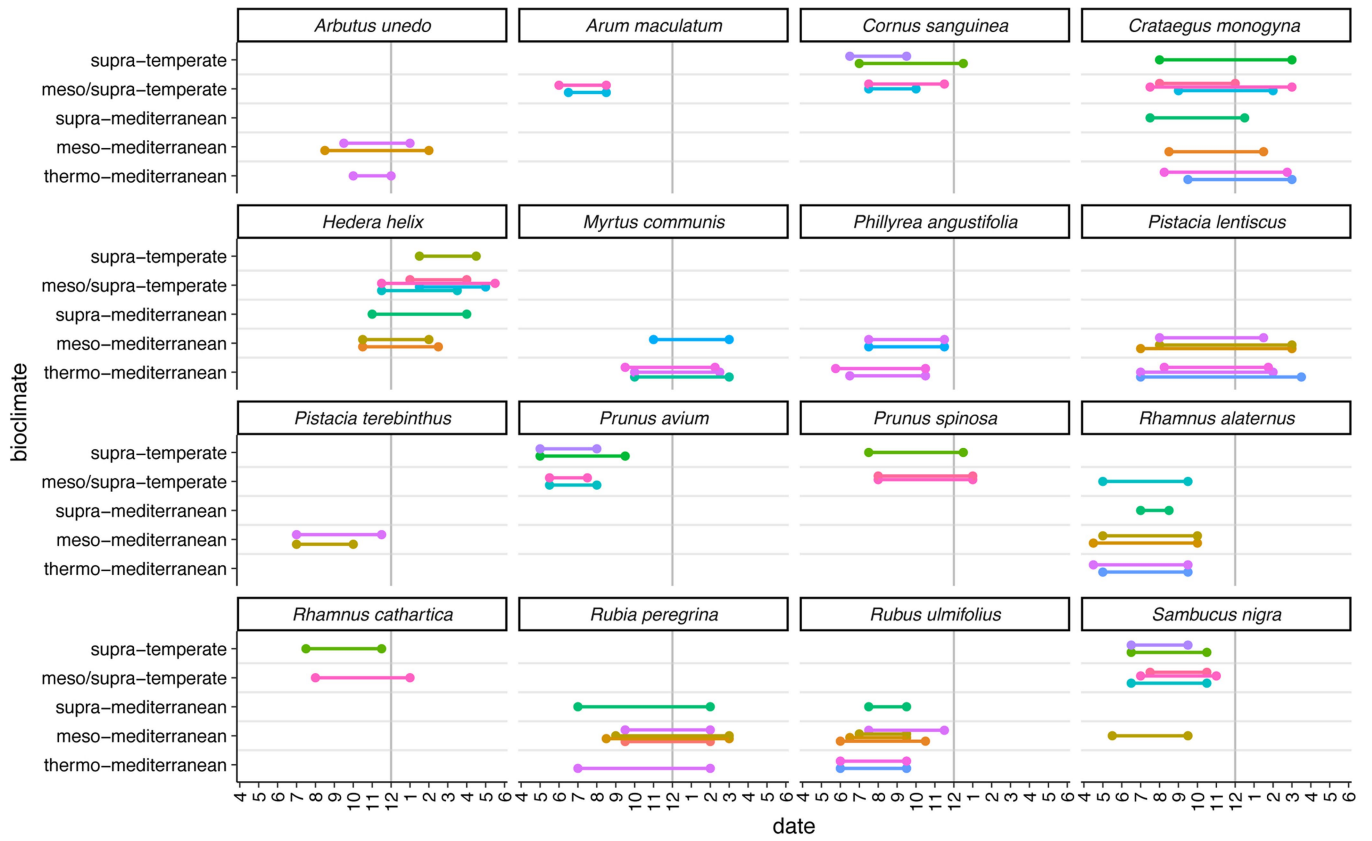


Extended Data Fig. 4 | Number of migratory bird species interacting with plants during migration per network in relation to migration direction and biome. Large dots and bars denote means \pm 95% confidence intervals estimated by a GLMM, whereas circles denote values for each seed-dispersal network ($n = 26$ observations, 13 networks \times 2 directions). Only migration direction had significant effects on the number of migratory bird species interacting with plants during migration in the GLMM (Poisson family and log-link function) testing the effects of migration direction (Wald $\chi^2 = 11.08$, $P = 0.0009$), biome (Wald $\chi^2 = 0.17$, $P = 0.6789$) and their interaction (Wald $\chi^2 = 0.02$, $P = 0.8921$). Model estimates \pm s.e.: intercept = 2.297 ± 0.156 ; direction (northward) = -0.500 ± 0.208 ; biome (temperate) = -0.091 ± 0.215 ; direction (northward) \times biome (temperate) = 0.039 ± 0.288 ; southward and Mediterranean were used as the reference categories (intercepts) for the factors direction and biome, respectively. A mean of 9.5 bird species per community dispersed plants during their southward migration, but only 5.9 species did so during the northward migration.



Extended Data Fig. 5 | Individual and cumulative bird species strengths accumulated across seed-dispersal subnetworks. a, Bird species strength accumulated across seed-dispersal subnetworks between plants and birds migrating southwards or northwards, and in Mediterranean and temperate biomes; species strength quantifies the relevance of a bird species across the entire fleshy-fruited plant community¹⁰⁴ ($n = 24$ species). Some bird species have stacked values from several subnetworks, whereas other species participated only in a single subnetwork. **b**, The cumulative species strength across the southward and northward subnetworks were significantly correlated in the Mediterranean (Kendall's $\tau = 0.396$, $P = 0.0129$) and the

temperate biome ($\tau = 0.588$, $P = 0.0006$), indicating that bird species generally display a proportional role in both migrations. However, the cumulative species strength in the Mediterranean and temperate biome were not correlated, neither across the northward ($\tau = 0.276$, $P = 0.1089$) nor across the southward subnetworks ($\tau = 0.263$, $P = 0.0764$) (correlation between left and right panels in **a**). These results indicate discordance between biomes in the identity of bird species contributions to community-wide seed dispersal during each migration. Pearson's r yielded qualitatively similar results, with higher coefficient values in the significant correlations ($r = 0.946$ and 0.847).



Extended Data Fig. 6 | Bioclimate-level plant phenology from several sources. Subset of 16 out of the 81 plant species present in the study networks illustrating how, in many cases, we obtained data on seed-dispersal phenology

from several sources for the same plant species–bioclimate combination. Colour codes denote different data sources. A vertical grey line divides the calendar year.

Extended Data Table 1 | Characteristics of the European seed-dispersal networks that we studied

N	Country	Network name	Source	Sampling type	Biome	Bioclimate	Latitude	Longitude	Years (n)	N_{plant}	N_{bird}	N_{int}
1	Spain	Hato Ratón	A	Mist-netting	Mediterranean	thermo-	37.1804	-6.3240	1981–1983 (2)	16	17	120
2	Spain	Nava Correhuelas	A	Observations	Mediterranean	supra-	37.9409	-2.7927	1997–1999 (2)	22	21	111
3	Spain	Garrapilos	B	DNA-barcoding	Mediterranean	thermo-	36.6589	-5.9493	2013–2015 (2)	14	21	56
4	Spain	Cabañeros	B	DNA-barcoding	Mediterranean	meso-	39.3213	-4.2896	2016–2017 (1)	16	14	44
5	Spain	Arbazal	B	DNA-barcoding	temperate	thermo/meso-	43.4313	-5.4971	2016–2017 (1)	14	14	52
6	Portugal	Vale Soeiro	C	Mist-netting	Mediterranean	meso-	40.3127	-8.4035	2012–2018 (6)	21	13	76
7	Italy	Ficuzza	B	DNA-barcoding	Mediterranean	meso-	37.8923	13.3749	2016–2017 (1)	13	12	30
8	UK	Buckinghamshire	D	Observations	temperate	meso/supra-	51.8910	-0.9120	1980–1985 (5)	29	19	204
9	UK	Wytham Woods	E	Observations	temperate	meso/supra-	51.7667	-1.3333	1979–1980 (1)	8	8	24
10	UK	Bradfield Woods	B	DNA-barcoding	temperate	meso/supra-	52.1808	0.8239	2016–2017 (1)	15	11	36
11	Germany	Hesse Highlands	F	Observations	temperate	supra-	51.3957	8.9427	1997–1999 (2)	28	18	128
12	Germany	Bauerbach	B	DNA-barcoding	temperate	supra-	50.7950	8.8230	2016–2017 (1)	10	9	30
13	Poland	Hebdów	B	DNA-barcoding	temperate	supra-	50.1429	20.4274	2016–2017 (1)	11	16	38

N_{plant} , N_{bird} and N_{int} denote the number of plant species, bird species and plant–bird interactions in each network, respectively. Network biomes were obtained from refs. ^{47,48}; network bioclimates were obtained from ref. ⁴⁸. Source A is ref. ⁴⁹; source B is this study (EU project MobileLinks); source C is this study (unpublished data provided by L.P.d.S. and R.H.H.); source D is ref. ⁵⁰; source E is ref. ²⁷; and source F is ref. ⁵¹.

Article

Extended Data Table 2 | List of bird and plant species of the 13 study networks

Bird species list		Plant species list			
Bird species	Bird family	Plant species	Plant family	Plant species	Plant family
<i>Alectoris rufa</i>	Phasianidae	<i>Amelanchier lamarckii</i>	Rosaceae	<i>Prunus avium</i>	Rosaceae
<i>Columba palumbus</i>	Columbidae	<i>Amelanchier ovalis</i>	Rosaceae	<i>Prunus domestica</i>	Rosaceae
<i>Corvus corax</i>	Corvidae	<i>Arbutus unedo</i>	Ericaceae	<i>Prunus mahaleb</i>	Rosaceae
<i>Corvus corone</i>	Corvidae	<i>Arum italicum</i> †	Araceae	<i>Prunus padus</i>	Rosaceae
<i>Corvus monedula</i>	Corvidae	<i>Arum maculatum</i> †	Araceae	<i>Prunus prostrata</i>	Rosaceae
<i>Cyanistes caeruleus</i>	Paridae	<i>Asparagus acutifolius</i>	Asparagaceae	<i>Prunus serotina</i>	Rosaceae
<i>Cyanopica cooki</i>	Corvidae	<i>Asparagus aphyllus</i>	Asparagaceae	<i>Prunus spinosa</i>	Rosaceae
<i>Dendrocopos major</i>	Picidae	<i>Berberis vulgaris</i>	Berberidaceae	<i>Pyrus amygdaliformis</i>	Rosaceae
<i>Emberiza calandra</i>	Emberizidae	<i>Bryonia dioica</i> †	Cucurbitaceae	<i>Pyrus bourgaeana</i> *	Rosaceae
<i>Erithacus rubecula</i>	Muscicapidae	<i>Cornus sanguinea</i>	Cornaceae	<i>Rhamnus alaternus</i>	Rhamnaceae
<i>Falco tinnunculus</i>	Falconidae	<i>Cotoneaster granatensis</i>	Rosaceae	<i>Rhamnus cathartica</i>	Rhamnaceae
<i>Ficedula hypoleuca</i>	Muscicapidae	<i>Crataegus monogyna</i>	Rosaceae	<i>Rhamnus lycioides</i>	Rhamnaceae
<i>Fringilla coelebs</i>	Fringillidae	<i>Daphne gnidium</i>	Thymelaeaceae	<i>Rhamnus saxatilis</i>	Rhamnaceae
<i>Gallinula chloropus</i>	Rallidae	<i>Daphne laureola</i>	Thymelaeaceae	<i>Ribes rubrum</i>	Grossulariaceae
<i>Garrulus glandarius</i>	Corvidae	<i>Euonymus europaeus</i>	Celastraceae	<i>Rosa canina</i>	Rosaceae
<i>Lanius excubitor</i>	Laniidae	<i>Ficus carica</i>	Moraceae	<i>Rosa sempervirens</i>	Rosaceae
<i>Luscinia megarhynchos</i>	Muscicapidae	<i>Fragaria vesca</i> †	Rosaceae	<i>Rubia peregrina</i> ‡	Rubiaceae
<i>Muscicapa striata</i>	Muscicapidae	<i>Frangula alnus</i>	Rhamnaceae	<i>Rubus fruticosus</i>	Rosaceae
<i>Oriolus oriolus</i>	Oriolidae	<i>Hedera helix</i> ‡	Araliaceae	<i>Rubus idaeus</i>	Rosaceae
<i>Parus major</i>	Paridae	<i>Hedera hibernica</i> ‡	Araliaceae	<i>Rubus ulmifolius</i>	Rosaceae
<i>Phasianus colchicus</i>	Phasianidae	<i>Ilex aquifolium</i>	Aquifoliaceae	<i>Ruscus aculeatus</i>	Asparagaceae
<i>Phoenicurus ochrurus</i>	Muscicapidae	<i>Jasminum fruticans</i>	Oleaceae	<i>Sambucus nigra</i>	Adoxaceae
<i>Phoenicurus phoenicurus</i>	Muscicapidae	<i>Juniperus communis</i>	Cupressaceae	<i>Sambucus racemosa</i>	Adoxaceae
<i>Pica pica</i>	Corvidae	<i>Juniperus oxycedrus</i>	Cupressaceae	<i>Smilax aspera</i> ‡	Smilacaceae
<i>Picus sharpei</i>	Picidae	<i>Juniperus phoenicea</i>	Cupressaceae	<i>Solanum dulcamara</i> †	Solanaceae
<i>Picus viridis</i>	Picidae	<i>Juniperus sabina</i>	Cupressaceae	<i>Solanum nigrum</i> †	Solanaceae
<i>Saxicola torquatus</i>	Muscicapidae	<i>Ligustrum vulgare</i>	Oleaceae	<i>Sorbus aria</i>	Rosaceae
<i>Sitta europaea</i>	Sittidae	<i>Lonicera arborea</i>	Caprifoliaceae	<i>Sorbus aucuparia</i>	Rosaceae
<i>Streptopelia decaocto</i>	Columbidae	<i>Lonicera caprifolium</i> ‡	Caprifoliaceae	<i>Sorbus torminalis</i>	Rosaceae
<i>Sturnus unicolor</i>	Sturnidae	<i>Lonicera etrusca</i> ‡	Caprifoliaceae	<i>Symphoricarpos albus</i>	Caprifoliaceae
<i>Sturnus vulgaris</i>	Sturnidae	<i>Lonicera periclymenum</i> ‡	Caprifoliaceae	<i>Dioscorea communis</i> †	Dioscoreaceae
<i>Sylvia atricapilla</i>	Sylviidae	<i>Lonicera xylosteum</i>	Caprifoliaceae	<i>Taxus baccata</i>	Taxaceae
<i>Sylvia borin</i>	Sylviidae	<i>Malus sylvestris</i>	Rosaceae	<i>Viburnum lantana</i>	Adoxaceae
<i>Sylvia cantillans</i>	Sylviidae	<i>Morus alba</i>	Moraceae	<i>Viburnum opulus</i>	Adoxaceae
<i>Sylvia communis</i>	Sylviidae	<i>Morus nigra</i>	Moraceae	<i>Viburnum tinus</i>	Adoxaceae
<i>Sylvia conspicillata</i>	Sylviidae	<i>Myrtus communis</i>	Myrtaceae	<i>Viscum album</i>	Santalaceae
<i>Sylvia curruca</i>	Sylviidae	<i>Olea europaea</i>	Oleaceae	<i>Vitis vinifera</i> ‡	Vitaceae
<i>Sylvia hortensis</i>	Sylviidae	<i>Osyris alba</i>	Santalaceae		
<i>Sylvia melanocephala</i>	Sylviidae	<i>Phillyrea angustifolia</i>	Oleaceae		
<i>Sylvia undata</i>	Sylviidae	<i>Phillyrea latifolia</i>	Oleaceae		
<i>Turdus iliacus</i>	Turdidae	<i>Phytolacca americana</i> †	Phytolaccaceae		
<i>Turdus merula</i>	Turdidae	<i>Pistacia lentiscus</i>	Anacardiaceae		
<i>Turdus philomelos</i>	Turdidae	<i>Pistacia terebinthus</i>	Anacardiaceae		
<i>Turdus pilaris</i>	Turdidae	<i>Polygonatum odoratum</i> †	Asparagaceae		
<i>Turdus torquatus</i>	Turdidae				
<i>Turdus viscivorus</i>	Turdidae				

We followed taxonomy from 'Birds of the World' (www.birdsoftheworld.org)⁷⁸ for birds and a previously published⁸⁶ phylogenetic tree (ALLMB) for plants. Plants are defined as herbs (†), woody vines (*) or trees and shrubs (all other species).

**Pyrus bourgaeana* (Iberian wild pear) was not present in ref. ⁸⁶ but 'World Flora Online' (www.worldfloraonline.org) considers this species as a synonym of *Pyrus communis* auct. iber. We thus matched *P. bourgaeana* to *P. communis* in the phylogenetic tree to test for phylogenetic signal.

Extended Data Table 3 | Significance of the fixed factors migration direction and biome, and their interaction, in GLMMs testing effects on seed-dispersal interactions of plants with migrating birds

Fixed-effects	(i) Proportion of plant species (Binomial, logit link)		(ii) Frequency of seed-dispersal interactions (Beta, logit link)		(iii) Number of bird species per plant (Poisson, log link)	
	χ^2	<i>P</i>	χ^2	<i>P</i>	χ^2	<i>P</i>
Hypothesis testing						
Direction (D)	51.02	2.0×10^{-16}	159.60	2.0×10^{-16}	5.75	0.0165
Biome (B)	0.09	0.7612	0.21	0.6452	0.67	0.4142
D × B	7.03	0.0080	6.51	0.0107	1.26	0.2623
Conditional model	Estimate ± se		Estimate ± se		Estimate ± se	
Intercept	1.414 ± 0.310		-0.418 ± 0.207		1.004 ± 0.125	
D (northward)	-1.734 ± 0.368		-1.842 ± 0.164		-0.307 ± 0.124	
B (temperate)	0.714 ± 0.426		-0.322 ± 0.274		0.085 ± 0.168	
D × B	-1.310 ± 0.494		0.642 ± 0.251		0.194 ± 0.173	
Dispersion model	Estimate ± se		Estimate ± se		Estimate ± se	
Intercept	-		1.028 ± 0.140		-	
D (northward)	-		1.874 ± 0.266		-	
B (temperate)	-		0.754 ± 0.198		-	
D × B	-		-1.575 ± 0.389		-	
Random effects	Variance		Variance		Variance	
Plant species: Network	0.368		3.4×10^{-9}		0.157	
Network	0.077		0.174		0.052	

Proportion of plant species interacting with birds during migration ($n = 434$ observations) (i) (Fig. 2a), frequency of seed-dispersal interactions with birds during migration whenever these interactions occurred (non-zero frequencies; $n = 260$ observations) out of the total interaction weight (ii) (Fig. 2b) and number of bird species dispersing each plant species during migration whenever these interactions occurred ($n = 260$ observations) (iii) (Fig. 2c). Family and link functions are shown in parentheses. All models included network identity and plant species nested within network as random factors to account for the repeated measures at these levels. Model (ii) also includes a dispersion model because the dispersion parameter Φ of the beta distribution was allowed to vary in response to the interactive effects of direction and biome⁹⁵. *P* values (two-sided) < 0.05 and significant model estimates ($P < 0.05$) are shown in bold. In all models, southward and Mediterranean were used as the reference categories (intercepts) for the factors direction (*D*) and biome (*B*), respectively.

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Extended Data Table 4 | Significance of the fixed factors migration direction and biome, and their interaction, in GLMMs testing effects on the proportion of migratory bird species that were Palaearctic migrants, and in the network-level frequency of seed-dispersal interactions with Palaearctic migrants

Fixed-effects	(i) Proportion of migratory bird species that were Palaearctic migrants (Binomial, logit link)		(ii) Interaction frequency during migrations with Palaearctic migrants (Beta, logit link)	
	χ^2	<i>P</i>	χ^2	<i>P</i>
Hypothesis testing				
Direction (D)	7.98	0.0047	32.47	2.0 × 10⁻¹⁶
Biome (B)	9.14	0.0025	12.98	0.0003
D × B	0.11	0.7458	7.12	0.0076
Conditional model	Estimate ± se		Estimate ± se	
Intercept	0.034 ± 0.363		1.188 ± 0.449	
D (northward)	1.004 ± 0.466		2.508 ± 0.503	
B (temperate)	1.429 ± 0.534		2.219 ± 0.531	
D × B	0.268 ± 0.825		-1.516 ± 0.568	
Dispersion model	Estimate ± se		Estimate ± se	
Intercept	–		1.318 ± 0.560	
D (northward)	–		3.404 ± 0.994	
B (temperate)	–		2.936 ± 0.742	
Random effects	Variance		Variance	
Network	0.336		0.088	

Family and link functions are shown in parentheses. Models included network identity as random factor to account for the repeated measures within networks ($n = 26$ observations, 13 networks × 2 directions). We used data only from Palaearctic migrants because the frequencies from both migrant types are fully interdependent (Fig. 3a, b). Model (ii) also includes a dispersion model because the dispersion parameter Φ of the beta distribution was allowed to vary in response to the additive effects of direction and biome⁹⁵. *P* values (two-sided) < 0.05 and significant model estimates ($P < 0.05$) are shown in bold. Results for the species richness of all migrant species pooled are provided in Extended Data Fig. 4. In all models, southward and Mediterranean were used as the reference categories (intercepts) for the factors direction (*D*) and biome (*B*), respectively.

Reporting Summary

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For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

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- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
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- For null hypothesis testing, the test statistic (e.g. F , t , r) with confidence intervals, effect sizes, degrees of freedom and P value noted
Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated

Our web collection on [statistics for biologists](#) contains articles on many of the points above.

Software and code

Policy information about [availability of computer code](#)

Data collection

Our study includes both data collection (field sampling and molecular analyses) and data compilation from previous studies. We used SEQUENCHER v. 4.9, BioEdit v. 7.0.9 and Chromas v. 2.5.1 for sequence alignment and editing. We used the online platforms BOLD (www.boldsystems.org) and BLAST (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>) for species identification from amplified DNA sequences (i.e. DNA barcoding).

Data analysis

We conducted all analyses in R version 3.5.2. We used the R packages glmmTMB (v. 0.2.3), car (v. 2.1-6), ape (v. 5.3), phytools (v. 0.6-99), lme4 (v. 1.1-19) and bipartite (v. 2.13). We made the figures using the R packages 'ggplot2' (v. 3.3.0) and cowplot (v. 0.9.4). The R scripts used to generate all results and figures are available through the Dryad Digital Repository (<https://doi.org/10.5061/dryad.15dv41nx3>).

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- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

All data used in the analyses are available through the Dryad Digital Repository (<https://doi.org/10.5061/dryad.15dv41nx3>). The dated phylogeny of seed plants (Spermatophyta) used to obtain our phylogenetic tree is available through GitHub (https://github.com/FePhyFoFum/big_seed_plant_trees/releases). Data on bird

Field-specific reporting

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Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

Study description	We combined phenological and migration information with data on 949 seed-dispersal interactions between 46 bird and 81 plant species from 13 woodland communities across Europe. The study design is factorial, as we tested the interactive effect between migration direction and community biome (direction x biome) on different response variables. The data structure is hierarchical because we have species-level data for plants at birds nested within sites. The data structure also has repeated (paired) measures, with a data points per migration direction (southward and northward).
Research sample	<p>Research sample includes seed-dispersal networks, that is, interacting communities of 46 frugivorous bird species that disperse the seeds of 81 fruiting plant species from 13 woodland communities across Europe (see Extended Data Table 2). Seed dispersal networks are expressed as interaction matrices where each row i represents a plant species and each column j represents a bird species. Elements in the matrices (w_{ij}) denote whether pairwise plant-bird interactions were observed ($w_{ij} > 0$) or not ($w_{ij} = 0$) and, if so, their value account for interaction weight. In this case, the quantity of seeds of each plant species dispersed by each bird species.</p> <p>The total sample size for plants, the main subject of the study, includes 434 observations (i.e. "plant-species / site / migration" combinations).</p> <p>Sex and age of the individuals is not relevant for the purposes of this study.</p>
Sampling strategy	All the study networks ($n = 13$) were sampled all year-round, a prerequisite to cover the entire fruiting periods of all local fleshy-fruited species, as well as the prenuptial and postnuptial migration periods of all migratory birds. Our study includes a combination of networks from previous studies ($n = 5$) and newly sampled networks ($n = 8$). The number of networks was evenly distributed between the Mediterranean ($n = 6$) and the temperate biomes ($n = 7$) of Europe, which allowed a sufficient sample size to test for biome effects.
Data collection	<p>Seven of the eight new networks were sampled through field sampling of bird-dispersed seeds and subsequent disperser identification by means of DNA-barcoding analysis. Individual seeds or droppings with seeds were sampled for DNA-barcoding analysis into 1.5- or 2.0-ml sterile tubes that were labelled and stored in a freezer at -20°C until DNA extraction. The molecular analyses for species identification was conducted by J.M.A. and J.C.I.</p> <p>The other new network was sampled through dietary analysis of birds captured in mist nets. Captured birds were individually placed in ringing bags for up to 30 min until they ejected droppings.</p> <p>The networks compiled from previous studies were obtained either through focal plant observations of birds feeding on fruits or through dietary analysis of birds captured in mist nets. Bird samples were collected in the field by J.P.G.-V., J.A., J.M.A., R.S.B., T.B., G.E.-A., N.F., D.G., J.C.I., P.J., P.K., W.J.S., E.V., L.P.d.S. and R.H. The networks from previous studies were compiled by B.I.S. Data on fruiting phenology and bird migrations was gathered by B.R., J.P.G.-V. and A.T.</p>
Timing and spatial scale	New data in this study was collected between 2012 and 2018. Data available from previous studies were collected between 1981 and 1999. Sampling consisted on periodical surveys during 1–6 years, generally every 2 weeks, in which seed traps were revised, birds were mist-netted or focal plants were observed. Such frequency of sampling during all-year-round allows recording interactions between all bird and plant species of each community, regardless the season. See details in Table S1. The data was collected in 13 sites across Europe and each site spans several square kilometers.
Data exclusions	No data were excluded.
Reproducibility	The analyses conducted in this study can be reproduced using the R code, which we will made publicly available once the article is accepted for publication. Moreover, we will make available in the DRYAD repository the complete dataset with the detailed information for each sample and its corresponding sequence obtained through DNA barcoding. The sequences include the sample code and all information associated: sampling date, site, seed species and number of seeds of the sample, bird species identified through DNA barcoding, percentage of similarity with best matched sequence and GenBank accession number of best matching sequence. In addition, we keep the DNA aliquots of all DNA extractions from each sample (voucher numbers correspond with 'sample_code') in our labs at University of Oviedo and EBD-CSIC (Spain). These aliquots are available under request. All seed samples are stored by J.P.G.-V at the laboratory of Botany in the University of Cádiz (Spain).
Randomization	Randomization does not apply as we did not conduct experiments. However, we did assess potential biases regarding the sampling methods of study networks and the approach used to obtain seed-dispersal periods (see Supplementary Discussion 2; Supplementary Figures 5–7).

Blinding

Did the study involve field work? Yes No

Field work, collection and transport

Field conditions

Location

Access & import/export - Cabañeros (Spain): national park; permission given by the director of the "Cabañeros National Park" to E.V. (June 2016).
- Arbazal (Spain): private site; permission given by local authority and owners to D.G. (February 2016).
- Vale Soeiro (Portugal): permission for mist-netting and bird ringing given by ICNF ("Instituto da Conservação da Natureza e das Florestas") to L.P.d.S. (117/2012, 126/2013, 130/2014, 137/2015, 140/2016, 146/2017 and 123/2018).
- Ficuzza (Italy): private site; permission given by owners to R.S.B. (May 2016).
- Bradfield Woods (UK): natural reserve; permission given by "Suffolk Wildlife Trust" to J.P.G.-V. and W.J.S (30 March 2016).
- Bauerbach (Germany): public site; permission by local authority to N.F. (6 April 2016).
- Hebdów (Poland): public and private sites; permission by local authority and owners to P.K. (May 2016)."/>

Disturbance

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Animals and other organisms

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Laboratory animals

Wild animals

Sex and age of the individuals is unknown (and not relevant for this study).

Field-collected samples

Seed samples collected in the field were stored in the freezer at -20°C until DNA-barcoding analyses.

Ethics oversight

No ethical approval was required as no bird was killed, injured or kept captive and we used normal procedures for mist-netting.

Note that full information on the approval of the study protocol must also be provided in the manuscript.