Importance of the lilac-crowned parrot in pre-dispersal seed predation of 

*Astronium graveolens* in a Mexican tropical dry forest

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(Received 21 September 2009)

**Abstract:** Parrots represent a large biomass of canopy granivores in tropical forests, and may be effective pre-dispersal seed predators. We evaluated the importance of the lilac-crowned parrot (*Amazona finschi*) as a pre-dispersal seed predator of *Astronium graveolens* (Anacardiaceae) in tropical dry forest. Seeds were collected in fruit-traps beneath 22 trees to compare pre-dispersal seed predation by parrots and insects, and determine whether intensity of seed predation was related to fruit-crop size or the aggregation of fruiting conspecifics around focal trees. Ground-level exclosures were established to compare post-dispersal seed predation by vertebrates and insects. The lilac-crowned parrot predated 43% of seeds pre-dispersal, while insects predated only 1.3%. Intensity of pre-dispersal seed predation by parrots was significantly greater in high-fruiting 0.79-ha resource patches, and was not related to fruit abundance of the focal tree. Foraging parrots also discarded immature fruits below the tree, causing a total 56% pre-dispersal loss of seed production, which was greater than post-dispersal removal by vertebrates, mainly rodents (51%) or insects (36%). Our results show that parrots play an important role as pre-dispersal seed predators in tropical dry forests. The reduction of parrot populations in tropical forests may have consequences for seed predation, affecting recruitment patterns of canopy trees.

**Key Words:** *Amazona finschi*, Chamela-Cuixmala, fruit crop size, granivory, Mexico, plant–animal interaction, post-dispersal seed predation, Psittacidae, resource patch, semi-deciduous forest

**INTRODUCTION**

Parrots (Psittacidae) constitute a high proportion of the biomass of canopy granivores in tropical forests (Terborgh et al. 1990), and consume a large number of seeds of canopy trees (Francisco et al. 2008, Galetti 1993, Renton 2001, 2006). This, combined with the tendency to form large feeding flocks, high mobility, and the ability to track fluctuations in food resource availability (Renton 2001), means that parrots may be effective pre-dispersal predators on canopy seed resources.

However, few data exist on the intensity of pre-dispersal seed predation by parrots, and only a few studies in humid forests provide estimates of 6.4–10% pre-dispersal seed predation for various parrot species (Galetti & Rodrigues 1992, Howe 1980, Trivedi et al. 2004). By comparison, in fragmented habitats, where food resources may be limited, parrots damage 20–67% of fruit crops (Coates-Estrada et al. 1993, Francisco et al. 2008, Galetti & Rodrigues 1992). Furthermore, no studies have compared seed predation by parrots to other sources of pre- and post-dispersal seed predation, particularly insects, which are important, specialist pre-dispersal seed predators (Janzen 1980).

Studies in tropical moist forests that have evaluated the relative roles of insects and vertebrates in pre- and post-dispersal seed predation found insects to be the main pre-dispersal seed predators (Curran & Leighton 2000, Forget et al. 1999, Nakagawa et al. 2005), while predation by arboreal vertebrates was sporadic (Curran & Leighton 2000, Nakagawa et al. 2005). However, vertebrates appear to play a greater role in post-dispersal seed predation, with rodents removing more seeds than insects (Anderson & MacMahon 2001, Curran & Leighton 2000, Westerman et al. 2003).

In contrast with humid forest, tropical dry forest presents a marked seasonality in rainfall and plant
phenology, where many tree species fruit during the dry season (Bullock & Solis-Magallanes 1990, Janzen 1967). Furthermore, the biomass of canopy arthropods in dry forest decreases by an average of 92% in the dry season compared with the rainy season (Lister & García-Aguayo 1992), including Coleoptera, which are among the main pre-dispersal seed predators (Janzen 1980). Therefore, seed predation patterns in tropical dry forest during the dry season may differ from those of humid forests.

Furthermore, dry-forest tree species tend to be clumped in distribution (Hubbell 1979), and present high synchrony in fruit production (Janzen 1967), affecting seed-resource abundance and dispersion. It has been found that either fruit-crop size or the local abundance of fruiting trees may influence foraging by parrot seed predators (Cameron & Cunningham 2006, Coates-Estrada et al. 1993, Francisco et al. 2008, Howe 1980). However, crop size of an individual tree may be less effective in attracting canopy vertebrates where there is high fruit production (Ortiz-Pulido & Rico-Gray 2000).

The dry-forest tree Astronium graveolens fruits at the end of the dry season (Pennington & Sarukhán 2005, Renton 2001), when insect abundance is low (Lister & García-Aguayo 1992), and its seeds are predominant in the diet of the lilac-crowned parrot (Renton 2001). Therefore, we hypothesize that canopy vertebrates, in particular the lilac-crowned parrot, rather than insects, will be the principal seed predators for A. graveolens, and tested this by comparing seed predation by vertebrates and insects at both the pre- and post dispersal stages. Furthermore, fruiting of A. graveolens occurs when few other food resources are available for the lilac-crowned parrot (Renton 2001), therefore we expect seed predation by the lilac-crowned parrot to be consistent between years. Finally, given the highly synchronized fruiting of dry-forest trees which may limit effectiveness of crop size, and clumped distribution of tree species creating resource patches, it may be expected that in tropical dry forest the local aggregation of fruiting trees has a greater influence on foraging by canopy vertebrates. We therefore predict that the intensity of pre-dispersal seed predation by parrots will be related to the aggregation of fruiting trees, rather than fruit-crop size, being highest where fruiting trees are more clumped in distribution.

STUDY SITE

We conducted the study during April to July of 2007 and 2008 in the tropical dry forest of the 13 142-ha Chamela-Cuixmala Biosphere Reserve (19°22′N–19°35′N, 104°56′W–105°03′W), on the Pacific coast of Jalisco, Mexico. Mean annual precipitation at the study site is 748 mm, with 85% of rainfall occurring from June to October, and a prolonged drought from mid-February to late May (Bullock 1986). The reserve has a hilly topography varying in elevation from 20 m to 520 m asl. The dominant vegetation type on the slopes is seasonally deciduous forest, with the great majority of trees dropping their leaves for 5–8 mo of the year (Rzedowski 1994). Small areas of semi-deciduous vegetation, in which tree species are evergreen or drop leaves for only 1–3 mo (Rzedowski 1994), occur along watercourses, known locally as ‘arroyos’, and in larger valleys (Lott 1993).

Astronium graveolens is a common tree species of semi-deciduous forest at the study site (Lott et al. 1987), and has a wide range from Mexico to South America (Pennington & Sarukhán 2005). Fruiting occurs during March to June at the end of the dry season when food resources are limited (Renton 2001, 2002), and coincides with the period when the lilac-crowned parrot is raising young (Renton 2002, Renton & Salinas-Melgoza 1999). Fruits are 1–1.5 cm long, containing a single seed of 9–10 mm long, and are wind-dispersed.

METHODS

Pre-dispersal seed predation

At the initiation of fruiting of A. graveolens, we set up four fruit-traps beneath the canopy of each of 19 trees in 2007, and 10 trees in 2008, seven of which were sampled in both years. Each fruit-trap was constructed of 1-mm nylon mesh with an area of 0.5 m², and was supported 1 m from the ground by aluminium posts (Nakagawa et al. 2005). We checked fruit-traps weekly throughout the fruiting period, and collected all fruits or remains of fruits that had fallen into the fruit-traps, placing them in separate paper bags for each fruit-trap by tree and collection date.

Each sample was then analysed in the laboratory to determine the number of ripe and immature whole fruits, and the number of fruits attacked by vertebrates (such as parrots), insects, or fungi. We discounted any fruits of <0.5 mm length as these had not yet formed a seed kernel that would be used by vertebrate or invertebrate seed predators. Ripe fruits of A. graveolens have a light-brown colour, in which the remains of the petals are open in a star-shape, with a yellow-coloured seed kernel. By contrast, immature fruits are green, with closed petals, and a green-coloured seed kernel. Immature whole seeds collected in the fruit-traps represent a waste of seed resources for the tree as they are unlikely to germinate.

Fruits attacked by parrots were easily identified by the characteristic half-moon beak bite-mark used to open the fruit and extract the seed kernel. The fruit remains are then dropped whole by the foraging parrot to fall in the fruit-traps below. Each A. graveolens fruit contains only one seed, therefore we could quantify the number of
seeds predated by parrots by the number of complete fruit remains collected in the fruit-traps.

All whole fruits were analysed under a stereoscopic microscope, and cut open to determine whether they contained healthy seeds, were empty, or had been attacked by invertebrates or fungi. We identified insect predation by small holes made on the outside of the fruit, or by the presence of insect faeces. Based on this analysis we determined the number and proportion of whole, ripe fruits with potential to germinate, as well as the number of complete, immature fruits wasted and dropped beneath the maternal tree, and the proportion of fruits attacked by vertebrates, insects, or fungi, or that had not produced a seed.

**Fruit-crop size and aggregation of fruiting trees**

We estimated initial fruit abundance for 18 focal trees in 2007 by visual counts of the number of fruits in the canopy, following the method described by Chapman et al. (1992). Visual counts were made of the number of fruits on each of five racemes in different areas of the canopy, using a 60× magnification telescope. We corroborated these estimates by counting the number of fruits on whole racemes collected from the tree. We then calculated the mean number of fruits per raceme, and estimated the number of racemes per branch of the tree, multiplied by the number of fruiting branches in the tree.

To determine the local aggregation of fruiting conspecifics around the focal tree, we calculated a neighbourhood fruiting index for 17 focal trees in 2007, and 10 focal trees in 2008. We established a circular plot of 50-m radius around the focal tree, based on the largest area in which it was logistically feasible to evaluate resource availability (Saracco et al. 2005), and the fact that semi-deciduous forest tends to occur in narrow bands along stream-beds at the study site (Lott 1993). Within each circular plot, we measured the distance from the focal tree to every other fruiting tree of *A. graveolens*, as well as the diameter at breast height (dbh) of each fruiting tree. We then calculated the neighbourhood fruiting index for each circular plot as: (1/mean distance to fruiting trees) × number of fruiting trees × mean dbh of fruiting trees. Hence, a focal tree with few or distant neighbours will have a low neighbourhood fruiting index, while a focal tree close to a number of other fruiting trees will have a high neighbourhood fruiting index.

**Post-dispersal seed predation**

We established an exclusion experiment in six quadrats to evaluate the probability of post-dispersal survival of seeds of *A. graveolens*, and determine the intensity of post-dispersal seed predation by insects or vertebrates. Three quadrats were located in semi-deciduous forest within the Chamela area of the reserve, while the other three quadrats were located along the Arroyo Limbo and the Cuixmala area of the reserve, which presents deciduous forest vegetation with elements of semi-deciduous vegetation.

Each quadrat contained 18 lots consisting of an area of 30 cm² of bare soil cleared of leaf-litter, with 10 *A. graveolens* fruits placed in each lot. The 18 lots represented six replicates of three treatment conditions: open, exclusion and closed. In the open treatment, intact fruits were placed on the bare soil patch without any physical protection to permit access by either insects or vertebrates. In the exclusion treatment, fruits were protected by a cylindrical, 1.5-cm-diameter wire-mesh of 40 cm high by 15 cm wide, to permit access by insects but restrict vertebrate access. In the closed condition, lots were covered by a plastic case of 10 cm wide × 4 cm high to prevent animal access to the fruits. The base of the plastic case was buried into the soil to prevent potential predators entering from below the base.

Each experimental quadrat was positioned around an existing *A. graveolens* tree that was not fruiting during the experimental period, and was separated by a mean 1.17 ± 0.08 km (range = 1.06–1.24 km) from another quadrat. Each quadrat comprised an area of 25 × 10 m, with the 18 lots organized in six lines and three columns. Each lot was separated by 5 m, and the three treatment conditions were randomly allocated to lots, giving six replicates of each treatment condition (Figure 1).

Experimental quadrats were checked daily during the first 2 wk, and thereafter were checked every 2 d. At each inspection, we recorded the number of fruits present or removed from each of the lots. The experiment was run for 45 d until the onset of the rainy season, when many fruits were either washed away in the rainfall or began to germinate.

**Statistical analyses**

We conducted preliminary analysis using the Kolmogorov–Smirnov test with Lilliefors significance level, to evaluate normality of distribution for the number and proportion of seeds predated by parrots (number: K-S = 0.11, n = 29, P = 0.20; proportion: K-S = 0.10, n = 29, P = 0.20) and by insects (number: K-S = 0.13; n = 29, P = 0.20; proportion: K-S = 0.09; n = 29, P = 0.20). To evaluate the relative importance of vertebrates and invertebrates in pre-dispersal seed predation, we applied a paired t-test comparing the number and proportion of seeds predated by parrots and by insects, paired by focal tree. We also applied paired t-tests to compare the intensity of parrot seed predation
between years for the number ($K-S = 0.17, n = 14, P = 0.20$) and proportion ($K-S = 0.21, n = 14, P = 0.08$) of seeds predated by parrots in seven trees that were sampled in both 2007 and 2008.

We applied linear regression to determine whether pre-dispersal seed predation by parrots and insects was related to the abundance of fruits in the focal tree, or to the neighbourhood fruiting index. The dependent variables were total number and proportion of seeds predated by parrots and insects in each tree. Independent variables were the initial number of fruits in the tree estimated by visual counts, and the neighbourhood fruiting index for the focal tree. We plotted the residuals against the predicted values of the linear regression to confirm homogenity of variances (Quinn & Keough 2002). We also calculated the leverage $h_i$ values, and Cook’s $D_i$ from the standardized deleted residuals, and did not detect outliers with undue influence on the regression fit (Quinn & Keough 2002).

We applied a two-way ANOVA to determine whether the number of fruits remaining at the end of 45 d post dispersal differed between the two sites of Chamela and Cuixmala, and among the experimental treatments. We evaluated the pattern of daily survival of fruits post dispersal using the Kaplan–Meier probability of survival (Pollock et al. 1989) for the open and exclusion treatments in the two sites of Chamela and Cuixmala. We used the log-rank test to compare the pattern of survival for each treatment condition between the two sites (Pollock et al. 1989). Where there was no difference in treatments between sites, we pooled the data for the two sites to obtain an overall survival probability for the open and exclusion treatments.

In order to estimate the potential rate of removal of fruits by vertebrates alone, for each sampling interval we subtracted the number of fruits removed by insects in the exclusion treatment from the total number of fruits removed in the open treatment (Kelt et al. 2004). Where more fruits were removed in the exclusion treatment we left the estimate for vertebrate removal at zero. We then conducted log-rank analysis to determine whether the pattern of survival varied between fruits subjected to insects in the exclusion treatment, and the estimated removal by vertebrates alone. Data are presented as mean and SD, and range values, and we applied the $P < 0.05$ significance level for all statistical tests.

**RESULTS**

**Pre-dispersal seed predation**

Duration of fruiting for the 29 focal trees had a mean 5 ± 1.4 wk (range = 3–8 wk); with a mean total of 2028 ± 1047 fruits (range = 546–4920 fruits) per tree collected in the fruit-traps. Of the total number of fruits collected in all the fruit-traps, 42.6% had their seeds predated by parrots, while only 1.3% of fruits were damaged by insects and 2.7% by fungi. In addition, 13.8% represented immature fruits discarded by foraging parrots, while 19.2% of the fruits did not contain a viable seed, and may have been aborted. Hence, of all the fruits collected...
in the fruit-traps only 21.3% were mature fruits capable of germinating. Considering both direct seed predation and waste of immature fruits discarded below the tree, the foraging activity of parrots resulted in a loss of germination potential for 56.5% of the fruits collected in the traps.

As predicted, in each tree there was significantly more pre-dispersal seed predation by parrots compared with insects in both number (paired \( t_{28} = 8, P < 0.001 \)) and proportion (paired \( t_{28} = 14, P < 0.001 \)) of seeds predated. Parrots predated a mean 40% ± 14.7% (range: 4–65%, \( n = 29 \)) of the seeds collected in the fruit traps per tree, whereas insects predated only 1.5% ± 0.8% (range: 0.01–2.7%, \( n = 29 \)) of the fruits. The level of seed predation by parrots also did not vary between years for those trees that were sampled in 2007 and 2008 (number: paired \( t_6 = 0.09, P = 0.93 \); proportion: paired \( t_6 = 0.66, P = 0.53 \)).

**Influence of fruit-crop size and aggregation of fruiting trees**

Visual estimation of initial abundance of fruits in the crown of each tree varied from 11 000 to 939 000 fruits (mean: 283 000 ± 295 000, \( n = 18 \)), and focal trees had a mean neighbourhood fruiting index of 5.9 ± 6.1 (range: 0–15.6, \( n = 27 \)). The intensity of pre-dispersal seed predation by parrots was significantly related to the neighbourhood fruiting index of the focal tree (Figure 2), for both total number (\( r^2 = 0.55, F_{1,25} = 31, P < 0.001 \)), and proportion (\( r^2 = 0.32, F_{1,25} = 12, P = 0.002 \)) of seeds predated. In contrast, initial fruit abundance did not influence either the number (\( r^2 = 0.11, F_{1,16} = 2.0, P = 0.18 \)) or proportion (\( r^2 = 0.15, F_{1,16} = 2.8, P = 0.12 \)) of seeds predated by parrots. Hence, resource dispersion influenced the intensity of pre-dispersal seed predation by parrots, with parrots predating a mean 52% ± 3.9% of seeds for nine trees with a high number of conspecifics fruiting nearby, whereas parrots predated only 26% ± 14.8% of seeds for six trees with no fruiting neighbours (Figure 2). By comparison, intensity of pre-dispersal seed predation by insects was not related to either initial fruit abundance (number: \( r^2 = 0.09, F_{1,16} = 1.5, P = 0.23 \); proportion: \( r^2 < 0.01, F_{1,16} < 0.01, P = 0.97 \)), or the neighbourhood fruiting index (number: \( r^2 = 0.04, F_{1,25} = 0.90, P = 0.35 \); proportion: \( r^2 = 0.05, F_{1,25} = 1.4, P = 0.24 \)).

**Post-dispersal seed predation**

During our inspections of the experimental quadrats, we occasionally found remains of *A. graveolens* seeds in the open treatments, with bite marks indicating attack by rodents. However, we never found seed remains in the exclusion treatments, suggesting that seeds had been removed whole through the wire mesh by invertebrates, possibly ants.

The number of fruits remaining at 45 d post dispersal differed significantly between open and exclusion treatments (\( F_{1,1} = 14.2, P < 0.001 \)), with greater survival in the exclusion treatments. There was no variation between the two sites of Chamela and Cuixmala in the number of fruits remaining at 45 d post dispersal (\( F_{1,1} = 0.12, P = 0.73 \)), and treatment effects were the same at both sites with no significant interaction site × treatment (\( F_{1,1} = 0.09, P = 0.76 \)). The Kaplan–Meier probability of survival to 45 d post dispersal also did not differ between the two sites for either the open treatments (Log rank: \( \chi^2_1 = 1.4, P > 0.05 \)) or the exclusion treatments (Log rank: \( \chi^2_1 = 0.003, P > 0.05 \)). We therefore combined data from the two sites for survival probability analysis between treatments.

The Kaplan–Meier probability of survival of *A. graveolens* fruits up to 45 d post dispersal differed significantly between treatments (Log rank: \( \chi^2_1 = 109, P < 0.001 \)), with an overall survival of 0.25 for the open treatment, compared with 0.64 for the exclusion treatment (Figure 3). Hence, we estimate that insects predated 36% of the seeds of *A. graveolens* post dispersal in the exclusion treatment. By subtracting seed removal in the exclusion treatment from that in the open treatment (Kelt *et al.* 2004), we obtained an estimated 0.49 Kaplan–Meier survival probability, suggesting a potential 51% post-dispersal predation by vertebrates. The pattern of seed removal estimated for vertebrates differed significantly from that estimated for insects in the exclusion treatment (Log rank: \( \chi^2_1 = 16, P < 0.001 \)). In particular, seed survival declined more rapidly at 10 d post dispersal when subjected to potential vertebrate predation, but when subjected to removal by insects, seed survival maintained a steady rate of decline increasing slightly towards the end of the post-dispersal period (Figure 3).

**DISCUSSION**

**Pre-dispersal seed predation**

The 43% pre-dispersal seed predation of *A. graveolens* by the lilac-crowned parrot was higher than pre-dispersal predation attributed to canopy vertebrates in other studies of insect and vertebrate predation (Curran & Leighton 2000, Nakagawa *et al.* 2005), as well as that estimated for other parrot species in humid forests (Galetti & Rodrigues 1992, Howe 1980, Trivedi *et al.* 2004). High pre-dispersal seed predation by parrots has previously only been recorded in fragmented habitats where parrots
may experience food-resource limitation (Coates-Estrada et al. 1993, Francisco et al. 2008, Galetti & Rodrigues 1992), though Hauagassen (2008) reports that macaws damaged 62% of the fruit crop of Couratari guianensis in Amazonian forest. The lack of research in this area may underestimate the impact of parrots on pre-dispersal seed loss. In addition to the direct predation of seeds, the lilac-crowned parrot also discarded immature fruits below the tree, causing a total 56% loss for the tree as a result of parrot foraging activity.

Furthermore, the intensity of pre-dispersal seed predation by parrots was consistent between the two years of study for those trees that were sampled in both years. This contrasts with pre-dispersal seed predation by arboreal vertebrates in tropical moist forest, which tends to be sporadic and of short duration (Curran & Leighton 2000, Nakagawa et al. 2005). The consistent, high level of pre-dispersal seed predation by the lilac-crowned parrot may be due to the fact that *A. graveolens* fruits during the months of March to June at the end of the dry season.
when there are few food resources available for parrots (Renton 2001), and coincides with the period when parrots are raising young (Renton 2002). The seeds of *Astronium graveolens* also present a high protein and lipid content (Gutiérrez-Pérez 2005), which are important for nestling growth (Klasing 1998). Hence, in addition to providing an abundant food resource, the seeds of *A. graveolens* may also provide a valuable nutritional contribution to parrot diets.

Compared with the high pre-dispersal seed predation by parrots, insects predated only 1.3% of the seeds of *A. graveolens*. This is contrary to that found by Nakagawa *et al.* (2005) for six species of dipterocarp in the tropical moist forest of Malaysia, where pre-dispersal seed predation by insects ranged from 25% to 78%, while vertebrate predation was only 1.4% to 25%. The limited insect predation of *A. graveolens* seeds may be due to the sharp decline in both biomass and abundance of canopy insects during the dry season (Lister & Garcia-Aguayo 1992). Furthermore, the leaves of *A. graveolens* contain a volatile monoterpenic hydrocarbon which is repellent to leafcutter ants (Chen *et al.* 1984), though it is not known whether the seeds also contain deterrent secondary compounds.

Finally, 19% of the fruits collected in the fruit-traps did not contain viable seeds, and appeared to have been aborted. Seeds may be aborted as a result of insect attack or unfavourable environmental conditions (Forget *et al.* 1999), or may not have developed due to failures in pollination or resource limitation (Leimu *et al.* 2002). This reduction in effective seed-crop size of *A. graveolens* further increases the relative impact of parrot pre-dispersal seed predation.

**Fruit-crop size and resource aggregation as predictors of pre-dispersal predation**

In our study, both the number and proportion of *A. graveolens* seeds predated by the lilac-crowned parrot were significantly related to the neighbourhood fruiting index of focal trees, being greater where the resource of fruiting trees was clumped in distribution, but was not related to the initial abundance of fruits in the tree. Other studies have indicated that macaws and cockatoos forage more frequently in areas with a higher concentration of fruiting trees (Berg *et al.* 2007, Cameron & Cunningham 2006). However, the explanatory power of our neighbourhood fruiting index, which gives greater weight to the distance of fruiting trees from the focal tree, suggests that it is not only the number of fruiting trees within a given area but the distance between fruiting conspecifics which influences intensity of seed predation. Coates-Estrada *et al.* (1993) also found that total loss of fruit crop from the foraging activity of the red-fronted parrot (*Amazona autumnalis*) was negatively correlated with distance to the nearest fruiting neighbour, seed loss being highest in trees closer to a conspecific. This proximity of fruiting conspecifics may facilitate the movement of foraging parrots between trees, leading to greater foraging efficiency, rather than travelling distances between potentially isolated trees, even though they have high fruit abundance.

It may be that both the local aggregation of fruiting trees and fruit-crop size play a role in avian resource selection within a hierarchical or temporal process (Cameron & Cunningham 2006, Sallabanks 1993). However, the interplay of these two aspects of resource abundance and
dispersion in influencing pre-dispersal seed predation by large-bodied canopy granivores may vary depending on the generalist or specialist nature of the seed predator (Leimu et al. 2002), and characteristics of the habitat (Forget et al. 1999). In landscapes that do not present high concentrations of fruiting trees it may be more efficient for parrots to forage on trees with the most abundant fruit (Francisco et al. 2008). The influence of resource dispersion on the intensity of seed predation by mobile, canopy vertebrates has implications for forest dynamics. Increased seed predation by parrots in areas where fruiting trees are aggregated means that the lilac-crowned parrot may play a role in regulating local dominance of A. graveolens.

**Post-dispersal seed predation**

Considering all sources of seed loss, the 75% post-dispersal seed predation uncovered here was higher than pre-dispersal seed predation, as determined for other systems (Hulme 2002, Hulme & Benkman 2002). Nevertheless, pre-dispersal predation and waste of immature seeds by foraging parrots was the single greatest contributor to seed loss for the tree. Vertebrate post-dispersal removal of A. graveolens seeds was most probably carried out by rodents, which are the principal post-dispersal seed predators (Crawley 1992). At our study site, the spiny rat (Lyomis pictus) is the most abundant vertebrate post-dispersal seed predator (Briones-Salas et al. 2006), removing seeds that are small and easily transported, such as A. graveolens.

The role of insects was greater at the post-dispersal stage than pre-dispersal, and was most likely carried out by ants or beetles (Kelt et al. 2004, Kerley 1991, Westerman et al. 2003). Vertebrates removed seeds faster than insects, though there was a slight increase in seed removal by insects in the exclusion treatment towards the end of the post-dispersal period. Insects such as ants range over shorter distances than rodents (Anderson & MacMahon 2001); hence it may take longer for insects to locate seeds post-dispersal. There may also be an increase in insect activity towards the end of the post-dispersal period with the onset of the rainy season. Nevertheless, the fact that seed predation by insects was greater at the post-dispersal, rather than pre-dispersal stage warrants further research.

Despide the high levels of pre- and post-dispersal seed predation, trees of A. graveolens may ensure the survival of sufficient seeds for propagation by producing a high number of fruits. We estimated a mean crop size of 283 000 fruits per tree. Considering the proportion of 21% of mature fruits in the fruit-traps, we estimate about 59 000 fruits per tree may potentially be dispersed. Furthermore, the 25% survival of fruits post-dispersal in the open treatments means that of the dispersed fruits, approximately 15 000 fruits per tree have the potential to germinate. As proposed by Janzen (1967, 1969), the high synchrony in fruit production by trees of A. graveolens, as well as a high abundance of fruit per tree, may serve to satiate potential seed predators, ensuring the survival of sufficient seeds for dispersal.

**Implications for plant–animal interactions**

The intensity of pre-dispersal seed predation by the lilac-crowned parrot in tropical dry forests was within the range of the average 36–47% pre-dispersal seed predation in tropical systems, usually carried out by insects (Hulme 2002, Hulme & Benkman 2002). Hence, it may be that canopy seed predators such as parrots adopt the functional role of insects as the main pre-dispersal seed predators in dry habitats, particularly during the dry season when arthropod abundances may be low.

While the present study focuses only on one tree species, the lilac-crowned parrot consumes the seeds of a variety of other tree species during the dry season. In particular, the seeds of tree species such as Brosimum alicastrum, Pirainea mexicana, Erythrina lanata and Comocladia engleriana are predominant in the diet of the lilac-crowned parrot in the dry season (Renton 2001). The lilac-crowned parrot also consumes the seeds of a variety of tree species in the rainy season (Renton 2001), and may be an important pre-dispersal seed predator for many of these trees. Very few studies have quantified pre-dispersal seed predation by parrots, or compared this with other sources of pre- and post dispersal seed predation. Further research is needed in this area to elucidate the functional importance of parrots as pre-dispersal seed predators in tropical forests. Many parrot species of seasonally dry forests are currently threatened or endangered (Collar & Juniper 1991). As in the case of defaunation of animal herbivores (Dirzo & Miranda 1991), the loss of parrot populations in these forests may have consequences for the overall process of pre-dispersal seed predation and the recruitment of canopy trees.

**ACKNOWLEDGEMENTS**

We are grateful for logistical support provided by the Fundación Ecológica de Cuixmala, and the Estación de Biología Chamela of the Instituto de Biología, Universidad Nacional Autónoma de México. EIV-S was supported by a postgraduate grant from the Consejo Nacional de Ciencia y Tecnología. The Secretaría del Medio Ambiente y Recursos Naturales granted permits for the research. We are especially grateful to F. Dentressangle, A. C. Mont and D. Vilchis for assistance in the field, and thank the
anonymous reviewers for constructive comments on the manuscript.

LITERATURE CITED


