The mutualism–antagonism continuum in Neotropical palm–frugivore interactions: from interaction outcomes to ecosystem dynamics

Caroline Marques Dracxler* and W. Daniel Kissling

Institute for Biodiversity and Ecosystem Dynamics (IBED), University of Amsterdam, P.O. Box 94240, Amsterdam, 1090 GE, The Netherlands

ABSTRACT

Frugivory, that is feeding on fruits, pulp or seeds by animals, is usually considered a mutualism when interactions involve seed dispersal, and an antagonism when it results in the predation and destruction of seeds. Nevertheless, most frugivory interactions involve both benefits and disadvantages for plants, and the net interaction outcomes thus tend to vary along a continuum from mutualism to antagonism. Quantifying outcome variation is challenging and the ecological contribution of frugivorous animals to plant demography thus remains little explored. This is particularly true for interactions in which animals do not ingest entire fruits, that is in seed-eating and pulp-eating. Here, we provide a comprehensive review of Neotropical palm–frugivore interactions, with a focus on how frugivore consumption behaviour (i.e. digestive processing, fruit-handling ability and caching behaviour) and feeding types (fruit-eating, pulp-eating and seed-eating) influence interaction outcomes at different demographic stages of palms. We compiled a total of 1043 species-level palm–frugivore interaction records that explicitly captured information on which parts of palm fruits are eaten by animals. These records showed consumption of fruits of 106 Neotropical palm species by 273 vertebrate species, especially birds (50%) and mammals (45%), but also fish (3%) and reptiles (2%). Fruit-eating involved all four taxonomic vertebrate classes whereas seed-eating and pulp-eating were only recorded among birds and mammals. Most fruit-eating interactions (77%) resulted in positive interaction outcomes for plants (e.g. gut-passed seeds are viable or seeds are successfully dispersed), regardless of the digestive processing type of vertebrate consumers (seed defecation versus regurgitation). The majority of pulp-eating interactions (91%) also resulted in positive interaction outcomes, for instance via pulp removal that promoted seed germination or via dispersal of intact palm seeds by external transport, especially if animals have a good fruit-handling ability (e.g. primates, and some parrots). By contrast, seed-eating interactions mostly resulted in dual outcomes (60%), where interactions had both negative effects on seed survival and positive outcomes through seed caching and external (non-digestive) seed dispersal. A detailed synthesis of available field studies with qualitative and quantitative information provided evidence that 12 families and 27 species of mammals and birds are predominantly on the mutualistic side of the continuum whereas five mammalian families, six mammal and one reptile species are on the antagonistic side. The synthesis also revealed that most species can act as partial mutualists, even if they are typically considered antagonists. Our review demonstrates how different consumption behaviours and feeding types of vertebrate fruit consumers can influence seed dispersal and regeneration of palms, and thus ultimately affect the structure and functioning of tropical ecosystems. Variation in feeding types of animal consumers will influence ecosystem dynamics via effects on plant population dynamics and differences in long-distance seed dispersal, and may subsequently affect ecosystem functions such as carbon storage. The quantification of intra- and inter-specific variation in outcomes of plant–frugivore interactions – and their positive and negative effects on the seed-to-seedling transition of animal-dispersed plants – should be a key research focus to understand better the mutualism–antagonism continuum and its importance for ecosystem dynamics.

Key words: Arecaceae, ecosystem functioning, ectozoochory, endozoochory, granivory, palm endocarp, seed caching, seed dispersal distance, seed predation, vertebrate

* Address for correspondence (Tel: +31 (0)20 525 6635; E-mail: carolinemdsbio@yahoo.com.br)
I. INTRODUCTION

Plant–frugivore interactions are widely considered mutualisms because the consumption of fruits by frugivores mostly results in seed dispersal, a process that is crucial for the establishment of animal-dispersed plant populations ( Jordano, 1987; Wang & Smith, 2002; Schupp, Jordano & Gómez, 2010; Snell et al., 2019). Nevertheless, it is also recognized that frugivores can positively influence seed viability by their direct or indirect effects on seeds (e.g. gut treatment of seeds or via other feeding types for both animals, the benefits come from food and energy resources provided by the plants whereas for plants the benefits depend on the effects of frugivores on various demographic stages during the seed-to-seedling transition. For instance, frugivores can positively influence seed viability by their direct or indirect effects on seeds (e.g. via gut treatment of seeds or seed caching), influencing whether seeds survive interactions or not ( Schupp et al., 2010). Moreover, frugivores also determine the sites of seed deposition, resulting in benefits for
plants if deposition sites match germination and/or establishment requirements (Cortes & Uriarte, 2013; Snell et al., 2019). At this stage, aspects such as dispersal distances and the quality of deposition microsites, which aggregate biotic conditions (e.g. conspecific and heterospecific seed density, levels of seed predation) and abiotic factors (e.g. luminosity and soil characteristics), determine whether outcomes are positive or negative for the plant (Schupp et al., 2010; Lopez-Toledo et al., 2013; Snell et al., 2019). These combined effects ultimately determine the influence of frugivory on seedling establishment (Beckman & Rogers, 2013). Importantly, frugivores can exert contrasting effects on different demographic processes, for instance when viable seeds (i.e. after a positive effect on seed viability) are deposited in places that are unsuitable for germination (i.e. via a negative effect at the seed-deposition stage). When interactions with frugivores involve only disadvantages for the plant (e.g. predation of seeds or seed dispersal only to sites which are unsuitable for germination), they are considered antagonistic (Bronstein, 2001). Nevertheless, most plant–frugivore interactions involve both benefits and disadvantages, and interaction outcomes can thus vary along a continuum from mutualism to antagonism (Thompson, 1988; Bronstein, 1994; Montesinos-Navarro et al., 2017; Rodriguez-Rodriguez, Jordano & Valido, 2017; Gomez et al., 2019).

A key element for determining the effects of frugivores on plants is how frugivores interact with fruits and/or seeds (hereafter ‘types of feeding interactions’ or ‘feeding guilds’). For instance, animals can ingest whole fruits (‘fruit-eaters’), they may only consume the pulp and thus discard the seeds (‘pulp-eaters’), or they may only feed directly on the seeds and thus discard the pulp (‘seed-eaters’). These feeding behaviours determine to what extent different plant demographic processes can be affected by frugivores and influence whether interaction outcomes are positive or negative. Interaction outcomes can vary further among different animal species within the same feeding guild. For instance, fruit-eaters often have a positive influence on seed viability and seed germination due to the gut passage of seeds, but mechanical and chemical effects associated with gut treatment may also decrease seed viability (Traveset, 1998; Través et al., 2002). Seed-eaters mostly have negative effects on seed survival because the seeds are their main food target (Janzen, 1971; Hulme & Benkman, 2002). Nevertheless, some seed-eaters such as scatter-hoarding rodents are known to be effective seed dispersers of some plant species because some of the seeds placed in caches can escape retrieval and predation if caches are neglected or abandoned (Vander Wall, 1990; Vander Wall & Beck, 2012; Zwolak & Crone, 2012; Lichi, Steele & Swihart, 2017; Mittelman et al., 2021). This might be particularly true for plants that produce large, long-lasting seeds that are perceived by rodents as good items for food storage. Pulp-eaters do not necessarily prevent or favour seed dispersal (potentially playing a neutral role), but pulp removal can favour seed escape and increase seed viability and/or seed germination (Loayza & Knight, 2010; Fedriani & Delibes, 2013). In some cases, pulp-eaters can also damage seeds during pulp consumption, therefore acting as seed predators because of their negative effects on seed viability (Simmons et al., 2018). Hence, the position of each frugivore species varies along the mutualism–antagonism continuum, with fruit-eaters being more likely located towards the mutualism end and seed-eaters being found more to the antagonistic end. However, interaction outcomes can even differ for the same animal species (Zwolak, 2018; Schupp et al., 2019) when feeding on different plant species or different parts of the fruits. This makes the quantification of ecological roles of frugivores a challenging task.

Here, we review the outcomes of plant–frugivore interactions in Neotropical palms (Areaceae) to assess our current knowledge of the ecological effects of frugivores on the seed-to-seedling transition and their placement along the mutualism–antagonism continuum. Palms are particularly abundant in the Neotropics (ter Steege et al., 2013; Muccarrella et al., 2020) and this biogeographic realm also contains a tremendous diversity of frugivores (Howe & Smallwood, 1982; Fleming, Breitwisch & Whitesides, 1987; Kissling, Böhning-Gaese & Jetz, 2009). We focus on palms because they are an excellent model system to understand the ecology and evolution of tropical rainforests and their biota (Henderson, 2002; Eiserhardt et al., 2011; Couvreur & Baker, 2013; Onstein et al., 2017), and because palms provide key food resources for many frugivores, including mammals, birds, reptiles, beetles, crabs, and fish (Zona & Henderson, 1989; Andreazzi, Pires & Fernandez, 2009; Muñoz, Trojelsgaard & Kissling, 2019). They are also a key resource for people and an important part of the livelihoods of rural populations, providing important ecosystem services (Cámara-Leret et al., 2017). With approximately 2,500 palm species distributed globally, and about 700 species in the Neotropics (Kissling et al., 2012), fruits and seeds of palms represent a large part of the diet of many animal species, and exhibit fruit traits that are clearly attractive for animals that feed on fruits, pulp and seeds (Kissling et al., 2019). We conducted a comprehensive literature review with the aim to identify different types of feeding interactions (i.e. fruit-eating, pulp-eating and seed-eating) and the influence of consumption behaviour (digestive processing, caching behaviour and fruit-handling ability) on outcomes of Neotropical palm–frugivore interactions at different palm demographic stages (Fig. 1). We gathered information on animal traits such as the digestive processing by fruit-eaters (i.e. regurgitation or defecation of seeds), seed caching behaviour by seed-eaters and fruit-handling ability of pulp- and seed-eaters. This allowed us to quantify how intra- and inter-specific variation of frugivory affects interaction outcomes. Specifically, we focus on the effects of frugivory on three key demographic processes of the seed-to-seedling transition of plants, namely seed viability, seed deposition and seedling establishment. Effects of frugivores on each of these three demographic processes can vary both intra- and inter-specifically, resulting in variation along the mutualism-antagonism continuum. We finally discuss important mechanisms of fruit-, pulp- and seed-eating, how they influence interaction outcomes and ultimately ecosystem
processes such as seed dispersal and seedling establishment, and which implications these interactions have for ecosystem dynamics such as carbon storage and nutrient cycling.

II. METHODS

(1) Compilation of interaction records and types of feeding interactions

We compiled interaction records of animals feeding on Neotropical palm fruits by screening articles from a comprehensive literature search on the Web of Science (WoS) conducted from October 2019 to July 2021. The Neotropical realm comprises South America, Central America, parts of Mesoamerica (Yucatan Peninsula), the Caribbean, and the southern parts of Texas and Florida in the USA (Udvardy, 1975). We used a combination of search terms that included ‘fruit removal’, ‘seed dispersal’, ‘seed predation’, ‘frugivory’, ‘granivory’, ‘seed’, ‘fruit’, ‘palm’, ‘Arecaceae’, ‘Neotropics’ and all Neotropical palm genera and country names (see online Supporting Information, Appendix S1, for the precise combination of search terms). We also conducted searches on the WoS using the same search terms translated into French, Portuguese and Spanish, since these languages are also used for publications in peer-reviewed journals in the Neotropics. This combination of terms allowed us to identify studies that included information on frugivore species feeding on the fruits and seeds of palm species in the entire Neotropics.

We obtained an initial list of 590 articles. All articles were manually and meticulously screened for information on interactions between frugivores and palm fruits. We also occasionally added additional literature that was cited in the screened articles. We extracted pairwise interaction records from an article when explicit information on animal species feeding on fruits, pulp and seeds of palm species was provided. This was most often referred to as frugivory, granivory, scatter-hoarding, fruit removal, seed dispersal, seed predation, fruit consumption, seed consumption or pulp consumption, but pairwise interactions could also be extracted when articles provided information on seed defecation, regurgitation, seed hoarding or seed caching. We compiled pairwise interactions regardless of how the interaction outcomes
were classified or which type of feeding the animals were displaying. We did not include interaction records when specific species involved in pairwise interactions were not clearly identified taxonomically, when evidence was only anecdotal, or when animal identity was only assumed by the authors (e.g. studies assessing fruit removal with no visual evidence of which animal species had actually removed the fruits). A total of 205 articles contained species-level information on pairwise interactions between animals and palm fruits, pulp and seeds.

Beyond simply recording the interaction between a palm and a frugivore, we aimed to identify the type of feeding in Neotropical palm–frugivore interactions (Fig. 1). We distinguished three different feeding interaction types: (i) fruit-eating (animals ingesting entire fruits), (ii) pulp-eating (animals feeding on fruit mesocarps only), and (iii) seed-eating (animals discarding fruit mesocarps and feeding only on seeds). This classification was done at the level of each individual interaction record to capture intra-specific variability because feeding behaviour of the same frugivore species may vary for different palm species. All species-level interaction records that specifically contained information on types of feeding interactions were considered eligible and included in this review (the data set is available from Dracxler & Kissling, 2021).

(2) Classification of interaction outcomes

We further classified the outcome of each interaction record (where possible) as (i) positive, (ii) negative or (iii) dual for all three feeding interaction types (Fig. 1). Evidence on interaction outcomes came from both quantitative and qualitative information in the screened articles, e.g. field observations or experimental quantification of seed dispersal, seed caching, seed predation, viability and germination of gut-passed or handled seeds, or seedling establishment (see criteria in Appendix S2). Available quantitative data on interaction outcomes included the number of seeds dispersed versus predated, viable versus non-viable gut-passed seeds, and success of seed germination or seedling establishment. From this information, we calculated the proportion of positive events (proportion of seeds dispersed, proportion of viable seeds after gut passage, proportion of seeds germinated, etc.) and classified interaction outcomes as positive if the proportion of positive events was ≥50%, and as negative if <50%. Since quantitative evidence is relatively scarce, we additionally extracted qualitative evidence on interaction outcomes. This reflected explicit mentioning in the articles that, for instance, an animal species acted as seed disperser (positive outcome), seed predator (negative) or both (dual). For dual interaction outcomes, seeds could be both predated and dispersed, without knowing the exact fate of the seeds.

(3) Consumption behaviour

We also compiled information on consumption-related behavioural traits of frugivores for all three feeding interaction types (fruit-, pulp-and seed-eating) because this can influence interaction outcomes (Simmons et al., 2018) (Fig. 1). For fruit-eating, we differentiated for each interaction record (whenever possible) how animals expel seeds (i.e. by seed regurgitation, defecation, or both) (Dracxler & Kissling, 2021) because this relates to the gut treatment of seeds which can influence, for instance, seed viability, the quality of seed deposition sites and dispersal distances (Schupp et al., 2010). For seed-eating, we distinguished whether seed-eaters show caching behaviour or not to evaluate if mechanisms related to seed caching (e.g. seed cleaning, dispersal and burial) influence interaction outcomes. Information on presence of seed-caching behaviour was compiled for each animal species involved in at least one seed-eating interaction (Dracxler & Kissling, 2021), either from articles containing the interaction records or from additional literature sources that described the behaviour of the animal at the species level. For seed-eating and pulp-eating, we further classified the animals’ fruit-handling ability into low, intermediate, or high (Fig. 1). Differences in fruit-handling ability can affect feeding efficiency and consumption rates, as well as dispersal and interaction outcomes (Levey, 1987; Feer et al., 2001; de Araújo & Marcondes-Machado, 2011; Fuzessy, Janson & Silveira, 2018). Examples of differences in fruit-handling ability are the direct consumption of pulp or seeds by tanagers (low fruit-handling ability), the holding of fruits or seeds by macaws with their feet (intermediate fruit-handling ability), and the use of forepaws and sitting behaviour to manipulate fruits and seeds by scatter-hoarding rodents such as agoutis or primates (high fruit-handling ability). Such information was also compiled at the animal species level (Dracxler & Kissling, 2021).

(4) Mutualism–antagonism continuum

To assess interaction outcomes along the mutualism–antagonism continuum, we aggregated outcomes (positive or negative) either at the species level (summarizing intra-specific variability) or at the family level (summarizing inter-specific variability) [see Gómez et al., (2019) for a variation of the continuum approach]. If the aggregated proportion of positive outcomes was ≥0.5, the frugivore species or family was considered to be mostly mutualistic, and if <0.5, then mostly antagonistic. We placed taxa along the mutualism–antagonism continuum using either qualitative or quantitative data on interaction outcomes extracted from the screened articles (see Section II.2). Qualitative data could be assessed at both the species and the family level whereas quantitative data were only available at the species level. We used a minimum of seven studies with qualitative outcomes to aggregate information at the family level along the continuum, and we only included families represented by at least two species (to show inter-specific variation of outcomes within families). For species-level continuums, we summarized outcomes for species with at least five outcome records. Species-level qualitative data on outcomes were abundant, but quantitative data could be summarized for only three species of frugivores for which...
≥5 interaction records reported in the screened articles provided evidence on interaction outcomes from field observations or experiments.

III. EXTENT OF THE LITERATURE REVIEW

Article screening allowed for the compilation of 3492 interaction records, from which 3230 described evidence of an animal species feeding on a particular palm species. A total of 1043 species-level records (32% of all records) recorded in 168 articles contained information on which parts of fruits animals feed upon (e.g. type of feeding interaction) (Dracxler & Kissling, 2021) and were therefore eligible for the classification of interaction outcomes and for inclusion in our review (Fig. 2). Interaction records with information on parts of fruits that animals consume were available from Argentina, Belize, Bolivia, Brazil, Colombia, Costa Rica, Ecuador, French Guiana, Guatemala, Honduras, Mexico, Panama, Peru, Trinidad and Tobago, Venezuela and the USA. Pairwise interaction records were obtained either from reviews (e.g. Beck, 2006; Gómez et al., 2019; Mendes, Koprowski & Galetti, 2019) or from original articles. Quantitative and/or qualitative information on outcomes available in the screened studies allowed us to classify interaction outcomes for 855 records (82%) with information on feeding interaction types (Fig. 2; Table 1) (Dracxler & Kissling, 2021).

IV. TYPES OF FEEDING INTERACTIONS AND THEIR TAXONOMIC COMPOSITION

About half of the records with information on the type of feeding interaction involved fruit-eating (52%), followed by seed-eating (37%) and pulp-eating (11%) (Dracxler & Kissling, 2021). These records included a total of 273 animal species recorded to feed on the fruits, pulp or seeds of 106 palm species (Table 2) belonging to 13 palm tribes and 40 palm genera (Appendix S3; Dracxler & Kissling, 2021). The majority of animal species recorded were birds and mammals (Table 2). Both groups also showed the largest amount of (total and unique) interaction records, with the number of interaction records being larger for mammals than for birds (Table 2). The diversity of animal families represented in records with data on feeding interaction types was largest in birds (N = 30 families), followed by mammals (N = 28), fish (N = 5) and reptiles (N = 5), while the diversity

---

![Flow diagram](image-url)  
**Fig. 2.** Flow diagram depicting the identification of articles, their manual screening, and reasons for eligibility of interaction records and their inclusion or exclusion. The structure of the flow diagram follows the PRISMA statement for transparent reporting of systematic reviews (Moher et al., 2009).
of animal genera was higher in mammals (N = 87 genera), followed by birds (N = 76), fish (N = 9) and reptiles (N = 5) (Dracxler & Kissling, 2021).

Birds acted mostly as fruit-eaters, but some families such as the parrots (Psittacidae) are recorded to feed mainly on pulp or seeds of palm fruits (Appendix S4). All records involving reptile and fish species suggested the ingestion of entire palm fruits (Fig. 3). By contrast, mammals predominantly fed on seeds, although consumption of entire fruits was also high (Fig. 3). Only birds and mammals acted as pulp-eaters (Fig. 3).

(1) Fruit-eaters
A total of 189 frugivore species were recorded in fruit-eating interactions with 80 palms, totalling 545 interaction records. The majority (58%) of fruit-eaters were birds, including toucans (Ramphastidae), parrots (Psittacidae), chachalacas, guans and curassows (Cracidae) and thrushes (Turdidae), which accounted for around 40% of all fruit-eating bird species observed ingesting entire palm fruits (Appendix S4). Mammals such as squirrels (Sciuridae), spiny rats (Cricetidae), primates (Atelidae and Cebidae) and carnivores (Canidae) accounted for 45% of all mammal species ingesting entire palm fruits, but ungulates (Tapiridae, Tayassuidae) and other carnivores (Procyonidae) were also important fruit-eaters (Appendix S4). The small number of fish and reptile species were also reported to feed on entire palm fruits (Appendix S4).

(2) Pulp-eaters
Pulp-eaters were represented by 32 bird species and 32 mammal species (Appendix S4), which were recorded to feed on the pulp of 27 palm species with a total of 115 interaction records. The two families with the highest number of pulp-eater species were bird families, namely tanagers (Thraupidae) and parrots (Psittacidae). The other pulp-feeding birds belonged to eight other families (Appendix S4). Among mammals, primates, marsupials and rodents were among the main mammalian palm-feeders of palms, including monkeys (Cebidae, Atelidae), opossums (Didelphidae), squirrels (Sciuridae), agoutis and acouchis (Dasyproctidae), and small rodents (Cricetidae). Each of these families was represented by 3–6 species (Appendix S4).

(3) Seed-eaters
A total of 73 animal species were recorded to feed on the seeds of 63 palm species (Appendix S4), with a total of 383 interaction records. This type of feeding interaction was largely dominated by rodent species (N = 62 spp.), which were involved in almost 95% of all seed-eating interaction records (Appendix S4). Other mammalian seed-eaters included primates and ungulates (Appendix S4). All five bird species involved in seed-eating interactions with palm seeds were parrots and macaws (Psittacidae) (Appendix S4).

V. INTERACTION OUTCOMES IN PALM–FRUGIVORE INTERACTIONS
We classified interaction outcomes as positive, negative, or dual for 73% of the available fruit-eating records, 76% of the pulp-eating records, and 97% of the seed-eating records (Table 1). The amount and frequency of interaction outcomes varied according to the feeding interaction type (Table 1), with positive outcomes being the most prevalent among fruit- and pulp-eaters and dual outcomes being the most prevalent outcome among seed-eaters. The interaction outcomes reflected a variety of effects on different stages during the seed-to-seedling transition of palms, including seed viability, seed deposition, and seedling establishment (Fig. 4). In the following, we synthesise for (i) fruit-eating, (ii) pulp-eating, and (iii) seed-eating interactions.
pulp-eating and (iii) seed-eating how interaction outcomes vary taxonomically and how they may influence plant demographic processes during the seed-to-seedling transition via effects on seed viability, seed deposition, and seedling establishment (Fig. 4). This includes, for instance, positive and negative effects of (i) fruit mastication, gut treatment, and...
endozoochorous seed dispersal of fruit-eaters, (ii) pulp removal and ectozoochorous seed dispersal of pulp-eaters, and (iii) ectozoochorous seed dispersal and seed caching of seed-eaters (Fig. 4).

(1) Fruit-eating

A total of 398 fruit-eating interaction records had information on interaction outcomes, capturing 251 unique interactions between 125 frugivore and 57 palm species. Almost 80% of the fruit-eating interaction records resulted in positive outcomes (Table 1), mainly involving birds and mammals (Fig. 5A). Negative and dual outcomes of fruit-eating interaction records were dominated by mammals (Fig. 5A). The type of digestive processing of fruits by animals (defecation, regurgitation, or both) was identified for 211 interaction records, involving 75 animal species and 45 palms (Draxler & Kissling, 2021). The main digestive process was defecation (N = 141, mostly mammals) and to a lesser extent regurgitation (N = 53, mostly birds), and sometimes both (N = 17, mostly birds) (Appendix S5). Most cases of fruit-eating for which the digestive process could be identified resulted in positive outcomes, regardless of how seeds were expelled (Fig. 6; Appendix S5).

(a) Fruit treatment in the beak or mouth

Many fruit-eaters ingest palm fruits without damaging the seeds, in part because palm endocarps offer a physical protection to seeds against mechanical and chemical damage, but also because many animal species do not mandibulate or masticate fruits in the mouth or beak, swallowing fruits whole. Fruit-eating without fruit processing in the oral cavity is seen, for example, in gulpers, that is birds that ingest whole fruits without processing in the bill (Levey, 1987). Toucans and toucanets (Ramphastidae) are good examples of gulpers and also important consumers of palm fruits, feeding on entire fruits of at least 11 palm species in interactions that...
are reported to result in positive outcomes. However, fruit-eaters can damage palm seeds in their mouth or beak when mandibulating or masticating fruits before swallowing them. The small number of negative outcomes of fruit-eating reported here are caused by seed damage derived from fruit mastication mainly by turtles (Chelidae), peccaries (Tayassuidae), tapirs (Tapiridae) (Zorz, 2009; Hibert et al., 2011) deer (Cervidae) and agoutis (Dasyproctidae) (Beck, 2006; Bodmer & Ward, 2006; Caputo & Vogt, 2008), but fruit-eating by fish and birds can also involve fruit mastication and lead to seed predation (Moegenburg & Levey, 2003; Piedade, Parolin & Junk, 2006; Correa et al., 2007).

Such damage during mastication is often caused when palm seeds have a relatively thin and fragile endocarp, such as those of *Euterpe* palms. However, fruit processing in the mouth or beak can also lead to damage of palm seeds that have stony endocarps, mainly such as those in the palm genera *Astrocaryum* and *Attalea* (Beck, 2006; Bodmer & Ward, 2006). Negative effects on seed viability due to mastication are further caused by the consumption of immature fruits, as reported for carnivores (Kays, 1999), rodents (Smythe, 1989; Mendes & Cândido-Jr, 2014; Acevedo-Quintero, Zamora-Abrego & Ortega-León, 2018) and primates (Smythe, 1989; Bowler & Bodmer, 2011).

(b) Gut treatment

After fruits are swallowed, they pass through the gut, and fruit-eaters either defecate or regurgitate the seeds. This may influence seed viability because fruit processing in the gut involves a series of chemical and mechanical actions that can affect the structure of the seed coat. This influence can be positive when fruit processing in the gut favours germination, for example due to the breakage of seed dormancy and facilitation of germination, or negative when seeds are destroyed as a result of harsh or long gut treatment (Torres, Castaño & Carranza-Quiceno, 2020). Whether seeds benefit from gut treatment or not also depends on specific traits of seeds such as seed size and hardness (Verdú & Traveset, 2004; Fuzessy et al., 2016; Fricke et al., 2019). Examples of regurgitation of viable palm seeds after fruit-eating (i.e. positive effects of gut treatment) include interactions between palm species and bird species of families such as Ramphastidae, Cotingidae and Cracidae (Pizo & Simão, 2001; Moegenburg & Levey, 2003; Sezen, Chazdon & Holsinger, 2009; Karubian et al., 2010, 2016; Campos, Steiner & Zilliéns, 2012; Ottewell et al., 2018). These regurgitated palm seeds have often a high germination success. Viable palm seeds have also been reported inside stomachs or in the dung of ungulates (e.g. Beck, 2006; Bodmer & Ward, 2006; Barcelos et al., 2013), carnivores (Kays, 1999; Campos et al., 2012), birds (Pizo & Simão, 2001; Cárdenas, Echeverry-Galvis & Stevenson, 2021), primates (Stevenson & Link, 2010; Chaves, Stoner & Arroyo-Rodriguez, 2012), turtles (Liu, Platt & Börg, 2004; Caputo & Vogt, 2008) and fish (Sório, Damasceno-Junior & Parolin, 2015; Barbosa & Montag, 2017). Our review shows that seeds of more than 30 palm species have been reported to benefit from gut treatment that involves seed defecation. In general, most palm seeds involved in fruit-eating seem to benefit from fruit processing in the gut of animals, regardless of whether they are defecated or regurgitated (Fig. 6; Appendix S5). Previous reviews have also shown that gut treatment of seeds has an overall positive effect on seed viability (Traveset & Verdú, 2002; Fuzessy et al., 2016; Torres et al., 2020), independent of whether seeds only pass through part of the animal gut (when regurgitated) or through the whole digestive system (when defecated). The results show that Neotropical palm seeds can have high levels of seed protection and thus are well adapted to fruit processing by fruit-eaters. Nevertheless, more studies addressing the effects of gut treatment on the viability of
palm seeds are needed, especially for some understudied fruit-eaters such as fish and reptiles [but see Correa et al. (2015) and Falcón, Moll & Hansen (2020) for reviews of frugivory by these animal groups].

(c) Long-distance seed dispersal by endozoochory

Fruit-eating always involves endozoochory (i.e. seed dispersal via ingestion). Subsequent dispersal distances can vary widely and depend on the mobility, space use, body size, gut retention time and digestive fruit-processing type of seed dispersers (Traveset & Verdú, 2002; Côrtes & Uriarte, 2013). When fruit-eaters move far after fruit consumption, long-distance dispersal events are facilitated, often with distances >100 m or even >1 km (Fig. 7; Appendix S6). This contributes to increasing the genetic diversity of palm populations (Sezen et al., 2009; Browne, Otewell & Karubian, 2015; Browne & Karubian, 2018). Long-distance seed dispersal by frugivores that eliminate viable palm seeds has been suggested for carnivores (Gatti et al., 2006; Silva et al., 2011; Quintela, Iob & Artioli, 2014; Menezes et al., 2018), fish (Anderson et al., 2011) and primates (Link & De Luna, 2004; Link & Di Fiore, 2006; Canale et al., 2016). Large bird species (e.g. Cotingidae, Cracidae, Ramphastidae, Steatornithidae) can deposit seeds far away from parent palms (Appendix S6), showing that avian fruit-eaters can have an ecological function similar to that of large-bodied mammalian fruit-eaters (Stevenson et al., 2021). For instance, long-wattled umbrellabirds (Cephalopterus penduliger) show high gut retention times (up to 105 min for seeds of the palm Oenocarpus bataua; Karubian et al., 2012) and have been shown to disperse intact seeds effectively (Karubian et al., 2010) over hundreds of meters away from fruit sources (Karubian et al., 2012, 2016). In rare cases like the oilbird (Steatornis caripensis), seed dispersal (e.g. of the palm Oenocarpus bataua) can even exceed 40 km (Stevenson et al., 2021).

(d) Seed deposition and seedling establishment

Interaction outcomes of fruit-eating are influenced by the microsites at which seeds are deposited. Seed deposition sites can be highly variable even when interactions involve the same frugivore species. For instance, latrines of the lowland tapir (Tapirus terrestris) are often located at upland forest sites, which are considered suitable sites for the germination of Attalea maripa seeds, but tapirs can also use latrines located in flooded areas, which can be disadvantageous for seed germination (Fragoso & Huffman, 2000; Quiroga-Castro & Roldán, 2001). Such intra-specific variability in the influence of fruit-eaters on the deposition of palm seeds can reduce the proportion of positive outcomes derived by palms from interactions with a specific fruit-eater. By contrast, germination and seedling establishment of palms is facilitated when fruit-eating animal species consistently deposit seeds at particular microsites with optimal conditions (i.e. directed seed dispersal, such as dispersal to a certain habitat type, at

![Fig. 7. Dispersal distances of fruit-eaters, pulp-eaters and seed-eaters involved in in Neotropical palm–frugivore interactions.](image-url)

(A) Distribution of mean dispersal distances (logarithmic axis) for fruit- (N = 3 records), pulp- (N = 25 records) and seed-eating (N = 34 records), involving a total of 26 frugivore species and 22 palm species. (B) Distribution of maximum dispersal distances (logarithmic axis) recorded for fruit- (N = 4 records), pulp- (N = 2 records) and seed-eating (N = 25 records), involving a total of 13 frugivore species and 18 palm species. Data on dispersal distances compiled from published articles (see Appendix S6).
preferred feeding trees, or specific breeding or sleeping sites) (Karubian et al., 2012, 2015; Canale et al., 2016). For example, long-wattled umbrellabirds (*Cephalopterus penduliger*) disperse a high proportion of the seeds of *Oenocarpus bataua* within breeding sites known as leks (Karubian et al., 2012), many of which are found to germinate and establish into seedlings (Karubian et al., 2016). However, this is not the case when palm seeds are dispersed to non-suitable sites, such as seeds that are deposited in caves by the oilbird *Steatornis caripensis* (Stevenson et al., 2017).

When viable palm seeds are deposited at sites that are suitable for germination, fruit-eaters facilitate the establishment of seedlings. Such positive associations between fruit-eating and seedling establishment of palms have been found for species of birds (Pizo & Simão, 2001; Sezen et al., 2009; Karubian et al., 2012), mammals (Fragoso, 1997; Fragoso & Huffman, 2000; Sica, Bravo & Giombini, 2014) and fish (Barbosa & Montag, 2017), involving palm species such as *Attalea maripa*, *Euterpe edulis*, *Inria reticulata*, *Oenocarpus bataua* and *Syagrus romanzoffiana*. For instance, Fragoso, Silvius & Correa (2003) showed that the aggregation of viable *Attalea maripa* seeds in tapir latrines favors local seedling establishment and contributes to generating patches of *A. maripa* adult individuals, an association that was also observed for the Atlantic Forest palm *Syagrus romanzoffiana* (Sica et al., 2014).

(2) **Pulp-eating**

For pulp-eaters, interaction outcomes could be identified for 87 out of 115 interaction records, including 29 mammal species and 15 bird species recorded to interact with 21 and 11 palm species, respectively. Pulp-eating records resulted mostly in positive interaction outcomes (Table 1).

Most mammalian species recorded in pulp-eating interactions with classified outcomes have a high fruit-handling ability (*N* = 15 spp.), but some also showed an intermediate (*N* = 7 spp.) or low (*N* = 7 spp.) ability to handle fruits while consuming the pulp (Dracxler & Kissling, 2021) (Appendix S7A). By contrast, bird species involved in pulp-eating showed low (*N* = 9 spp.) or intermediate (*N* = 6 spp.) ability to handle fruits (Dracxler & Kissling, 2021). Around half (44%) of all pulp-eating records involved animals with low handling ability, predominantly mammals (Appendix S7A). This suggests that pulp-eating might be a feeding strategy that predominantly occurs among animals that cannot skillfully manipulate palm fruits. Positive outcomes generally dominated, regardless of the level of fruit-handling ability (Fig. 6; Appendix S7B), suggesting that most pulp-eaters of palm fruits are mutualists.

(a) **Pulp removal**

The way pulp is removed by animals depends on how animals can handle fruits. Pulp-eating usually involves pulp extraction in the mouth and subsequent spitting (e.g. ungulates), fruit manipulation with the help of hands or forepaws and the mouth or beak (e.g. primates and macaws), or pulp-pecking by animals that cannot ingest entire fruits or manipulate them otherwise. The removal of palm mesocarps by pulp-eaters can negatively affect seed viability when animals (e.g. parrots and peccaries) damage seeds during fruit handling and pulp extraction (Keuroghlian & Eaton, 2009; Villalobos & Bagno, 2012). Our review suggests that, in most cases, pulp-eaters consume the pulp and discard palm seeds intact, a process that benefits seed survival and germination because the pulp often has chemical inhibitors that prevent seeds from germinating (Robertson et al., 2006). Effects of pulp-eating on seed viability and germination also depend on the amount of pulp that is removed, because remaining pulp can favor attacks by pathogens and invertebrate predators or lead to a faster deterioration of seeds (Herrera, 1982). Among pulp-eaters of palms, partial pulp removal is mainly observed with animals that have a low handling ability, including birds such as tanagers and blackbirds (e.g. Matos & Watkinson, 1998; Villalobos & Bagno, 2012) or mammals such as peccaries (Fragoso, 1997). Pulp-eaters with intermediate or high handling ability (e.g. macaws or primates) generally remove most of the pulp and then discard the well-cleaned palm seeds (e.g. Fragoso, 1997; Villalobos & Bagno, 2012; Franco-Quimbay & Rojas-Robles, 2015; Filho, Andrade & Bezerra, 2021). This often leads to high palm seed survival and germination, and thus a positive interaction outcome. Yet, cleaned palm seeds that are discarded by pulp-eaters can be subsequently eaten by other animals, as is the case for the exotic palm *Elaeis guineensis* in the Atlantic Forest in northeastern Brazil, where the red-back agouti *Dasyprocta iacki* consumes seeds left over after pulp-eating by the blonde capuchin *Sapajus flavius* (Filho et al., 2021). This highlights the importance of considering secondary fates of seeds discarded by pulp-eaters.

(b) **Seed dispersal by ectozoochory**

Seeds discarded after pulp consumption are often found below or near parent palms (Kays, 1999; Campos, Steiner & Zillikens, 2012; Franco-Quimbay & Rojas-Robles, 2015). However, some mammalian pulp-feeders with high mobility and a high ability to handle fruits (e.g. primates, rodents, marsupials, cingulates and ungulates) transport palm fruits to other sites or feeding trees, away from parent palms, where they feed on the pulp and then discard the viable and cleaned palm seeds (e.g. Cintra & Horna, 1997; Fragoso et al., 2003; Pimentel & Tabarelli, 2004; Beck, 2006; Keuroghlian & Eaton, 2009; Brown, 2011; Silva et al., 2011; Acevedo-Quintero & Zamora-Abrego, 2016; Canale et al., 2016; Junges et al., 2018). The dispersal distances are typically rather short (e.g. <50 m), but some large birds such as macaws and parrots (Psittacidae) can transport seeds over long (>200 m) distances (Sazima, 2008; Villalobos & Bagno, 2012; Santos & Ragusa-Netto, 2014; Baños-Villalba et al., 2017; Blanco et al., 2019) (Fig. 7; Appendix S6). These defleshed seeds...
(i.e., the endocarps) are left on the ground surface (without burial, unlike hoarding seed-eaters) and do not have protective excreta around the seeds (as provided by fruit-eaters) (Ríos & Pacheco, 2006). Nevertheless, they benefit from pulp removal and dispersal, and seeds can successfully germinate and establish as seedlings (e.g., Campos et al., 2012). This might be particularly beneficial if the abiotic conditions at the deposition sites (e.g., humidity or luminosity) match the germination requirements of the seeds, or if desiccation associated with seed exposure facilitates the break of dormancy, germination and establishment (Broschat, 1998). Hence, pulp-eaters such as macaws can act as legitimate, long-distance seed dispersers of palms and thus play an important role in connecting palm populations between habitat fragments (Baños-Villalba et al., 2017).

(3) Seed-eating
A total of 370 seed-eating records contained information on interaction outcomes, covering 186 unique interactions involving 70 animal species feeding on the seeds of 62 palm species. More than 60% of these seed-eating records (N = 222) showed dual outcomes and almost 30% resulted in negative outcomes (Fig. 5C; Table 1). Some seed-eating interactions had positive outcomes (N = 41). Interaction outcomes of seed-eaters were dominated by mammals, mainly involving agoutis and squirrels (Appendix S4).

Most species of seed-eaters had an intermediate ability to manipulate fruits, including 32 rodent species and five parrots (Dracxler & Kissling, 2021). High handling ability was observed for 27 rodent species and two primate species, while a low handling ability was identified for only four species of rodents and two peccary species. The largest proportion of interaction records involved mammalian seed-eaters with high handling ability (Appendix S8A), partly due to the large number of interaction records involving agoutis (Dasyprocta spp.). Seed-eating interactions mostly resulted in dual outcomes regardless of the handling ability of animals (Appendix S8B).

For 61 out of the 70 seed-eater species (56 mammals and 5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absence of caching animals included birds (5 bird species) we could classify the presence or absent...
However, seed predation rates can increase when food is limited, increasing negative outcomes. This has been documented, for example, for the palm *Astrocaryum aculeatum* and rodents in the Brazilian Amazon (Jorge & Howe, 2009).

Large and long-lasting seeds are cached more often compared to small seeds with thin endocarps because hoarding animals perceive them as the most valuable food items (Jansen *et al*., 2002; Galetti *et al*., 2010; Mittelman *et al*., 2021). This suggests that interactions involving large, stony palm seeds are more likely to result in positive outcomes because those seeds are preferentially cached instead of immediately consumed. Seed caching can also indirectly favour seed viability by facilitating escape from seed predation by invertebrates such as bruchid beetles (Smythe, 1989; Kuprewicz, 2015). This happens because rodents reduce chances of seed encounter by bruchids, which tend to infest seeds near parent palms. They further exert a top-down control over bruchid populations by consuming bruchid larvae (i.e., grubivory, *sensu Silva*, 2002) or removing oviposited larvae during seed cleaning (Peguero *et al*., 2017). This secondary benefit of interactions with seed-caching animals can be crucial for some palm species that would otherwise experience high invertebrate predation if seeds were left near parent palms (Dracxler, Pires & Fernandez, 2011).

Seed caching usually involves seed dispersal because animals tend to move and cache seeds away from parent palms (Smythe, 1989; Lichti *et al*., 2017). Dispersal distances of palm seeds depend on a combination of factors such as frugivore traits (e.g., body size, movement patterns and home ranges of seed-caching animals (Carvajal & Adler, 2008; Lichti *et al*., 2017)], fruit availability (Gávez *et al*., 2009) and palm traits such as seed mass (Galetti *et al*., 2010). The Central-American agouti *Dasyprocta punctata* can move seeds of *Astrocaryum standleyanum* hundreds of meters away from parent palms because cache owners and thieves re-cache the seeds multiple times (Jansen *et al*., 2012). Nevertheless, most reports of caching-associated seed dispersal suggest that scatter-hoarders tend to disperse seeds to distances <50 m (Forget & Wenny, 2005) (Fig. 7; Appendix S6). For palm species that are exclusively dispersed by scatter-hoarding rodents, dispersal distances associated with seed caching can strongly limit how far a population can expand (Charles-Dominique *et al*., 2003). However, commonly used methods to track seeds are limited in how far dispersal distances can be measured (Forget & Wenny, 2005; Mittelman *et al*., 2021).

Seed-caching rodents may select different microsites to deposit the seeds (Hoch & Adler, 1997). For instance, scatter-hoarding animals often use spatial cues to recover their caches. They cache seeds near large objects such as fallen trunks (Kiltie, 1981; Smythe, 1989; Hoch & Adler, 1997; Pires & Galetti, 2012), move seeds to forest gaps (Carvajal & Adler, 2008), or deposit them in areas with low density of conspecific adults (Brewer & Webb, 2001; Hirsch *et al*., 2012b). Importantly, deposition microsites of seed-caching animals are not always beneficial for the palms. Squirrels, for example, can place seeds in arboreal caches, a microsite that might not be favourable for the development of palms (Carvajal & Adler, 2008). Nevertheless, seed caching is mostly associated with positive outcomes at the seedling establishment stage because caching behaviour involves palm seed cleaning, storage and burial at depths that favour seed germination and seedling emergence (Vander Wall, 1990).

VI. MUTUALISM–ANTAGONISM CONTINUA: QUANTIFYING INTRA- AND INTER-SPECIFIC VARIATION

Section V documented how outcomes in palm–frugivore interactions can vary among fruit-eaters, pulp-eaters, and seed-eaters and according to their consumption behaviours and traits. In this section we synthesise available evidence on where specific frugivore families and species fall along the mutualism–antagonism continuum and how this position varies along the continuum (Fig. 8). This was done by summarizing variation in interaction outcomes from the best qualitative and quantitative studies available for palms, focusing on the proportion of positive outcomes within individual frugivore species (intra-specific variability) and across different frugivore species within families (inter-specific variability).

(1) Qualitative data

A total of 17 frugivore families had ≥7 studies with qualitative data available at the family level to summarize the proportion of positive outcomes along the mutualism–antagonism continuum (Fig. 8A). These families were represented by a total of 107 animal species recorded in 495 interaction records, and included rodents (Dasyproctidae, Echimyidae, Sciuridae and Cricetidae), ungulates (Tapiroidea, Tayassuidea and Cervidae), primates (Atelidae and Cebidae), carnivores (Canidae and Procyonidae), marsupials (Didelphidae) and small (Turdidae) and large birds (Ramphastidae, Cracidae, Cotingidae and Psittacidae). Twelve frugivore families (75%) were classified as mainly mutualistic along the continuum (Fig. 8A), indicating that the majority of species in these families (95%, N = 74) provided positive outcomes for palms. Most of the interactions with these animals provided benefits for palms through increased seed survival and seedling establishment after seed treatment. This included even families such as carnivores (Canidae and Procyonidae), parrots (Psittacidae) and rodents (Dasyproctidae) that are rarely perceived as mutualists. In each of two families (Psittacidae, Dasyproctidae) that were classified as mutualistic, two species fall on the antagonistic side of the continuum: two parrots (*Pyrrhura frontalis* and *Triclaria malachitacea*) and two agoutis (*Dasyprocta leporina* and *D. iacki*). This highlights the large inter-specific variability within parrots and rodents. Other mammalian families with mostly positive outcomes for palms included marsupials
(Didelphidae), primates (Atelidae and Cebidae) and ungulates (Tapiridae). Four bird families ranked as highly mutualistic, including toucans (Ramphastidae), thrushes (Turdidae), chachalacas, guans and curassows (Cracidae) and cotingas (Cotingidae). A total of 33 bird species were represented by these four bird families, with a total of 117 interaction outcomes represented in the continuum, all of which involved fruit-eating and positive outcomes. Canidae and Didelphidae were also classified as fully mutualistic and represented by 11 animal species.

Only five frugivore families (including spiny rats, peccaries, deer and squirrels) fell on the antagonistic side of the continuum, with the average proportion of positive outcomes ranging from 0.17 to 0.36 (Fig. 8A). However, even those families were not exclusively antagonistic because interaction outcomes reported in the literature also indicated positive interaction outcomes for some species. Negative outcomes were mainly explained by the predation of seeds through mastication in fruit-eating events or by direct consumption (and destruction) of seeds in seed-eating interactions.

Fig. 8. The mutualism–antagonism continuum in Neotropical palm–frugivore interactions. The proportion of positive interaction outcomes from qualitative and quantitative evidence reported in the literature is summarized for frugivore families and species along the mutualism–antagonism continuum (x-axis). The left side of the continuum mostly shows antagonistic families or species (total or mean values in red; proportion of positive outcomes <0.5) and the right side mostly mutualistic species or families (total or mean values in blue; proportion of positive outcomes ≥0.5). Grey dots represent the proportion of positive outcomes for species within families (A) or individual studies within species (C), with grey horizontal lines indicating the range. (A) Qualitative family-level data summarizing the proportion of positive outcomes across species within 17 frugivore families from 124 papers, with a total of 495 interaction records. The number of species (2–12) and the number of interaction records (mean ± SD: 29.1 ± 23.3, range: 7–83 outcome records) varies per family. (B) Qualitative species-level data showing the average proportion of positive outcomes for 34 frugivore species with ≥5 outcome observations (mean ± SD: 11.9 ± 11.8, range: 5–53 outcome records) from 111 papers, with a total of 406 interaction records. (C) Quantitative species-level data for three species of frugivores (the lowland tapir *Tapirus terrestris*, the red-rumped agouti *Dasyprocta leporina*, and the hyacinth macaw *Anodorhynchus hyacinthinus*) summarizing evidence on the proportion of positive outcomes (five records per species) from six scientific studies. Outcome data are provided in a supplementary file containing interaction records and outcomes extracted from the sources (Continuum data; Dracxler & Kissling, 2021).
A total of 34 frugivore species had ≥5 studies available with qualitative data available at the species level to summarize the proportion of positive interaction outcomes along the mutualism–antagonism continuum (Fig. 8B). Twenty-seven species (79%) were predominantly mutualists (with proportion of positive outcomes ≥0.5; Fig. 8B). Sixteen of those were fully mutualistic (proportion of positive outcomes = 1), including two toucans (Ramphastos vitellinus and R. dicolorus), the oilbird (Steatornis caripensis), two toucanets (Pteroglossus baiioli and Selenidera maculirostris), five thrush species (Turdus albicollicollis, T. flavipes, T. leucomelas, T. rufiventris and T. amaurochalinus), the dusky-legged guan (Penelope obscura), the long-wattled umbrellabird (Cephalopterus penduliger), the greater rhea (Rhea americana), the crab-eating fox (Cerdocyon thous), the domestic cow (Bos taurus) and the ring-tailed coati (Nasua nasua) (Fig. 8B). The eleven other frugivore species placed on the mutualistic side of the continuum (including two tapirs, two macaws, two monkeys, two agoutis, a paca, a squirrel and a spiny rat) were mutualistic because of their positive role as fruit-eaters, besides dispersing seeds by ectoozoochory, feeding on pulp and discarding intact seeds, and caching seeds. Seven species (21%) were classified as mostly antagonists, including peccaries, squirrels, a spiny rat, an agouti and a turtle species (Fig. 8B). Antagonistic outcomes involving these animals were mainly derived from fruit- and seed-eating interactions, in which seeds are damaged due to fruit mastication or direct seed predation. However, the continuum reveals that none of these species are fully antagonistic because palms can also derive benefits from interactions with these animals. In general, aggregation of outcomes at the species level shows that most animal species fall along the mutualism–antagonism continuum and are neither obligate mutualists nor exclusively antagonists, that is that intra-specific variation of outcomes is common for most animals interacting with palm fruits.

(2) Quantitative data

A total of three frugivore species had ≥5 studies available with quantitative information to allow us to summarize trends and intra-specific variability in the proportion of positive outcomes along the mutualism–antagonism continuum (Fig. 8C). For the lowland tapir (Tapirus terrestris), quantitative data were available on seed viability and seedling establishment, representing the proportion of seeds found viable in dung and the germination success of seeds in dung piles, respectively (Rodrigues, Ollas & Galetti, 1993; Fragoso, 1997; Fragoso & Hulman, 2000; Giombini, Bravo & Martinez, 2009). For the hyacinth macaw (Anodorhynchus hyacinthinus), quantitative data reflected information on seed damage, either as the proportion of seeds found undamaged after seed dispersal or as the proportion of predated seeds (Tella et al., 2020). For the red-rumped agouti (Dasyprocta leporina), quantitative data on seed predation were available as the proportion of seeds predated versus cached (Galetti et al., 2010). Among these three species, the lowland tapir showed the highest proportion of positive outcomes (mean proportion of positive outcomes = 0.68; Fig. 8C) and was thus clearly a mutualistic frugivore. However, some evidence also indicated a potential antagonistic role in some cases (range of outcomes = 0.28–0.98; Fig. 8C). By contrast, the red-rumped agouti and the hyacinth macaw predominantly had an antagonistic role for palms (mean proportions of positive outcomes <0.5), although some studies indicated a mutualistic role (Fig. 8C).

The available quantitative evidence of the position and range of species along the mutualism–antagonism continuum shows that outcomes of interactions involving the same frugivore species can vary widely, even for taxa such as the tapir which are generally perceived as mutualistic frugivores. This might explain the lack of consensus in the literature about the ecological roles of some frugivores (Tella et al., 2015; Gómez et al., 2019; van Leeuwen, Tella & Green, 2020). The red-rumped agouti provides an interesting example of the variability in interaction outcomes because some studies report the predation of virtually all seeds of some palms (e.g. Euterpe edulis, Syagrus pseudococos and S. romanzoffiana) whereas another study shows a high positive outcome for another palm (66% of Astrocaryum aculeatissimum seeds are cached; Fig. 8C) (Galetti et al., 2010). Similarly, the hyacinth macaw typically predates the majority of seeds (resulting in a largely antagonistic role) but can also be a mutualist when it discards a large number of undamaged seeds of specific palms (e.g. Acrocomia totai, Attalea barreirensis and Attalea eichleri) below perching trees (Tella et al., 2020).

VII. PROSPECTS

To our knowledge, our review provides the most comprehensive evidence to date on the inter-specific and intra-specific variability in interaction outcomes of a particular plant–frugivore system. Despite the many studies on palm–frugivore interactions that we have reviewed here, evidence on the positive and negative effects of fruit-eating, pulp-eating and seed-eating on palm demography during the seed-to-seedling transition remains scarce. This is due to two main reasons. First, it is often not clear whether animal species recorded in frugivory interactions feed on entire fruits, only on the pulp or only on seeds. This lack of detail on frugivory interactions makes it difficult to gather evidence on the effects of different types of feeding interactions on seed-to-seedling transitions. Second, assessing quantitative and qualitative effects of interactions on seed viability, seed deposition, germination and seedling establishment is logistically difficult, time-consuming and methodologically constrained, and thus rarely conducted (Schupp et al., 2010; Beckman et al., 2020). Nevertheless, a range of field and laboratory methods are available to study and quantify interaction outcomes (Fig. 9).
Visual observations at focal trees allow identifying which species are involved in interactions and which type of feeding and fruit-handling ability they have (Fig. 9), for example, whether animals feed on fruits, pulp or seeds and how they handle the fruits and seeds. Repeating this for different types of plant species and different individuals of the same plant species will provide insights into the variability of feeding interaction types within and across frugivorous animal species. Visual observations can further provide quantitative information on fruit removal and interaction frequencies (Schupp et al., 2010; Meiga & Christianini, 2020). However, experimental approaches such as cafeteria plots (e.g. Jansen, Bongers & Hemerik, 2004) or exclosure experiments (Hulme, 1994) have an advantage over visual observations because they provide a semi-controlled setting to evaluate the number of fruits or seeds taken by animals. When used in combination with camera traps, such experiments are effective ways for evaluating species-specific visitation and fruit or seed removal rates, as well as in situ pulp consumption (Jansen et al., 2012; Acevedo-Quintero & Zamora-Abrego, 2016; Campos et al., 2018; Meiga & Christianini, 2020; Selwyn et al., 2020).

Following seed survival and seed fates after fruits are taken by frugivores (Fig. 9) is challenging because visual observations are limited in measuring seed viability and dispersal distances when seeds are deposited away from parent plants. Large-bodied fruit-eaters can move long distances before they deposit seeds (Nathan, 2006) and scatter-hoarding rodents may move seeds in multiple steps (Jansen et al., 2012), making it difficult to assess outcomes after fruit removal. Misrepresentations of interaction outcomes can emerge if fruit or seed removal is assumed (rather than measured) to result either in mutualisms or antagonisms. For instance, fruit-eating is often assumed to be equivalent to effective dispersal (mutualism) and seed or fruit removal by seed-eaters is assumed to be equivalent to seed predation (antagonism) (Gómez et al., 2019). However, such conclusions can only be drawn accurately through careful evaluation of individual seeds sampled after interactions have taken place. Measuring seed fate can be done by tracking individual seeds (Fig. 9), e.g. via marking seeds through stable isotope enrichment of fruiting trees (Carlo, Tewksbury & del Río, 2009) or by tagging seeds, e.g. with spool-and-line or telemetric thread tags (Forget & Wenny, 2005; Hirsch, Kays & Jansen, 2012a). These methods also provide:

**Fig. 9.** Examples of methods for assessing the effects of frugivores on seed-to-seedling transitions, separated by feeding interaction types. The suggested methods allow for assessing interaction aspects such as interaction frequency (e.g. number of visits), fruit removal and consumption behaviour of frugivores, and their effects on plant population dynamics (seed survival and seed fate) and demography (seed germination and seedling establishment). Blue, yellow and orange circles show methods applicable to fruit-eating (‘fruit’), pulp-eating (‘pulp’) and seed-eating (‘seed’), respectively.

(1) **Quantification of interaction outcome variation**

Visual observations of interaction outcomes are crucial for understanding the dynamics of plant-animal interactions. Visual observations can provide quantitative information on fruit removal and interaction frequencies, but they are limited in measuring seed viability and dispersal distances when seeds are deposited away from parent plants. Large-bodied fruit-eaters can move long distances before they deposit seeds, and scatter-hoarding rodents may move seeds in multiple steps, making it difficult to assess outcomes after fruit removal. Misrepresentations of interaction outcomes can emerge if fruit or seed removal is assumed to result in mutualisms or antagonisms. For instance, fruit-eating is often assumed to be equivalent to effective dispersal (mutualism) and seed or fruit removal by seed-eaters is assumed to be equivalent to seed predation (antagonism). However, such conclusions can only be drawn accurately through careful evaluation of individual seeds sampled after interactions have taken place. Measuring seed fate can be done by tracking individual seeds, e.g. via marking seeds through stable isotope enrichment of fruiting trees, or by tagging seeds, e.g. with spool-and-line or telemetric thread tags. These methods also provide...
information on dispersal distances (Forget & Wenny, 2005; Jansen et al., 2012), although fixed-tagging methods may reduce seed movement and constrain the evaluation of dispersal distances. Applying DNA-barcoding protocols to identify disperser species from samples of regurgitated or defecated seeds is also a powerful method which allows linking disperser identity to fruit trait choices and seed deposition patterns (González-Varo, Arroyo & Jordano, 2014). Assessing whether individual retrieved seeds (e.g. in caches, on the forest floor, or in animal dung) are intact after gut treatment by fruit-eaters, after pulp removal by pulp-eaters or if seed-eaters ultimately leave seeds undamaged is necessary to inform the effects of frugivores on seed survival. This might also require manually opening endocarps or using X-rays to assess endosperm integrity because apparently intact endocarps can actually be infested by bruchid larvae (Brancalion et al., 2011; Dracxler et al., 2011). Controlled assessments of the effects of mouth or beak and gut treatment on seeds and gut retention times (Fig. 9) – including captive frugivore feeding trials combined with germination experiments – are also useful for estimating the effects of fruit-eaters on seed viability (Samuels & Levey, 2005; Fricke et al., 2019), and for quantifying seed dispersal kernels (Schurr et al., 2009; Pires et al., 2018; Sorensen et al., 2020).

A number of methods exist to assess how germination and seedling establishment is affected by the seed treatment of frugivores (Fig. 9) (Snell et al., 2019). Experimental approaches in the laboratory or in the field can evaluate germination success of gut-passed, handled (e.g. defleshed) or cached seeds (Samuels & Levey, 2005; Kuprewicz, 2015), and when conducted at mid- or long-term, can also show effects of frugivores on seedling survival and transitions into later ontogenetic stages (e.g. the sapling stage). Field observations (e.g. mapping or monitoring of seedlings) can further inform about the net effect of frugivores on seed-to-seedling transitions (e.g. seedlings established from seeds found in dung or buried in caches) (Fragoso, 1997; Beckman & Rogers, 2013; Sica et al., 2014; Dracxler & Forget, 2017). Finally, an approach that combines a spatial assessment of the distribution of seedlings and adults (e.g. source trees) with molecular techniques such as genetic parentage analysis (Sezen et al., 2009; Choo, Juenger & Simpson, 2012; Giombini, Bravo & Tosto, 2016; Giombini et al., 2017; Browne & Karubian, 2018; Diaz-Martin & Karubian, 2021) is a way to shed light on how frugivory influences the spatial and genetic structure of palm populations. The non-exhaustive list of methods provided here (Fig. 9) illustrates exciting opportunities for deepening our knowledge about the effects of frugivores on interaction outcomes and plant demography.

(2) Influence of fruit traits on feeding types and interaction outcomes

Variation of interaction outcomes within types of feeding interactions might partly be driven by variation in palm fruit and seed traits (Fig. 10). For example, fruits with stony endocarps are found in palm genera such as *Astrocaryum*, *Attalea* and *Syagrus*, sometimes with needle-like spines on the exocarp, a fibrous mesocarp, and a lipid-rich endosperm (Fig. 10C). These are mainly consumed by scatter-hoarding rodents (i.e. seed-eaters), but because these palm seeds are ideal for storage (i.e. long-lasting, large nutrient-rich endocarps), the interactions often result in mutualisms rather than antagonisms due to high caching rates (Jorge & Howe, 2009; Galetti et al., 2010; Klinger & Rejmánek, 2010; Jansen et al., 2012; Dracxler & Forget, 2017). On the other hand, when rodents feed on small fruits that have rather thin endocarps – such as the fruits of *Euterpe edulis* which have a fleshy-fibrous pulp and are mainly consumed by birds (Fig. 10A) – interactions with rodents may predominantly result in seed predation rather than seed caching (Pizo, Von Allmen & Morellato, 2006; Galetti et al., 2013). Similarly, seeds enclosed in thick endocarps are more likely to survive mouth, beak and gut treatment of fruit-eaters compared to fruits with thin seed coats (Bodmer & Ward, 2006). The type of pulp in palm fruits (e.g. farinaceous, fibrous or fleshy) might also interfere with how easily seeds are defleshed by pulp-eaters (Fig. 10B), in turn influencing whether seeds are left undamaged after pulp removal. Future studies should also focus on understanding how palm traits might influence associations with different groups of animals and/or feeding guilds and whether the emergence of positive and negative outcomes can be predicted by palm traits. A workable hypothesis is that fruits sharing specific combinations of traits (e.g. red- and purple-coloured fruits, < 2 cm in diameter, and with fleshy pulp such as in *Euterpe*, *Geonoma* and *Oenocarpus*; Kissling et al., 2019) that have evolved in response to selection by certain groups of frugivores (e.g. large-bodied birds) are more likely to benefit from interactions with such groups of frugivores than with other types of frugivores (e.g. rodents). Combining different plant traits (Gautier-Hion et al., 1985; Nascimento et al., 2020) with data on trait-matching relationships (Bender et al., 2018) and interaction outcomes (as reviewed here) may provide novel insights into how mutualisms and antagonisms in plant–frugivore interactions have evolved (Guimarães Jr, 2020) and how they could be used to project future changes in communities of interacting species (Schleuning et al., 2020).

(3) Influence of frugivory on plant and ecosystem dynamics

In recent years, studies have helped to consolidate functional and trait-based approaches that link frugivore species to ecosystem dynamics and ecosystem functioning (Schleuning, Frield & Garcia, 2015). Emerging evidence supports a positive effect of large-bodied frugivores on carbon storage in tropical forests and thus emphasizes an important role of fruit-eaters in biogeochemical cycles (Bello et al., 2015). Animals such as ungulates and rodents further may affect other biogeochemical cycles such as the nitrogen cycle (Villar et al., 2020). To date, the importance of seed- and pulp-eaters for biogeochemical cycles in tropical ecosystems remains little studied, although it is known

---

*Biological Reviews* 97 (2022) 527–553 © 2021 The Authors. *Biological Reviews* published by John Wiley & Sons Ltd on behalf of Cambridge Philosophical Society.
that seed-caching animals tend to cache large seeds and predate small ones, thus selecting for palm species that are likely to store more carbon (Mittelman et al., 2021). Scatter-hoarding rodents such as agoutis (*Dasyprocta* spp.) can disperse seeds larger than those dispersed by megaherbivores such as tapirs and large-bodied monkeys (*Tapirus* spp., *Brachyteles* spp. and *Ateles* spp.) (Mittelman et al., 2021). Particular seed-eaters might thus contribute more to carbon storage than the remaining extant Neotropical megaherbivores. The type of feeding interaction can influence the animal’s contribution to carbon storage if fruit and seed size are positively correlated with carbon storage capacity of trees (Fig. 11). Fruit-eating is often characterized by size matching, that is large-bodied fruit-eaters disperse both large and small palm seeds whereas small-bodied fruit-eaters can only disperse small palm seeds. However, pulp- and seed-eaters are able to disperse seeds that are larger than their mouth or beak because they can transport seeds externally (either by ectozoochory or synzoochory) (Fig. 11). This suggests that animal species which are typically considered as non-mutualistic can actually contribute to palm population dynamics and ecosystem functions such as carbon storage. This contribution, however, depends on the interaction outcomes with frugivores, emphasizing the need to assess intra-specific variation of the effects of frugivores on the several stages of the plant life cycle (Fig. 9).

Assessing how disturbances and novel communities can influence interaction outcomes of palm–frugivore interactions is of fundamental importance for predicting the consequences of habitat loss and modification for palm population dynamics (Jorge & Howe, 2009; Klinger &
Rejmánek, 2010; Schupp et al., 2019; Escobar et al., 2020; Lamperty, Karubian & Dunham, 2021). Habitat and climate change can also alter palm phenology (e.g. Andreazzi Lamperty, Karubian & Dunham, 2021). Habitat and climate change can have cascading effects on palm regeneration. Quantifying how seed dispersal distances vary according to palm fruit traits and feeding guilds of frugivores is also crucial for forecasting palm species distributions under climate change (Butler & Larson, 2020; Sales et al., 2021). Pulp-eaters and seed-eaters may not be able to disperse palm seeds as far as fruit-eaters can, but certain frugivores in these feeding guilds provide highly relevant long-distance dispersal services for several palms (Fig. 7). Ongoing defaunation might mean that seed- and pulp-eaters can take on the role as main dispersers of some large-fruited palm species (e.g. Meiga & Christianini, 2020). Since palms are particularly abundant in the Neotropics (ter Steege et al., 2013; Muscarella et al., 2020), understanding the interactions with different feeding guilds and projecting the consequences for palm demography and ecosystem dynamics and functioning should be of high priority.

VIII. CONCLUSIONS

(1) Frugivory involves both advantages and disadvantages for animal-dispersed plants. Because the effects of frugivory on the seed-to-seedling transition in plants vary both within and among animal species, assessing intra- and inter-specific variation of interaction outcomes along a mutualism–antagonism continuum is essential for understanding the consequences of frugivory for plant demography and population and ecosystem dynamics.

(2) Frugivory can involve fruit-eating, pulp-eating and seed-eating. Each of these types of feeding interactions can have different influences on seed viability, seed deposition and seedling establishment of plants. Based on a comprehensive compilation of 1043 Neotropical palm–frugivore interaction records that capture variation of feeding interaction types – covering 273 species of birds, mammals, fish and reptiles feeding on the fruits, pulp and seeds of 106 palm species (40 palm genera) – we show that the majority of interactions involve fruit-eating (545 records; birds, mammals, fish and reptiles), followed by seed-eating (383 records; birds and mammals) and pulp-eating (113 records; birds and mammals).

(3) All types of feeding interactions can result in positive, negative and dual outcomes for palms. However, the majority of fruit-eating (77%) and pulp-eating (91%) results in mutualistic outcomes whereas seed-eating mostly results in dual outcomes (60%). Variation in interaction outcomes can partly be explained by the consumption behaviour of animals, e.g. in relation to digestive processing (seed defecation versus regurgitation), caching behaviour (caching versus non-caching) and fruit-handling ability (high, intermediate, low). Positive interaction outcomes can be derived from mechanisms such as gut treatment (e.g. leading to increased seed viability and better seed germination after gut passage), endozoochory (e.g. long-distance seed dispersal by large-bodied fruit-eaters), ectozoochory and pulp removal (e.g. discarding of viable and defleshed seeds after dispersal by pulp-eaters), or seed caching (e.g. facilitation of seedling establishment after seeds are forgotten or neglected by cache owners). Negative outcomes emerge when seeds are damaged during fruit, pulp or seed consumption, or when seeds are deposited in sites that are non-suitable for germination and establishment.

(4) Mutualism–antagonism continuums illustrate how outcomes of plant–frugivore interactions vary intra- and inter-specifically. The mutualism–antagonism continuums for palm–frugivore interactions reveal that most animal species can act as both mutualists and antagonists. The frequency of such outcomes ultimately determines the predominant contribution of frugivores to palm population dynamics and their position along the continuum. These species-level continuums show that ungulates can be highly mutualistic (e.g. the lowland tapir Tapirus terrestris), but also predominantly antagonistic (e.g. the white-lipped peccary Tayassu pecari and the collared peccary Pecari tajacu). Similarly, rodents fall both on the mutualistic side (e.g. the pacu Caninus pacu, the Central-American agouti Dasyprocta punctata and the red-tailed squirrel
**The mutualism-antagonism continuum**

*Notocnus granatensis* and on the antagonistic side (e.g. the red-rumped agouti *D. leporina* and the Brazilian squirrel *Guerlinguetus brasiliensis*). Most bird species, on the other hand, fall on the mutualistic side of the continuum. Family-level continuums further highlight the contribution of overlooked mutualists such as carnivores (Canidae and Procyonidae) and marsupials (Didelphidae) and reinforce the mostly antagonistic nature of animal groups such as certain rodents or ungulate families.

(5) Evaluating the full range of positive and negative effects of fruit-eating, pulp-eating and seed-eating on plant demography during the seed-to-seedling transition is challenging. Nevertheless, a range of methods exists to quantify how animals affect seed viability, seed deposition and seedling establishment of plants. Besides visual observations at focal trees, experimental approaches in the laboratory and field can quantify fruit removal rates (e.g. with cafeteria plots or enclosure experiments) and the effects of frugivores on seed survival and seedling establishment (e.g. with germination experiments of gut-passed, handled or cached seeds). The application of relatively recent and novel methods such as camera traps, individual seed tracking with telemetric thread tags, or molecular techniques such as genetic parentage analysis considerably expand the toolbox for assessing the effects of frugivores on plant population dynamics and demography. Trait-based approaches will further help to elucidate the link between fruit-, pulp- and seed-eaters and ecosystem functions such as carbon storage.

(6) Our review is a first step towards synthesizing the effect of different consumption-related behavioural traits and feeding guilds of vertebrate fruit consumers on plant regeneration and ecosystem functioning. While a large amount of evidence is available for fruit- and seed-eaters, little is still known about how pulp-eaters or certain groups of fruit-eaters (e.g. frugivorous fish and reptiles) influence seed-to-seedling transitions in animal-dispersed plants. Moreover, quantitative evidence of intra- and inter-specific variability in outcomes of plant–frugivore interactions along the mutualism–antagonism continuum remains limited, as does information on seed dispersal distances. Filling this gap will require more studies that link feeding behaviour and animal traits to the effects of frugivores on seed viability, seed deposition and seedling establishment, and the consideration of how different feeding types and their interaction outcomes affect ecosystem functions and dynamics.

**IX. ACKNOWLEDGMENTS**

W.D.K. acknowledges funding from the Netherlands Organization for Scientific Research (824.15.007) and the University of Amsterdam (via a starting grant and through the Faculty Research Cluster ‘Global Ecology’). C.M.D thanks Christian Ziegler and Mathias Pires for kindly sharing pictures of frugivores feeding on palm fruits. We thank Mauro Galetti, one anonymous referee and the Assistant Editor Alison Cooper for their valuable feedback during the revision. We further thank the members of the Biogeography & Macroecology (BIOMAC) lab at the University of Amsterdam for fruitful discussions. Finally, we thank all authors involved in original research on palm–frugivore interactions in the Americas for their scientific contributions and their outstanding fieldwork under often difficult conditions.

**X. REFERENCES**

References identified with an asterisk are used in the online Supporting Information and associated Dryad dataset (Dracxler & Kissing, 2021).

Acevedo-Quintero, J. F. & Zamora-Árreguía, J. G. (2016). Role of mammals on seed dispersal and predation processes of *Mauritia flexuosa* (Arecaceae) in the Colombian Amazon. *Revista de Biología Tropical* 64, 5–15.

Acevedo-Quintero, J. F., Zamora-Árreguía, J. G. & Ortega-León, Á. M. (2018). The prickles of *Artocarpus altilis* as a structural defence to avoid seed predation. *Food Weeds* 16, e00088.

Adler, G. H. & Kentell, D. W. (1998). Fates of Neotropical tree seeds influenced by spiny rats (*Proechimys semispinus*). *Biotaica* 30, 677–681.

Aliaga-Rosell, E. & Fragoso, J. M. (2013). Deforestation affects *Artocarpus altilis* (Arecaceae) seed survivorship in a sub-montane tropical forest. *Revista de Biología Tropical* 63, 57–67.

Alvés, B. C., Mendes, C. P. & Ridero, M. C. (2018). Queen palm fruit selection and foraging techniques of squirrels in the Atlantic Forest. *Biotaica* 50, 274–281.

Anderson, J. T., Nuttall, T., Saldaña Rojas, J. S., Pendergast, T. H. & Flecker, A. S. (2011). Extremely long-distance seed dispersal by an overfished Amazonian frugivore. *Proceedings of the Royal Society B: Biological Sciences* 278, 3329–3335.

Alvarado, P. C., Mota, J. V. L. & Carvalho, A. A. F. (2013). Interações mutualistas entre aves frugívoras e plantas em um fragmento urbano de Mata Atlântica, Salvador, BA. *Revista Brasileira de Ornitologia* 19, 63–73.

Andradez, C. S., Pires, A. S. & Fernandes, F. A. (2009). Maníforas e palmeiras neotropicas: interações em paisagens fragmentadas. *Oecologia Brasiliensis* 13, 554–574.

Andradez, C. S., Pimenta, C. S., Pires, A. S., Fernandes, F. A. S., Oliveira-Santos, L. G. & Menezes, J. F. S. (2012). Increased productivity and reduced seed predation favor a large-seeded palm in small Atlantic Forest fragments. *Biota* 44, 237–245.

*Barbosa, P. S., Castro, M. F., Ojeda, R. A. & Navarro, J. L. (2015). Diet of wild boar (*Sus scrofa*) in a protected area of Argentina: the importance of baiting. *Manuscript Research* 60, 81–87.

*Barroso-Villalba, A., Blanco, G., Diaz-Luque, J. A., Dener, F. V., Hiraldo, F. & Tell, J. L. (2017). Seed dispersal by macaws shapes the landscape of an Amazonian ecosystem. *Scientific Reports* 7, 7373.

Barrons, T. A. P. & Montag, L. F. A. (2017). The role of *Lithodon dorsalis* (Sphagnetidae: Dendrophylax) as seed disperser in Eastern Amazon. *Neotropical Ichthyology* 15, e160061.

Barcellos, A. R., Borowicz, P. E. D., Sansaatti, T. M. & Gribe, R. (2013). Seed germination from lowland tapir (*Tapirus terrestris*) focal samples collected during the dry season in the northern Brazilian Amazon. *Integration Zoology* 8, 63–73.

Beck, H. (2006). A review of peccary-palm interactions and their ecological ramifications across the Neotropics. *Journal of Mammalogy* 87, 519–530.

Beckman, N. G., Aslan, C. E., Rogers, H. S., Kogan, O., Bronstein, J. L., Bullock, J. M., Hartig, F., Hille Ris Lambers, J., Zhou, Y., Zurell, D., Bronis, J. F., Bruno, E. M., Cantrell, R. S., Becker, R. R., Eshom, E. & et al. (2020). Advancing an interdisciplinary framework to study seed dispersal ecology. *Adu Planta* 12, e0484.

* Beckman, N. G. & Muller-Landau, H. C. (2007). Differential effects of hunting on pre-dispersal seed predation and primary and secondary seed removal of two Neotropical tree species. *Biotaica* 39, 328–339.

* Biological Reviews 97 (2022) 527–553 © 2021 The Authors. Biological Reviews published by John Wiley & Sons Ltd on behalf of Cambridge Philosophical Society.
Butler, C. J., Galetti, M., Pizo, M. A., Magnago, L. F. S., Rocha, M. F., Lima, R. A., Peres, C. A., Ovaskainen, O. & Jordano, P. (2015). Defaunation affects carbon storage in tropical forests. Science Advances 1, e1501105.

Bender, I. M. A., Kissling, W. D., Blendinger, P. G., Börnhöft-Gaese, K., Hensen, I., Köhn, I., Muñoz, M. C., Neuschulz, E. L., Nowak, L., Queller, D. C., Sánchez, F., Santéllez, M., Streit, M. L., Wiegand, T., Delling, D. M., et al. (2018). Morphological trait matching shapes plant-plant frugivore networks across the Andes. Ecology 94, 1910–1919.

Blanco, G., Tell, J. L., Díaz-Luque, J. A. & Hiraldo, F. (2019). Multiple external seed dispersers challenge the megafaunal syndrome and the six-vector ecological function of livestock. Frontiers in Ecology and Evolution 7, 328.

Bodmer, R. & Ward, D. (2006). (Frugivory in large mammalian herbivores. In Large Herbivore Ecology: Ecosystem Dynamics and Conservation (eds K. Daniel, P. Duncan, R. Bergerown and J. P. Pastor), pp. 232–260. Cambridge University Press, Cambridge.

Brown, D. D. (2013). Fruit-eating by an olive invertebrate: palm fruit consumption in wild northern tamarindas (Tamarindus mexicana) in Panama. Etologia 12, 63–65.

Brown, L. & Karubian, J. L. (2018a). Habitat loss and fragmentation reduce effective gene flow by disrupting seed dispersal in a neotropical palm. Molecular Ecology 27, 3053–3060.

Brown, L. & Karubian, J. L. (2018b). Rare genotype advantage promotes survival and genetic diversity of a tropical palm. New Phytologist 218, 1658–1667.

Brown, L., Ottewell, K. & Karubian, J. (2015). Short-term genetic consequences of habitat loss and fragmentation for the neotropical palm Oenocarpus bancana. Hereditas 113, 389–395.

Butler, C. J. & Larson, M. (2020). Climate change winners and losers: the effects of climate change on five palm species in the southeastern United States. Ecology and Evolution 10, 10408–10425.

Carballo, S. D. O., Freitas, I. D. S., Morlanes, V., Katzenberger, M. & Calarco, C. (2019). Potential seed dispersers: a new facet of the ecological role of bats in a tropical rainforest. Linnarum 1786, Brotas Nototricha 19, e20180626.

Cáceres, L., Rudzki-Surowiak, M., de Granville, J. J., Bicca-Marques, J. C., De Granville, J. J. & Bicca-Marques, J. C. (2017). Fundamental species traits explain provisioning services of tropical forest. Trends in Ecology & Evolution 32, 180–187.

Carneiro, C. B. & Marchand-Machado, L. O. (2011). Diet and feeding behavior of the yellow-faced parrot (Aratinga auratus) in Brazil. Okapia J. J. Trop. Zool. 40, 255–272.

Cid, B., Oliveira-Santos, L. G. R. & Mourão, G. (2013). Seasonal habitat use of agoutis (Dasyprocta aguti) driven by the palm Attalea planifrons in Brazilian Pantanal. Biotropica 45, 380–305.

Cid, B., Figueiredo, L. A. T. E. M. M., Bibeau, A. F. & Fernández, F. A. (2014). Short-term success in the reintroduction of the red-humped agouti (Dasyprocta spixii), an important seed disperser, in a Brazilian Atlantic forest reserve. Tropical Conservation Science 7, 939–950.

Correa, S. B., Vinhèmiller, K. O., López-Fernández, H. & Galetti, M. (2007). Evolutionary perspectives on seed production and dispersal by fishes. Biomeca 37, 749–756.

Cortés, M. C. & Uriarte, M. (2013). Integrating frugivory and animal movement: a review of the evidence and implications for scaling seed dispersal. Biological Reviews 88, 255–272.

Couvreur, T. L. & Baker, W. J. (2013). Tropical rainforest evolution: palms as a model group. BMC Biology 11, 1–12.

De Almeida, L. B. & Galetti, M. (2007). Seed dispersal and spatial distribution of Attalea geoffroyana (Arecales) in two remnants of Cerrado in southeastern Brazil. Acta Oecologica 32, 180–187.

De Araújo, C. B. & Marchand-Machado, L. O. (2011). Diet and feeding behavior of the yellow-faced parrot (Aratinga auratus) in Brazil. Okapia J. J. Trop. Zool. 40, 255–272.

Dettl, W. B., Steiner, T. & Adler, G. H. (2015). Seed dispersal by rodents in a lowland forest in Central Panama. Journal of Tropical Ecology 31, 403–412.

Dobey, S., Masters, D. V., Scheick, B. K., Clark, J. D., Pelton, M. R. & Sunquist, M. E. (2005). Ecology of Florida black bears in the Okefenokee–Osceola ecosystem. Wildlife Monographs 158, 1–41.

Donatti, C. I., Golmares, F. R. & Galetti, M. (2009). Seed dispersal and predation in the endemic Atlantic rainforest palm, Attalea odoratissima, across a gradient of seed disperser abundance. Ecological Research 24, 1187–1195.

Dos Santos, J., Varasini, I. G. & Macher, V. C. (2018a). Effects of neighborhood on pollination and seed dispersal of a threatened palm. Acta Oecologica 89, 95–101.

Dos Santos, J., Varasini, I. G. & Macher, V. C. (2018b). Survey dataset on mutualistic interactions among Euterpe edulis Mart./Arecales and floral and frugivorous visitors considering influence of neighborhood plant density and availability of resources. Data in Brief 21, 2015–2019.
The mutualism-antagonism continuum

Dracxler, C. M. & Forget, P.-M. (2017). Seed caching by rodents favours seedling establishment of two palm species in a lowland Atlantic forest remnant. *Journal of Tropical Ecology* 33, 228–231.

Dracxler, C. M. & Kissling, W. D. (2021). Data from: the mutualism-antagonism continuum in Neotropical palm-frugivore interactions: from interaction outcomes to ecosystem dynamics. Dryad Dataset. https://doi.org/10.5061/dryad.61g1pjw3.

Forget, P.-M., Pires, A. S. & Fernández, F. A. (2011). Invertebrate seed predators are not all the same: seed predation by bruchine and scolytine beetles and ants. *Biota Neotropica* 11, 11.

Eguiarte, L. E., Burkez, A. Rodriguez, J., Martínez-Ramos, M., Sarukhan, J. & Pinedo, D. (1993). Direct and indirect estimates of neighborhood and effective population size in a tropical palm, *Astronium macrostemon*. *Evolution* 47, 75–87.

Espinosa, D. M., Kissling, W. D. & Baisley, H. (2011). Geographical ecology of the palms (Arecaceae): determinants of diversity and distributions across spatial scales. *Annuals of Botany* 108, 1391–1416.

Ezcurra, S., Rezende, L. H. & Basilius, H. (2011). Post-dispersal seed removal in a large-seeded palm by frugivorous mammals in Guiana: comparison with Myrrocoptes coelis. *Tropical Ecology* 32, 155–167.

Fain, S., Fleury, M., Donatti, C. I. & Galetti, M. (2009). Effects of frugivore impairment and seed predators on the recruitment of a keystone palm. *Acta Oecologica* 35, 188–196.

Falcón, W., Moll, D. & Hassan, D. M. (2020). Frugivory and seed dispersal by chelonians: a review and synthesis. *Biological Reviews* 95, 142–166.

Federici, J. M. & Delibes, M. (2013). Pulp feeders alter plant interactions with subsequent animal associates. *Journal of Ecology* 101, 1581–1588.

Federici, J. M., Zwyer, M. & Delibes, M. (2012). Thieves or mutualists? Pulp feeders enhance endochochore local recruitment. *Ecology* 93, 573–587.

Fer, F., Henry, O., Forget, P.-M. & Gavoty, M. (2001). Frugivory and seed dispersal by terrestrial mammals. In *Neotropical: Dynamics and Plant-Animal Interactions in a Neotropical Rainforest* (eds F. Bongers, P. Charles-Dominique, P.-M. Forget and M.汤), pp. 227–232. Klwer Academic, Dortrecht.

Filho, R. F., Andrade, B. M. T. & Bezerra, B. (2021). Trash, tasty and healthy: the red-back antiu (Daucus carota) feed on leaflets from blouse: capsicums (*Sapium flavum*). *Tropical Ecology* 62, 149–152.

Fleming, T. H., Brettinwisch, R. & Whitesides, G. H. (1987). Patterns of tropical vertebrate frugivore diversity. *Annual Review of Ecology and Systematics* 18, 91–109.

*Fleury, M.* & *Galetti, M.* (2006). Forest fragment size and microhabitat effects on palm seed predation. *Biological Conservation* 131, 1–13.

*Forget, P.-M.* (1991). Seed-scatterhoarding of the last Neotropical megafaunal element in Amazonia, the tapir. *Biota Neotropica* 3, 135–167.

*Forget, P.-M.* (1991). Distributions across spatial scales. *Annals of Botany* 108, 1391–1416.

*Ezcurra, S.* (2011). Post-dispersal seed removal in a large-seeded palm by frugivorous mammals in Guiana: comparison with *Myrrocoptes carolinus*. *Tropical Ecology* 32, 155–167.

Fain, S., Fleury, M., Donatti, C. I. & Galetti, M. (2009). Effects of frugivore impairment and seed predators on the recruitment of a keystone palm. *Acta Oecologica* 35, 188–196.

Falcón, W., Moll, D. & Hassan, D. M. (2020). Frugivory and seed dispersal by chelonians: a review and synthesis. *Biological Reviews* 95, 142–166.

Federici, J. M. & Delibes, M. (2013). Pulp feeders alter plant interactions with subsequent animal associates. *Journal of Ecology* 101, 1581–1588.

Federici, J. M., Zwyer, M. & Delibes, M. (2012). Thieves or mutualists? Pulp feeders enhance endochochore local recruitment. *Ecology* 93, 573–587.

Ferr, F., Henry, O., Forget, P.-M. & Gavoty, M. (2001). Frugivory and seed dispersal by terrestrial mammals. In *Neotropical: Dynamics and Plant-Animal Interactions in a Neotropical Rainforest* (eds F. Bongers, P. Charles-Dominique, P.-M. Forget and M. 汤), pp. 227–232. Klwer Academic, Dortrecht.

Filho, R. F., Andrade, B. M. T. & Bezerra, B. (2021). Trash, tasty and healthy; the red-back antiu (Daucus carota) feed on leaflets from blouse: capsicums (*Sapium flavum*). *Tropical Ecology* 62, 149–152.

Fleming, T. H., Brettchinsch, R. & Whitesides, G. H. (1987). Patterns of tropical vertebrate frugivore diversity. *Annual Review of Ecology and Systematics* 18, 91–109.

*Fleury, M.* & *Galetti, M.* (2006). Forest fragment size and microhabitat effects on palm seed predation. *Biological Conservation* 131, 1–13.

*Forget, P.-M.* (1991). Seed-scatterhoarding of the last Neotropical megafaunal element in Amazonia, the tapir. *Biota Neotropica* 3, 135–167.

*Forget, P.-M.* (1991). Distributions across spatial scales. *Annals of Botany* 108, 1391–1416.

*Ezcurra, S.* (2011). Post-dispersal seed removal in a large-seeded palm by frugivorous mammals in Guiana: comparison with *Myrrocoptes carolinus*. *Tropical Ecology* 32, 155–167.
HENDERSON, A. (2002). Evolution and Ecology of Palms. Bronx, NY: New York Botanical Garden Press.

Henderson, A. (2011). Internal dispersal of seed-inhabiting insects by vertebrate frugivores: a review and prospects. Integrative Zoology 6, 213–221.

HERRERA, C. M. (1982). Defence of ripe fruit from pests: its significance in relation to plant-disperser interactions. The American Naturalist 120, 218–241.

HIBERT, F., SABATIER, D., ANDROVIT, J., SCOTTI-SANTAGNE, G., GONZALEZ, S., PRIEVOY, M. F., GREENAND, P., CHEVE, J., CABON, H. & RICHARD-HANSSON, C. (2013). Botanical. Acta Oecologica 44, 141–145.

HIBERT, F., KAYS, R. & JANSSEN, P. A. (2012). Directed seed dispersal towards areas with low conspecific tree density by a scat-dispersing rodent. Ecology Letters 15, 1423–1429.

HOCH, G. A. & ADLER, G. H. (1997). Removal of black palms (Arecaceae: Stannulataeae) seeds by humans. Plant Ecology 134, 201–228.

HULME, P. E. (1994). Post-dispersal seed predation in grassland: its magnitude and sources of variation. Journal of Ecology 82, 645–652.

HULME, P. E. & FERGUSON, S. (2007). Seed dispersal, destruction and demography. In Seed Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds D. J. LEVY, W. R. SILVA and M. GALETI), pp. 257–274. CAB Publishing, Wallingford.

HULME, P. E. & BENKMAN, C. W. (2002). Granivory. In Plant-Animal Interactions: An Evolutionary Approach (eds C. M. HERBEE and O. PELLISSIER), pp. 132–156. Oxford University Press, Oxford.

IZAWA, K. & MIZUNO, A. (1977). Palm-fruit cracking behavior of wild black-capped capuchin (Sapajus spilops). Primates 18, 773–792.

JANSSEN, P. A., BARTHOLOMUESEN, M., BORGERS, F., ELIZA, J., DEN OUDER, J. & VAN WIJRE, S. E. (2002). The role of seed size in dispersal by a scat-dispersing rodent. In Seed Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds D. J. LEVY, W. R. SILVA and M. GALETI), pp. 209–225. CAB International, Wallingford.

JANSSEN, P. A., BONGERS, F. & HEMERIK, L. (2004). Seed mass and mast seeding influence of relatedness, conspecific density and microhabitat on seedling establishment. Annual Review of Ecology and Systematics 35, 465–499.

JANSSEN, P. A., ELSCHOT, K., VIERHEIL, P. & WRIGHT, S. J. (2010). Seed predation and defleshing in the agouti-dispersed palm Arecaceae: Stannulataeae. Journal of Tropical Ecology 26, 473–480.

JANSSEN, P. A., HIRCH, B. T., EMMENS, W. J., ZAMORA-GUTIERREZ, V., WERKELIS, M. & KAYS, R. (2012). Thieving rodents as substitute dispersers of megaasial seedlings. Proceedings of the National Academy of Sciences of the United States of America 109, 12610–12615.

JANSSEN, P. A., VISKER, M. D., JOSOPH WRIGHT, S., RUTGEN, G. & MULLER-LANDAU, H. C. (2014). Negative density dependence of seed dispersal and seedling recruitment in a Neotropical palm. Ecology Letters 17, 1111–1120.

JANSSEN, D. H. (1971). Seed predation by animals. Annual Review of Ecology and Systematics 2, 465–492.

JERZOLINSKAS, A., RIBEIRO, M. B. N. & MARTINS, M. (2009). Are tortoises important seed dispersers in Amazonian forests? Acta Oecologia 36(1), 517–528.

JORDAN, P. P. (2007). Significance of mutualistic interactions in pollination and seed dispersal: connectance, dependence asymmetries, and coevolution. The American Naturalist 129, 657–677.

JORGES, M. L. S. P. & HOWE, H. F. (2009). Can forest fragmentation disrupt a neotropical scatter-hoarding rodent. In Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds P. A. JANSSEN, W. E. OLANDER, J. GALETI & P. J. KUPREWICZ), pp. 89–107. CAB International, Wallingford.

JORGES, M. L. S. P. & HOWE, H. F. (2009). Can forest fragmentation disrupt a neotropical scatter-hoarding rodent. In Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds P. A. JANSSEN, W. E. OLANDER, J. GALETI & P. J. KUPREWICZ), pp. 89–107. CAB International, Wallingford.

JORGES, M. L. S. P. & HOWE, H. F. (2009). Can forest fragmentation disrupt a neotropical scatter-hoarding rodent. In Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds P. A. JANSSEN, W. E. OLANDER, J. GALETI & P. J. KUPREWICZ), pp. 89–107. CAB International, Wallingford.

JORGES, M. L. S. P. & HOWE, H. F. (2009). Can forest fragmentation disrupt a neotropical scatter-hoarding rodent. In Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds P. A. JANSSEN, W. E. OLANDER, J. GALETI & P. J. KUPREWICZ), pp. 89–107. CAB International, Wallingford.

JORGES, M. L. S. P. & HOWE, H. F. (2009). Can forest fragmentation disrupt a neotropical scatter-hoarding rodent. In Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds P. A. JANSSEN, W. E. OLANDER, J. GALETI & P. J. KUPREWICZ), pp. 89–107. CAB International, Wallingford.

JORGES, M. L. S. P. & HOWE, H. F. (2009). Can forest fragmentation disrupt a neotropical scatter-hoarding rodent. In Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds P. A. JANSSEN, W. E. OLANDER, J. GALETI & P. J. KUPREWICZ), pp. 89–107. CAB International, Wallingford.

JORGES, M. L. S. P. & HOWE, H. F. (2009). Can forest fragmentation disrupt a neotropical scatter-hoarding rodent. In Dispersal and Frugivory: Ecology, Evolution, and Conservation (eds P. A. JANSSEN, W. E. OLANDER, J. GALETI & P. J. KUPREWICZ), pp. 89–107. CAB International, Wallingford.
The mutualism-antagonism continuum

551

*Menegaro, A. L. & Pivello, V. R. (2012). Phenology and fruit traits of Archontophoenix cunninghamiana, an invasive palm tree in the Atlantic forest of Brazil. *Ecotropica* **18**, 45–54.

*Meriney, S. D., Bischler, C. M., Santos-Guimaraes, M. R. O. P. & Pires, A. S. (2021). Sowing forests: a synthesis of agouti (*Dasyprocta* spp.) seed dispersal and predation and their influence on plant communities. *Biological Reviews*. https://doi.org/10.1111/brr.12761.

Moenenburg, S. M. & Levey, D. J. (2003). Do frugivores respond to fruit harvest? An experimental study of short-term responses. *Ecology* **84**, 2660–2663.

*Moher, D., Liberati, A., Tetzlaff, J., Altman, D. G. & Prisma Group (2009). Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *PLOS Medicine* **6**, e1000097.

Montesinos-Navarro, A., Hiraldo, F., Tellas, J. L. & Blanco, G. (2017). Network structure enhancing mutualism-antagonism continua increases community robustness. *Nature Ecology & Evolution* **1**, 1661–1669.

*Montes-Sánchez, J. J., Huato-Sobarian, L., Buntin-Dios, S. E. & León-de La Luz, J. L. (2020). The feral pig in a poor impacted ecosystem: analysis of diet composition and its utility. *Rangeoland* & *Ecology Management* **73**, 703–711.

Morini, M. S. C., Diniz, A. O., Lewis, S. L., Silva, R. R. & Kato, L. M. (2003). Non-specific interaction and fruit dispersal by ants (*Hypermelitta; Formicidae*) and fruits of *Syagrus romanzoffiana* (*Arecaceae*) in an area of the Brazilian Atlantic Forest. *Soilbiology* **42**, 663–673.

Munóz, G., Trojelgaard, K. & Kissling, W. D. (2019). A synthesis of animal-mediated seed dispersal of palms reveals distinct biogeographical differences in seed intervals. *Journal of Biogeography* **46**, 466–484.

Muscarella, R., Emilio, T., Phillips, O. L., Lewis, S. L., Silva, R. R., Baker, W. J., Couvreur, T. L. P., Eisert, W. H., Svending, J. C., Assunção-Fauroe, K., Aiba, S.-I., Almeida, E. C., Almeida, S. S., Oliveira, E. A., Álvarez-Dávila, E., et al. (2020). The global abundance of tree palms. *Global Ecology and Biogeography* **29**, 403–414.

Nascimento, L. F. D., Guimaraes, P. R. Jr., Onstein, R. E., Kissling, W. D. & Pires, M. M. (2020). Associated evolution of fruit size, fruit colour and spines in Neotropical palms. *Journal of Evolutionary Biology* **33**, 838–860.

Nathan, R. (2006). Long-distance dispersal of plants. *Science* **313**, 786–788.

*Obidul, G., Silva, C. M. & Campom-Arceit, A. (2015). Frugivory and seed dispersal by tapirs: an insight on their ecological role. *Integrative Zoology* **8**, 4–17.

*Olsson, F., Pardini, R., Bouhloola, R. L. P., Bürger, R. & Morello, C. (2019). Do tapirs steal food from palm seed predators or give them a lift? *Biota* **31**, 373–375.

Onstein, R. E., Baker, W. J., Couvreur, T. L., Faury, S., Svending, J. C. & Kissling, W. D. (2017). Frugivory-related traits promote speciation of tropical palms. *Nature Ecology & Evolution* **1**, 1903–1911.

Ottewell, K., Browne, L., Carrera, D., Olivero, J. & Karubian, J. (2018). Genetic diversity of dispersed seeds is highly variable among gods of the long-winded umbrella-tortoise. *Acta Oecologica* **86**, 31–37.

*Palacios, E., Rodriguez, A. & Defler, T. R. (1997). Diet of a group of Callicebus torquatus lambei (Humboldt, 1812) during the annual resource bottleneck in Amazonian Colombia. *International Journal of Primatology* **18**, 503–522.

*Pacheco, L. F., Lopes, M. C. & Campos-Arceit, A. (2019). Spatial genetic structure, genetic diversity and pollen dispersal in a harvested population of *Arfocactus acutifolius* in the Brazilian Amazon. *BMC Genet* **16**, 1–10.

*Ribeiro, L. F., Conde, L. O. M. & Tabarelli, M. (2010). Predação e remoção de sementes de cinco espécies de palmeiras por *Guaratinga ingrami* (Thomas, 1901) em um fragmento de floresta costeira Atlântica. *Revista Amanã* **04**, 637–649.

Rios, R. S. & Pacheco, L. F. (2006). The effect of dung and dispersal on post-seedling establishment of *Atalaia phalacraea* (Arecaceae) by brachial beetles. *Biota* **38**, 778–781.

Robertson, A. W., Trass, A., Leadbe, J. J. & Kelly, D. (2006). Assessing the benefits of frugivory for seed germination: the importance of the dehisulation stage. *Functional Ecology* **20**, 58–66.

*Roberts, J. R., Traveset, A., Santamaria, E. & Alves, I. R. (2018). Non-speciation in plants: putting the fruit in the acronym species. *Flora* **293**, 1–15.

*Rodrigues, M., Olmos, F. & Galetti, M. (1993). Seed dispersal by tapirs in southeastern Brazil. *Mammalia* **57**, 460–461.

Rodríguez-Rodríguez, M. C., Jardel, P. & Valido, A. (2017). Functional consequences of plant-animal interactions along the mutualism-antagonism gradient. *Ecology* **98**, 1266–1276.

*Rojo-Robles, G., Gayo, N. S. & Muñoz-Sara, V. (2012). Frugivory and seed dispersal of *Oncosperma hatusa* (Arecaceae) in a forest from the Colombian Andes. *Revista de Biología Tropical* **60**, 1451–1461.

*Roldán, A. I. & Simenett, J. A. (2001). Plant-plant interactions in tropical Bolivian forests with different hunting pressures. *Conservation Biology* **15**, 617–623.

Roman, C., Neto, L. T. & Cáceres, N. C. (2010). Fruit manipulation of the palm *Syagrus romanzoffiana* by vertebrates in southern Brazil. *Neotropical Biodiversity and Conservation* **5**, 101–105.

*Russo, S. E., Portnay, S. & Auspurgter, C. K. (2006). Incorporating animal behavior into seed dispersal models: implications for seed shadows. *Ecology* **87**, 3160–3174.

Sales, L. P., Kissling, W. D., Galetti, M., Naimi, B. & Pires, M. M. (2021). Climate change reshapes the co-evolutionary dynamics of a Neotropical seed-dispersal system. *Global Ecology and Biogeography* **30**, 1129–1138.

Samuels, I. A. & Levey, D. J. (2005). Effects of gut passage on seed germination: do experiments answer the questions they ask? *Functional Ecology* **19**, 365–368.

*Sanos, N. N. D. (2017). Estudio da dispersão do Bactra cucuia (L.) (Lepidoptera: Saturniidae) entre árvores de *Syagrus romanzoffiana* (Cham.) *Glasmann* (*Arecaceae)*. *Revista da Biologia Tropical* **15**, 945–956.

*Rodrigues, M., Olmos, F. & Galetti, M. (1993). Seed dispersal by tapirs in southeastern Brazil. *Mammalia* **57**, 460–461.

*Sanos, N. N. D. (2017). Estudo da dispersão do Bactra cucuia (L.) (Lepidoptera: Saturniidae) entre árvores de *Syagrus romanzoffiana* (Cham.) *Glasmann* (*Arecaceae)*. *Revista da Biologia Tropical* **15**, 945–956.

*Sanos, N. N. D. (2017). Estudo da dispersão do Bactra cucuia (L.) (Lepidoptera: Saturniidae) entre árvores de *Syagrus romanzoffiana* (Cham.) *Glasmann* (*Arecaceae)*. *Revista da Biologia Tropical* **15**, 945–956.
Smythe, N., Silva, F. R., Stevenson, P. R., Sezen, U. U. (2020). Biological Reviews 55, 2020–2037.

Schupp, E. W., Jordano, P., & Gómez, J. M. (2010). Seed dispersal effectiveness revisited: a conceptual review. New Phytologist 188, 333–353.

Schupp, E. W., Zvolak, R., Jones, L. R., Snell, R. S., Beckman, N. G., Aman, C., Cavazos, B. R., Effiom, E., Frick, E. C., Montaño-Centellas, F., Poulsen, J., Razafindratsima, O. H., Sandor, M. E. & Sheh, K. (2019). Intrinsic and extrinsic drivers of intraspecific variation in seed distribution of Butia odorata (Arecaceae) seedlings in savanna-like palm tree formations in southern Brazil. Austral Ecology 44, 1398–1409.

Schupp, E. W., Jordano, P., & Gómez, J. M. (2010). Seed dispersal effectiveness revisited: a conceptual review. New Phytologist 188, 333–353.

Schupp, E. W., Zvolak, R., Jones, L. R., Snell, R. S., Beckman, N. G., Aman, C., Cavazos, B. R., Effiom, E., Frick, E. C., Montaño-Centellas, F., Poulsen, J., Razafindratsima, O. H., Sandor, M. E. & Sheh, K. (2019). Intrinsic and extrinsic drivers of intraspecific variation in seed distribution of Butia odorata (Arecaceae) seedlings in savanna-like palm tree formations in southern Brazil. Austral Ecology 44, 1398–1409.

Schupp, E. W., Zvolak, R., Jones, L. R., Snell, R. S., Beckman, N. G., Aman, C., Cavazos, B. R., Effiom, E., Frick, E. C., Montaño-Centellas, F., Poulsen, J., Razafindratsima, O. H., Sandor, M. E. & Sheh, K. (2019). Intrinsic and extrinsic drivers of intraspecific variation in seed distribution of Butia odorata (Arecaceae) seedlings in savanna-like palm tree formations in southern Brazil. Austral Ecology 44, 1398–1409.

Schupp, E. W., Zvolak, R., Jones, L. R., Snell, R. S., Beckman, N. G., Aman, C., Cavazos, B. R., Effiom, E., Frick, E. C., Montaño-Centellas, F., Poulsen, J., Razafindratsima, O. H., Sandor, M. E. & Sheh, K. (2019). Intrinsic and extrinsic drivers of intraspecific variation in seed distribution of Butia odorata (Arecaceae) seedlings in savanna-like palm tree formations in southern Brazil. Austral Ecology 44, 1398–1409.

Schupp, E. W., Zvolak, R., Jones, L. R., Snell, R. S., Beckman, N. G., Aman, C., Cavazos, B. R., Effiom, E., Frick, E. C., Montaño-Centellas, F., Poulsen, J., Razafindratsima, O. H., Sandor, M. E. & Sheh, K. (2019). Intrinsic and extrinsic drivers of intraspecific variation in seed distribution of Butia odorata (Arecaceae) seedlings in savanna-like palm tree formations in southern Brazil. Austral Ecology 44, 1398–1409.

Schupp, E. W., Zvolak, R., Jones, L. R., Snell, R. S., Beckman, N. G., Aman, C., Cavazos, B. R., Effiom, E., Frick, E. C., Montaño-Centellas, F., Poulsen, J., Razafindratsima, O. H., Sandor, M. E. & Sheh, K. (2019). Intrinsic and extrinsic drivers of intraspecific variation in seed distribution of Butia odorata (Arecaceae) seedlings in savanna-like palm tree formations in southern Brazil. Austral Ecology 44, 1398–1409.
XI. Supporting information

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

**Appendix S1.** Combination of search terms for literature compilation in the Web of Science.

**Appendix S2.** Interaction types and criteria to classify interactions into positive, negative or dual outcomes based on quantitative and qualitative evidence from published articles on palm–frugivore interactions.

**Appendix S3.** List of palm species by tribe and genera recorded in interactions with frugivores in the data set with types of feeding interactions.

**Appendix S4.** Number of animal species and interaction records in different types of feeding interactions (fruit-, pulp-, and seed-eating), summarized for each family, order and class.

**Appendix S5.** The role of digestive-processing types for fruits in interactions between fruit-eaters and palms.

**Appendix S6.** Dispersal distances recorded for interactions between vertebrate frugivores and Neotropical palm species.

**Appendix S7.** The role of fruit-handling ability in interactions between pulp-eaters and palms.

**Appendix S8.** The role of handling ability and seed-caching behaviour in interactions between seed-eaters and palms.

**Appendix S9.** Fruit size of palm species recorded in interactions with frugivores, according to parts of fruits consumed by animals.

(Received 10 March 2021; revised 14 October 2021; accepted 18 October 2021; published online 1 November 2021)