

Insect assemblages attacking seeds and fruits in a rainforest in Thailand

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1	Insect assemblages attacking seeds and fruits in a rainforest in Thailand
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- 44 Abstract

46	Insect seed predators are important agents of mortality for tropical trees, but little is known
47	about the impact of these herbivores in rainforests. During three years at Khao Chong
48	(KHC) in southern Thailand we reared 17,555 insects from 343.2 kg or 39,252 seeds/fruits
49	representing 357 liana and tree species. A commented list of the 243 insect species
50	identified is provided, with details about their host plants. We observed that: (1) about 43%
51	of identified species can be considered pests. Most were seed eaters, particularly on dry
52	fruits. (2) About 19% of parasitoid species (all Opiinae) for which we could determine
53	whether their primary insect host was a pest or not (all Bactrocera spp. breeding in fruits)
54	can be considered beneficials. (3) The seeds/fruits of about 28% of the plant species in this
55	forest were free of attack. Phyllanthaceae, Rubiaceae, and Meliaceae were attacked
56	relatively infrequently; in contrast, Annonaceae, Fabaceae, Sapindaceae, and
57	Myristicaceae were more heavily attacked. There was no apparent effect of plant
58	phylogeny on rates of attack but heavily attacked tree species had larger basal area in the
59	KHC plot than rarely attacked tree species. (4) Insects reared from fleshy fruits were more
60	likely to exhibit relatively stable populations compared to insects reared from dry fruits, but
61	this was not true of insects reared from dipterocarps, which appeared to have relatively
62	stable populations throughout the study period. We tentatively conclude that insects
63	feeding on seeds and fruits have little effect on observed levels of host abundance in this
64	forest.
65	

- **Key words:** dipterocarp, parasitoid, pest, seed predation, Tephritidae.

68 INTRODUCTION

69 Insect seed predators represent important agents of mortality for tropical rainforest trees because they often kill the plant embryo, or make the fruit unsuitable for seed dispersers 70 (Janzen 1970; Lewis & Gripenberg 2008). Insects feeding internally on fleshy fruits can 71 also cause significant loss of plant fitness and economic damage, via, notably, fruit 72 73 abortion (Stephenson 1981). There is an abundant literature on seed predators as pests of economic plants (e.g. Zehnder et al. 2007) or on seed- and fruit-feeding insects in 74 75 temperate areas (e.g. Turgeon et al. 1994) but in comparison little is known about these in tropical rainforests where community-level studies of insects feeding on seeds (dry fruits, 76 77 achenes) and fleshy fruits are extremely rare. This is because it is difficult to survey the extremely diverse range of potential host plants with adequate spatial and temporal 78 79 sampling effort, particularly with regard to pre-dispersal seed predation (Ctvrtecka et al. 2014). To the best of our knowledge, there are currently only six such examples which are 80 81 relatively comprehensive. (1) Janzen studies of insect seed predation in Guanacaste

 Janzen-Connell hypothesis, explaining the coexistent resulting from negative density-dependence proce Hosaka and their colleagues have studied insect seed locations in Malaysia (Nakagawa <i>et al.</i> 2003, 2005; 2017). (3) Copeland <i>et al.</i> (2009) made a broad surv Kenya, targeting tephritids (4) Ramírez and Traves 	ce of tree species in tropical forests as
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 87 2017). (3) Copeland <i>et al.</i> (2009) made a broad surv 88 Kenya, targeting tephritids (4) Ramírez and Traves 	Hosaka et al. 2009, 2011; Iku et al.,
88 Kenya targeting tephritids (4) Ramírez and Traves	ey of insects feeding on wild fruits in
20 survey of insect and produces in different behitets i	et (2010) published a comprehensive
 90 patches of forest. (5) Ctvrtecka and colleagues studi 	ied insects feeding on both seeds and
91 fruits with high sampling effort in a lowland forest o	f Papua New Guinea (Ctvrtecka <i>et al</i> .
92 2014, 2016; Sam <i>et al.</i> 2017). (6) More recently, G	Gripenberg et al. (2018, unpubl. data)
93 conducted a similar survey on Barro Colorado Island	l in Panama.
94 The present contribution adds the first study	in Thailand. We have summarized the

95 higher faunal composition of the insects reared from seeds and fruits at this location

97 contribution, we attempt to answer various questions related to three general hypotheses 98 that are particularly relevant to the identity *per se* of the plants surveyed and insect species 99 reared. First, forests may act as reservoirs of both fruit/seed-feeding pests and their 100 101 parasitoids. For example, most research on frugivorous insects from wild fruit is 102 specifically concerned with discovering the range of reservoir hosts of fruit flies (Tephritidae), which are major pests of commercial fruit crops (Allwood et al. 1999; 103 Copeland et al. 2009). Given that most insect herbivores in tropical rainforests are 104 105 reasonably host-specific (Novotny et al. 2002), it is not immediately clear whether a 106 relatively pristine forest may contribute significantly as a reservoir of pests of cultivated 107 plants, or of potential parasitoids of such pests. Further, forest pests attacking the seeds of 108 ecologically and economically important species of timber trees, such as many species of Dipterocarpaceae (Lyal & Curran 2000), may spread into plantations of these species. The 109

(Basset et al. 2018) and intend to discuss interaction networks in detail elsewhere. In this

110	forest may also potentially act as a reservoir of pests of stored products because these
111	insects usually feed on a resource low in water (Subramanyam 1995), similar to that of
112	seed predators of dry fruits (achenes; Janzen 1980). Rainforests might also act as reservoirs
113	of beneficial insects, such as parasitoids of pest species (Aluja et al. 2014). The enemy
114	hypothesis states that predatory insects and parasitoids are more effective at controlling
115	populations of herbivores in diverse systems of vegetation than in simple ones (Russell
116	1989). For example, there is evidence that diverse wet and dry forests in Mexico and
117	Central America act as reservoirs of parasitoids attacking fruit flies in fruit orchards. This
118	mechanism contributes to the value of tropical tree conservation in Mexico (Aluja et al.
119	2014).
120	Second, the identity of the plants and insects involved in interactions is crucial for
121	two reasons. The identity of plants whose levels of seed/fruit attack stand out from the rest
122	of the local vegetation (i.e. rarely or heavily attacked) is important because it can shed light
123	on patterns of insect host shifts and use (Janzen 1985) and, ultimately, to practical measures

124	of crop protection. The identity (or absence of) of the enemies of seed eaters, such as insect
125	parasitoids, is also important because some granivores and frugivores may be relatively
126	free of enemies, perhaps suggesting effective defenses. The nasty host hypothesis proposes
127	that insect herbivores feeding on plant hosts with strong and/or distinctive chemical
128	defenses may support a reduced load of parasitoids because their tissues may be more toxic
129	to parasitoids (Gauld et al. 1992). Given the potential importance of insect seed predators
130	in tropical tree mortality (Lewis & Gripenberg 2008), this hypothesis may have
131	consequences for the local distribution of tree species and the dynamics of their
132	populations.
133	Finally, seed predators are thought to be satiated by mass production of seeds,
134	which promotes escape from predation. The satiation hypothesis has been well-studied in
135	dipterocarp forests of Malaysia (Curran & Webb 2000). The whereabouts of seed-predators
136	of mast-fruiting trees, such as dipterocarps in many forests, in-between periods of masting,
137	which can be as long as several years, is crucial for these specialized insects (Hosaka et al.

138	2011). The extent of annual fluctuations of seed predators in tropical rainforests has not
139	been well-studied, with the exception of dipterocarp seed predators, which may maintain
140	populations by prolonged dormancy and/or alternative hosts (Hosaka et al. 2011). This
141	issue could help understanding patterns of insect attack on particular plant species, and
142	their local distribution and abundance. Here again the identity of both plants and insects are
143	crucial to evaluate potential patterns.
144	The general aims of this paper are to document (as far as possible) the identity of
145	insects attacking seeds and fruits, as well as their main parasitoids, in a lowland rainforest
146	in Thailand. Our specific questions are as follows:
147	
148	1) Does this forest represent a potential reservoir of pests for seed and fruit crops or seeds
149	of valuable timber trees, such as dipterocarps, in Thailand?
150	2) Does this forest represent a reservoir of parasitoids potentially able to control pests of

151 seeds and fruits in Thailand?

152 3) Which taxa of seed/fruit-feeding insects are relatively free of parasitoids?

- 4) Which tree species suffer unusual rates of seed/fruit attack in this forest? Are these tree
- 154 species particularly rare or abundant in this forest?
- 155 5) Which insect species maintain relatively high and stable populations during the study

156 years?

157

158 MATERIALS AND METHODS

159 Study site

Our study site included the 24 ha ForestGEO permanent vegetation plot (https://forestgeo.si.edu/; see below) at Khao Chong (KHC; 7° 32' N, 99° 47' E, altitude 162 120-330 m) and the surrounding forest (i.e. an area of ca. 1,500 ha). This permanent plot is 163 located in the protected lowland seasonal evergreen forest of the Khao Ban Thad Wildlife 164 Sanctuary in Southern Thailand and is described in detail by Anderson-Teixeira *et al.*

165 (2014). Mean annual rainfall is 2,665 mm and mean daily maximum air temperature is

166	27.1°C. KHC experiences a 2 to 3 months seasonal drought from January to March
167	(drought defined as any month receiving <100 mm of rainfall: Baltzer & Davies 2012). In
168	the ForestGEO plot, all trees with a diameter at breast height (DBH) of 1 cm or greater
169	have been mapped and identified to species (Anderson-Teixeira et al. 2014). There are 593
170	tree species, representing 285 tree genera and 82 plant families in the plot, with
171	approximately 300 species per ha (Baltzer & Davies 2012). The proportion of plant species
172	with dry fruits (achenes) is 26.0% and total seed rain is 7.0 dry g x m ⁻² x yr ⁻¹ (Basset <i>et al.</i>
173	2018). Although 13 dipterocarp species grow at KHC (representing 11.8% of stems and
174	23% of the basal area in the ForestGEO plot; Bunyavejchewin et al. 2011), phenological
175	studies demonstrated that the reproductive phenology of the KHC forest was more similar
176	to tropical forests with similar rainfall seasonality in other parts of the world than it was to
177	dipterocarp-dominated forests in ever wet regions of Southeast Asia (Kurten et al. 2017).
178	

179 Survey of plants and rearing of insects

180 Plant surveying and the rearing of insects from seeds and fruits are detailed in Basset *et al.*

181 (2018). Briefly, in 2013 we surveyed seeds and fruits of locally abundant tree, shrub and

- liana (more rarely herb) species. During 2014 and 2015, we restricted our sampling effort
- to 10 plant families, which represented the most common families at KHC. We refer to

these families as focal families and they included: Annonaceae, Arecaceae, Ebenaceae,

- 185 Euphorbiaceae, Fabaceae, Lauraceae, Meliaceae, Phyllanthaceae, Rubiaceae and
- 186 Sapindaceae. Unless specified, results are detailed for all host plant species. Seeds and
- 187 fruits collected on plants or freshly fallen (without apparent decomposition) were targeted,
- thus focusing on pre-dispersal attack (i.e. on insects attacking developing or mature seeds
- in the canopy of trees). Host plants were identified and their seeds/fruits assigned to the
- 190 following seed and fruit "syndromes" (hereafter seed syndromes for brevity; see Basset *et*
- 191 *al.* 2018 for more details): A1.1, fleshy drupe with thick mesocarp (>5 mm); A1.2, fleshy
- drupe with thin mesocarp (<5 mm); A2.1, non-fleshy drupe with thick mesocarp (>5 mm);

193	A2.2, non-fleshy drupe with thin mesocarp (<5 mm); B1, fleshy indehiscent fruit with
194	multiple seeds; B2, non-fleshy dehiscent fruit with multiple seeds; C1, dry winged seed
195	that does not develop in capsule; and C2, multiple dry seeds (with or without wings) that
196	develop in a capsule/pod (opening across one axis). These categories were recombined in
197	some analyses as just 'fleshy fruits' (= A1.1, A1.2, B1) or just 'dry fruits' (achenes= A2.1,
198	A2.2, B2, C1, C2).
199	Rearing sample units included clusters of conspecific seeds/fruits of similar size
200	collected from the same trees. We targeted as many individuals as possible for each plant
201	species, typically > 5 . These sample units were weighed (fresh weight) and stored in
202	individual plastic pots. Pots were lined with tissue paper and covered with very fine netting
203	for ventilation and to avoid subsequent colonization/contamination of fruits by, notably,
204	drosophilid flies (Copeland et al. 2009). Rearing pots were stored under semi-natural

206 twice weekly, and any emerging insects were collected, preserved, mounted and then

205

conditions in covered but ventilated sheds under the forest canopy. They were checked

identified (see below). Seeds/fruits were stored for 3 months, and then dissected to ensure
that there were no developing larvae inside. Seed/fruits with live larvae were reared for
longer, while other seeds/fruits were discarded.

210

211 Insect identification

212	The level of identification was unequal among insect orders owing to the availability of
213	specialists on particular insect groups. In general, beetle and moth families were identified
214	mostly to species level, whereas for Diptera and Hymenoptera only Stratiomyidae,
215	Tephritidae and Ichneumonoidea were sorted to species level. We obtained DNA
216	Cytochrome c oxidase subunit I (COI, 'DNA barcode') sequences from legs of
217	representative specimens, and we used Barcode Index Numbers (BINs) derived from insect
218	sequences to delineate species (Ratnasingham et al. 2013). Unfortunately, most of the
219	original high-quality DNA samples were spoiled in the sequencing laboratory of the
220	Smithsonian Tropical Research Institute, and in the meantime the remaining specimens

221	had been exposed to high humidity, so we were unable to obtain DNA sequences from all
222	species. Data were deposited in the Barcode of Life projects KHCSP and KHCTE (398
223	sequences). Full specimen data for specimens sequenced (including those that failed),
224	including images and host plants, are available on BOLD (www.boldsystems.org),
225	accessible from the data set KHCFRUIT using a DOI
226	(dx.doi.org/10.5883/DS-KHCFRUIT). Morphological identification of specimens, when
227	possible, was performed by RT, SEM, JWB, DLJQ, MK, PP, MS, and by colleagues cited
228	in the Acknowledgements. For Lepidoptera, nomenclature follows Holloway (2011) and
229	Holloway et al. (2001). Insect vouchers are deposited at the Thai Department of National
230	Parks, Wildlife and Plant Conservation, Bangkok, Thailand, and the National Museum of
231	Natural History, Smithsonian Institution, Washington, D.C.
232	Insects reared from seeds/fruits were assigned to a guild system at the family,
233	subfamily, or in some cases at the generic or specific level (details in Basset et al. 2018).
234	Here we only consider three guilds: seed eaters (coded as SE: larva feeding mostly on seed

235	tissue), pulp eaters (PU: larva feeding mostly on mesocarp tissue), and parasitoids (PA:
236	larva feeding on insect hosts). Members of the moth families Blastobasidae and Tineidae,
237	which are predominantly scavengers, were not included in the analyses, but when available,
238	we nevertheless provided basic information about them.
239	Assessing the pest status of insect species identified is not an easy task. For
240	Lepidoptera, we examined the list of species of economic importance compiled by Zhang
241	(1994). We further considered for pest species the number of citations occurring in the
242	Review of Applied Entomology (up to 1994) as an indication of the severity of the pest
243	(Zhang 1994). Additionally, we considered the host records of Kuroko and Lewvanich
244	(1993) for Thailand. For Tephritidae we followed the nomenclature and pest status as
245	indicated in Doorenweerd et al. (2018). The pest status of Scolytinae was inferred from
246	Browne (1961) and other sources indicated in Appendix S1, as for the rest of beetles.
247	Finally, we also considered the species listed as pests and beneficial insects in Thailand
248	(Hutacharern & Tubtim 1995).

249	Regarding the parasitoids, we considered interactions between members of the
250	Braconidae and Icheumonidae (both Ichneumonoidea) and their insect hosts; these two
251	families represented most of the parasitoids that we reared. Unlike with the host plants, our
252	interpretations of the hosts of the reared parasitoids only reflect 'high expectations of
253	interactions', not documented interactions. This is because parasitized hosts were not
254	isolated and reared individually, the parasitoids instead being reared from samples
255	including relatively high numbers of seeds and fruits. To assign putative hosts to each
256	parasitoid species, we applied three simple rules in decreasing number of importance: (1)
257	since many ichneumonoid lineages are rather conservative in host use, we followed Quicke
258	(2015) to select the most likely host order or family; (2) we then examined for each
259	parasitoid species, the co-occurrence of primary consumers in each sample from which this
260	parasitoid species was reared; and finally (3) we considered the highest abundance of
261	putative host reared in samples in which the parasitoid species was also reared. We
262	emphasize that our host assignments must not be taken as definite records (Shaw 1994).

264 Statistical analyses

265	A main host plant/insect was defined if 80% of reared individuals originated from this host.
266	Sampling effort for a particular plant species may be assessed as either the number of
267	samples obtained, or the sum of seeds collected, or the total weight of seeds. To examine
268	which plant species were rarely attacked by insects, we considered species with a high
269	number of seeds collected but none attacked (i.e. no insect reared from the seeds), as this
270	variable is more directly relevant to the regeneration of the plant species. We considered
271	the distribution of the number of seeds free of attack for each tree species, ranked in
272	decreasing number. Host species 'rarely attacked' were defined as species belonging to the
273	first quartile of this distribution. It was more challenging to define host species 'heavily
274	attacked' and for this we considered insect load on their hosts both in term of species
275	richness and abundance. With regard to insect species richness, we considered for each
276	host species, the number of insect species reared from a main host, excluding insect

277	singletons. With regard to insect abundance, we considered the number of insects reared
278	per seed (per unit seed), to reduce the effect of sampling effort, and calculated these values
279	for hosts relatively well sampled (for which > 75 seeds were collected). We compared the
280	abundance in the KHC permanent plot of rarely vs. heavily attacked tree species
281	(abundance not defined for liana species) with Mann-Whitney tests for the variables
282	Number of stems (i.e. number of individuals per tree species) and Basal area (i.e. total
283	cross-sectional area of all stems in the plot measured at breast height).
284	
201	Our analyses about insect inter-annual variation in abundance are limited by only
285	3 years of data, but motivated by the lack of data for tropical species other than those
285 286	3 years of data, but motivated by the lack of data for tropical species other than those attacking dipterocarp seeds (i.e. Nakagawa <i>et al.</i> 2003). We used the stability index of
285 286 287	3 years of data, but motivated by the lack of data for tropical species other than those attacking dipterocarp seeds (i.e. Nakagawa <i>et al.</i> 2003). We used the stability index of Wolda (1983) to estimate the magnitude of change in insect abundance between study
285 286 287 288	3 years of data, but motivated by the lack of data for tropical species other than those attacking dipterocarp seeds (i.e. Nakagawa <i>et al.</i> 2003). We used the stability index of Wolda (1983) to estimate the magnitude of change in insect abundance between study years (2013-2015). The index is calculated as the natural logarithm of the variance in the
285 286 287 288 289	3 years of data, but motivated by the lack of data for tropical species other than those attacking dipterocarp seeds (i.e. Nakagawa <i>et al.</i> 2003). We used the stability index of Wolda (1983) to estimate the magnitude of change in insect abundance between study years (2013-2015). The index is calculated as the natural logarithm of the variance in the natural logarithms of the abundances (+1) of the individuals species. We included insect

291	considered the average number of insects reared per seed among samples obtained each
292	year as a measure of insect abundance. We tested for differences in the average stability
293	index of species (a) of pulp vs. seed eaters, (b) reared from dipterocarps vs.
294	non-dipterocarps, and (c) reared from fleshy vs. dry fruits with Mann-Whitney tests. For (b)
295	and (c) we considered only insects reared from a main host, in order to relate unequivocally
296	insect species to either plant family or seed syndrome. Raw data (abundance per year) for
297	insect species are indicated in Appendix S1.
298	We evaluated the influence of host plant phylogeny on our results as follows. First,
299	we estimated the phylogenetic relationships between host species present at KHC using the
300	software package Phylomatic (Webb & Donoghue 2005; details in Basset et al. 2018).
301	Second, we tested for phylogenetic signal for all tree species attacked, for trees rarely or
302	heavily attacked, and for host trees from which Ichneumonoidea were reared. We
303	calculated the D statistic for phylogenetic signal in a binary trait (Fritz & Purvis 2010). The
304	value of the D statistic is based on the sum of changes between sister clades across the

305	phylogeny. Highly clumped traits tend to have lower D values, closer to 0. We compared
306	the scaled value of the observed D statistic to values generated under a simulated Brownian
307	model of phylogenetic structure and one resulting from no phylogenetic structure (each
308	with 10,000 permutations) using the R package 'Caper' (Orme 2013). We used a
309	complementary significance-based approach to provide further support for these results, by
310	testing for phylogenetic signal according to the mean phylogenetic distance (MPD)
311	between tree species. We used standardized effect sizes of MPD generated under null
312	models of tip label randomization (999 runs) as implemented in the R package 'Picante'
313	(Kembel <i>et al</i> . 2010).
314	
315	RESULTS
316	
317	Faunal composition and occurrence of pests and beneficial insects
318	During the three-year study, we collected 1,970 samples comprising 343.2 kg or 39,252

319 seeds/fruits from 357 liana and tree species (and a few herbs) representing 66 plant families.

From these samples we reared 17,555 insects (8,851 individuals from the 10 focal plant
families). There was a relatively high incidence of Alysiinae (Hymenoptera: Braconidae)
and a relatively low incidence of Bruchinae (Coleoptera: Chrysomelidae), Baridinae
(Coleoptera: Curculionidae), Cosmopterigidae (Lepidoptera), and Sesiidae (Lepidoptera)
in comparison with sites in Panama and Papua New Guinea (Basset et al. 2018). Appendix
S1 details the 243 species (totaling 8,949 individuals) in the guilds of seed/pulp eaters and
parasitoids that we were able to identify or morphotype. About 71% of the morphospecies
could be identified to genus and 28% of them to species. This material included mostly
beetles, with Curculionidae (53 spp. and 5,644 individuals; including 22 spp. and 4,262
individuals of Scolytinae) and Anthribidae (8 spp. and 396 individuals) predominating.
Tephritidae and Stratiomyidae represented 26 and 8 species, and 814 and 464 individuals,
respectively. Moths were dominated by Tortricidae (16 spp., 337 indivdiuals), Crambidae

- 332 (15 spp., 321 individuals) and Pyralidae (14 spp., 390 individuals), while Braconidae were
- 333 represented by 54 species and 344 individuals (Appendix S1). Most of the insects reared

334	were pulp eaters (127 spp., 73.7% of individuals), followed by seed eaters (55 spp., 22.5%)
335	and parasitoids (62 spp., 4%; Appendix S1). Among pulp eaters, two species of
336	Coccotrypes were the most abundant and reared from numerous hosts, whereas the most
337	abundant seed eater was an unidentified species of Aclees reared mostly from Mucuna
338	phaseoleae (Fabaceae). Note that the scolytines C. carpophagus, C. dactyliperda and C.
339	gedeanus may be considered as seed eaters rather than pulp eaters (Appendix S1). In
340	addition, 796 specimens of Tineidae and Blastobasidae were reared from 56 host species,
341	but the larvae of these families are more likely to be scavengers. We reared at least one
342	species of Lateantenna (Blastobasidae, L. inana (Butler, 1881)), one of Opogona
343	(Tineidae), three of <i>Phaeoses</i> (Tineidae), and one of <i>Tineovertex</i> (Tineidae).
344	Of the 69 taxa identified to species-level, 30 (43%) may be considered pests
345	(Appendix S1). This includes two ambrosia beetles that usually do not breed in seeds. The
346	insect taxa in which the proportions of reported pest species to species identified were
347	highest included: Nanophyidae (100%), Crambidae (67%), Tortricidae (55%), Scolytinae

348	(36%), and Tephritidae (26%). The origin of these pest species is summarized in Fig. 1.
349	Most pests were seed eaters, and were reared mostly from Dipterocarpaceae and from hosts
350	with seed syndromes C1 (dry winged seed) and A1.2 (fleshy drupe with thin mesocarp).
351	Most pest species and individuals were reared from dry fruits as opposed to fleshy fruits
352	(Fig. 1). Only one pest of stored products, <i>Pyralis pictalis</i> , was reared from the seeds and
353	fruits collected in the Khao Chong forest.
354	We obtained 57 samples from seven of 13 dipterocarp species growing at KHC,
355	totaling 1,240 seeds (10.3 kg; 3.1% of total seeds reared), which yielded 425 insects (14
356	samples lacked insects). Out of these, we obtained 236 weevils and moths whose
357	individual larvae likely feed on and kill a single seed (Hosaka et al. 2009). This suggests
358	that about 19% of dipterocarp seeds were lost to weevils and moths. Insects reared from
359	dipterocarp seeds included at least 26 species of seed and pulp eaters (Appendix S2),
360	mostly belonging to the Curculionidae, Nanophyidae and Tortricidae. The most abundant
361	species were an unidentified species of Alcidodes (Curculionidae) reared from Parashorea

stellata, and Andrioplecta shoreae reared from four dipterocarp hosts. In comparison 362 Nakagawa et al. (2003) reared 1,419 insects representing 51 species from 20,215 seeds of 363 364 24 dipterocarp species in Sarawak. Only four species were in common between their study 365 and ours (Appendix S2). In Pasoh, Malaysia, Hosaka et al. (2009) recorded at least 32 insect species from two consecutive mast-fruiting events of 15 species of dipterocarps 366 (3,779 insects reared from 27,483 seeds). Senthilkumar et al. (2009) studied seed predation 367 in Dipterocarpus retusa in Assam, India, and recorded nine species of seed predators. In 368 Thailand, at least 12 species of seed predators have been recorded from dipterocarps 369 370 (Hutacharern & Tubtim 1995; DNP 2018). Because of incomplete identifications, different 371 taxonomists studying the insect material and inconsistent use of DNA barcoding, it is 372 difficult to compare the lists of taxa provided by these dipterocarp studies. Nonetheless, 373 they suggest a relatively low overlap with the fauna feeding on dipterocarp seeds at KHC. The densities of reared insect individuals per dipterocarp seed appears to be higher at Khao 374 Chong during the study period (0.34 insect per seed) compared with Lambir Hills (0.07 375

376	insect per seed; Nakagawa et al., 2003) or Pasoh (0.14 insect per seed; Hosaka et al. 2009),
377	during periods of mast fruiting. One species of Blastobasidae and two species of Tineidae
378	were reared from Dipterocarpaceae at KHC.
379	Of 27 parasitoid species for which we could identify the main insect hosts and
380	verify whether the host was considered a pest of fruits or seeds, 5 species (18.5%) could be
381	considered beneficial (Appendix S1). All these species were Opiinae attacking Bactrocera
382	pests (Tephritidae) breeding in the fruits of many host plant species. In addition, the larvae
383	of Hermetia illucens recycle manure, so this species can be also considered beneficial
384	(Appendix S1).
385	
386	Levels of parasitism of insects attacking seeds and fruits
387	Our data allowed us to present only crude estimates of the level of parasitism due to

- 388 Ichneumonoidea (mostly Braconidae, Appendix S1 and Table 1). Overall, about 8.2% and
- 389 2.9% of insect species and individuals were parasitized, respectively. The level of

390 parasitism was not notably different between pulp and seed eaters (Fisher exact test, p = 391 0.483). Tephritidae was the most commonly attacked family by Braconidae, followed by 392 Curculionidae (not including Scolytinae). Bactrocera irvingiae and Andrioplecta shoreae, reared from several host plant species, appeared to be the species most commonly 393 parasitized by braconids. Insect taxa that appeared to be infrequently parasitized (Table 1: 394 395 Stratiomyidae, Pyralidae, Crambidae, Scolytinae) may be under attack by parasitoids other than braconids. For example, *Coccotrypes* spp. (Scolytinae) are known to be attacked by 396 the braconid genera Spathius, Bracon and Diospilus (Quicke, 2015). These genera were 397 398 infrequently reared at Khao Chong and obtained from other putative hosts. We also note 399 that there was no obvious correlation between the number of species of parasitoids and 400 prey reared from particular plant families (only main hosts considered: Spearman rank 401 correlation, $r_s = 0.112$, p > 0.25, n = 31 plant families). Finally, most species of parasitoids were reared from main host plant species with syndrome A1.2 (40.9% of species), B1 402 (25.0%) and A2.2 (18.2%). 403

Rates of seed attack

406	Of 357 plant species surveyed, seeds/fruits of 101 were free of attack (28.3%). The first
407	quartile of the distribution of these species represented 71% of the total number of seeds
408	not attacked. Antidesma neurocarpum (Phyllanthaceae) was the most avoided plant species,
409	with 344 seeds not attacked (Fig. 2). Other tree species rarely attacked (first quartile of the
410	distribution in Fig. 2) included 11 Rubiaceae, 9 Annonaceae, 9 Arecaceae, 7 Meliaceae,
411	and 6 Euphorbiaceae, Fabaceae and Phyllanthaceae each. Plant families with a high
412	proportion of seeds not attacked (> 15%) included Apocynaceae, Clusiaceae, Meliaceae,
413	Anacardiaceae, Rubiaceae, Celastraceae, Phyllanthaceae, Sapotaceae (Fig. 2). Of those,
414	Phyllanthaceae, Rubiaceae and Meliaceae were species-rich and collected with a high
415	sampling effort, and hence, may be considered as families relatively infrequently attacked
416	by insects. Seed syndrome B2 (non-fleshy) also had a relatively high proportion of seeds
417	free of attack (Fig. 2).

418	The main hosts of insects at KHC (as defined in the methods) belonged to 40
419	species and 16 plant families. Only Parashorea stellata (Dipterocarpaceae) and
420	Lepisanthes rubinigosa (Sapindaceae) supported more than two insect species.
421	Dipterocarpaceae, Annonaceae and Fabaceae had a relatively high load of insect species,
422	as well as seed syndromes B1, A1.2 and C1, a mixture of dry and fleshy fruits (Fig. 3a).
423	The 25 most heavily attacked host species (in terms of insect abundance) often belonged to
424	Annonaceae, Fabaceae, Sapindaceae and Myristicaceae (Fig. 3b). The highest numbers of
425	insect reared were obtained from Mezzettia parviflora (Annonaceae). Overall densities of
426	insects were also relatively high on Meliaceae and Anacardiaceae (Fig. 3c). On average the
427	highest densities of insect reared per seed and plant species were obtained from hosts with
428	Syndrome C2 (multiple dry seeds). There was no significant difference between the
429	number of stems in the plot of tree species rarely and heavily attacked (Mann-Whitney U =
430	192.5, $p = 0.808$). However heavily attacked tree species had significantly larger basal
431	areas in the plot than rarely attacked tree species (U= 309.0, p < 0.001; mean \pm s.e. = 6.08

432 $m^{-2} \pm 1.145$ and 1.28 $m^{-2} \pm 0.439$, respectively).

433

434 **Insect fluctuation during study years**

Overall the highest densities per unit seed over the three-year study were attained by 435 several species of Scolytinae (Appendix S1). There was no significant difference between 436 437 the average stability index of pulp-eating species and that of seed-eating species (Mann-Whitney test, U = 1481.5, p = 0.927). However, the average stability index of 438 species reared from dipterocarp hosts was significantly smaller (more stable) than that of 439 species reared from non-dipterocarp hosts (U= 710.0, p = 0.027; Fig. 4a). Further, the 440 average stability index of species reared from fleshy fruits was significantly smaller (more 441 stable) than that of species reared from dry fruits (U=313.0, p=0.010; Fig. 4b). 442

443

444 Host plant phylogenetic signals

445 Fig. 5 provides a visual interpretation of how all/rarely/heavily attacked plant species, and

446	from which Ichneumonoidea were reared, clustered across the whole plant phylogeny at
447	KHC. The three first categories showed a limited phylogenetic signal with the D statistic
448	relatively high (all plant species attacked: D=0.862, p(D>0)=0.0001, p(D<1)=0.0001;
449	species rarely attacked: D=0.781, p(D>0)=0.005, p(D<1)=0.0023; species heavily attacked:
450	D=0.855, p(D>0)=0.025, p(D<1)=0.0001). For plant species hosting Ichneumonoidea,
451	there was clearly no phylogenetic signal (D= 0.994, p(D>0)=0.418, p(D<1)=0.0001).
452	Significance tests of phylogenetic signal according to MPD indicated that all categories
453	were not clumped across plant phylogeny (all species: MPD observed = 358.9, MPD
454	random mean = 342.5, p = 0.92; species rarely attacked: MPD observed = 305.3, MPD
455	random mean = 328.1, p = 0.23; species heavily attacked: MPD observed = 364.8, MPD
456	random mean = 329.6, $p = 0.88$; species hosting Icheumonoidea: MPD observed = 355.9,
457	MPD random mean = 330.3 , p = 0.78).
458	
459	DISCUSSION

461	Insect assemblages feeding on seeds and fruits in tropical rainforests are challenging to
462	study, primarily because of low rates of attack, high plant diversity, and the high sampling
463	effort required to rear sufficient numbers of insect specimens to provide meaningful
464	statistics (Ctvrtecka et al. 2014). Further, the taxonomic knowledge of insects reared from
465	native seeds and fruits of tropical countries is often limited (Nakagawa et al. 2003; Miller
466	et al. 2014). Regarding the questions asked in this study, we observed that (1) about 43% of
467	species identified could be considered pests. Most were seed eaters, particularly on dry
468	fruits (but only a single pest of stored products was recorded), belonging to Nanophyidae,
469	Tortricidae, Crambidae, Scolytinae and Tephritidae. (2) About 19% of parasitoid species
470	for which we could assess whether the main insect host is a pest could be considered
471	beneficial. All these species were Opiinae with Bactrocera pests breeding in fruits as main
472	hosts. (3) Overall about 8% of insect species reared from seeds/fruits were parasitized by
473	Ichneumonoidea, with Tephritidae being the family most commonly attacked. (4) The
474	seeds/fruits of about 28% of plant species in the KHC forest were free of attack. The

475	seeds/fruits of Phyllanthaceae, Rubiaceae and Meliaceae were attacked relatively
476	infrequently by insects. In contrast, fruits and seeds of species of Annonaceae, Fabaceae,
477	Sapindaceae and Myristicaceae were more likely to be heavily attacked, with multiple dry
478	seeds (Syndrome C2) often well attacked. There was no apparent effect of plant phylogeny
479	on rates of attack but heavily attacked tree species had larger basal area in the KHC plot
480	than rarely attacked tree species. (5) The highest densities per unit seed over the three study
481	years were attained by several species of Scolytinae, as these beetles may produce large
482	brood inside fruits. Insects reared from fleshy fruits were more likely to exhibit relatively
483	stable populations compared to insects reared from dry fruits, except for insects reared
484	from dipterocarps, which appeared to have relatively stable populations during the study
485	years at KHC.
486	The proportion of pest species recorded in our study is probably inflated because

488 reared from native seeds and fruits (Miller et al. 2014). We encountered two general

487

in the tropics insect pests are far better known than native forest insects, especially those

categories of pests: (1) various beetles species breeding in the dry seeds of dipterocarps that 489 appear to be rather specific (Nakagawa et al. 2003) and (2) polyphagous species of 490 491 Tephritidae breeding in fleshy fruits. The former species could be of concern because modest dipterocarp plantations have been established in Thailand since the 1980s 492 (Weinland 1998). However, densities of the most common pest feeding on dipterocarps, 493 Alcidodes sp. 15, were rather low, reaching 0.16 insect per seed on average during the 494 three-year study. Bactrocera irvingiae was the most commonly reared tephritid from fleshy 495 fruits, but this species is not considered a pest. Dacus longicornis, a pest of Cucurbitaceae, 496 497 reached densities of 0.44 flies per fruit on our focal hosts, but was not very abundant when 498 all plant species surveyed were considered. We conclude that during our study years the 499 KHC forest did not support insect pests in densities that may cause concern to timber 500 species (dipterocarps) or fruit crops. Less than 20% of parasitoid species appeared to have insect pests as hosts. Since we have little evidence that the KHC forest acts as a reservoir of 501 insect seed/fruit pests, it is difficult to argue that the same forest acts as a reservoir of 502

beneficial insect species. A better test of this issue would be to compare parasitoid and seed 503 504 insect assemblages in commercial crops contiguous with natural forests, such as in Mexico 505 (Aluja et al. 2014). However, in Thailand such situations are rare, with habitats contiguous to natural forests represented primarily by buffalo fields, maize plantations, or holiday 506 resorts (DJ Quicke, pers. obs.). 507 A more interesting question related to parasitoids is whether some seed insects 508 may be relatively free of ichneumonoid parasitoids. In Costa Rica, Janzen (1980) observed 509 that Bruchinae seed predators are rarely attacked by parasitoids. At KHC Bruchinae are 510 replaced by Anthribidae and Curculionidae (Basset et al. 2018), whose species frequently 511 512 were attacked (except for Scolytinae, Table 1). Further, many of the Tephritidae species were attacked by braconids. We reared about 50% fewer individuals of Stratiomyidae 513 514 (Appendix S1) but did not record any braconid attacks on these flies. There are very few Ichneumonoidea parasitoids of Stratiomyidae (Quicke 2015), which are attacked only as 515 eggs by various Chalcididae and Trichogrammatidae (Robertson 1987). We also note that 516

517	there was no obvious correlation (negative or positive) between the number of prey and
518	parasitoids reared from particular plant families, and that there was no phylogenetic signal
519	relating host plant species from which Ichneumonoidea were reared. Although these
520	represent weak tests of the nasty host hypothesis (Gauld et al. 1992), these observations do
521	not appear to support it (and see Quicke 2012 for other considerations). Our rearing scheme,
522	albeit imperfect to obtain reliable data about the identity of parasitoid hosts and level of
523	parasitism, nevertheless suggests that the action of parasitoids at KHC may be too
524	infrequent to induce strong differences in seed/fruit crops, with possible consequences on
525	local tree abundance.
526	There are certainly different reasons for seeds of particular plant species to be
527	attacked less frequently by insects. First, plant chemistry may be an important determinant;
528	because seeds represent the most valuable part of the plant, they are usually well protected
529	(Janzen 1969; Ramírez & Traveset 2010). At present we lack data for most KHC plant
530	species to provide a context for discussing plant chemistry (see Gripenberg et al. 2018 for

531	such a discussion). Our phylogenetic tests indicated only limited phylogenetic signal for
532	the categories of plant species attacked by seed and pulp eaters, as well as for plant species
533	rarely of heavily attacked. This suggests that insects overall may not be very selective
534	regarding attacking or avoiding particular clades of plant species, even if they may be
535	reasonably host specific. Second, sample size is certainly important (Ctvrtecka et al. 2014),
536	but among our focal plant families, we could nevertheless crudely assign species to the
537	categories rarely and heavily attacked. The next important variable is probably local host
538	abundance. We found that host species heavily attacked have on average a higher basal area
539	(but not number of stems) in the KHC plot than rarely attacked host species. This suggests
540	that seed and pulp eaters are influenced primarily by seed/fruit production, which is probably more dependent on basel area than on number of stems. It seems less likely that
542	seed and pulp eaters are directly limiting the local abundance of heavily attacked tree
543	species.
	-

544 The observations that dipterocarp mast fruiting does not occur at Khao Chong

(Kurten et al. 2017), and insect densities in dipterocarp seeds during the study years were 545 higher than in Malaysian dipterocarp forests experiencing mast fruiting (Nakagawa et al. 546 547 2003; Hosaka et al. 2009), support the hypothesis of satiation of seed predators by mast fruiting (Curran & Webb 2000). However, it is not clear why insects reared from 548 dipterocarp seeds at KHC should have more stable populations than insects reared from 549 non-dipterocarp hosts. This may be related to easy host-switching and alternative hosts for 550 insects feeding on dipterocarp seeds (Nakagawa et al. 2003). The low faunal turnover 551 between dipterocarp insects at Khao Chong and in Malaysia is also of interest, suggesting 552 that different insect assemblages may be well adapted to either mast-fruiting events or the 553 554 lack of these events. We also strongly suspect that low host specificity in insects breeding 555 in fleshy fruits may explain the more stable populations of these species as opposed to 556 those breeding in dry fruits. This issue will be explored elsewhere with more adequate data. In conclusion, most of the evidence (often indirect) suggests that insects feeding 557

on seeds and fruits at Khao Chong have a limited impact on host abundance in this forest.

559	Insect densities were low, as was the number of confirmed insect pests, and heavily
560	attacked tree species were not notably less abundant than other species. This situation
561	appears similar to that described for a lowland rainforest in Papua New Guinea (Ctvrtecka
562	et al. 2014; Sam et al. 2017). This could be a consequence of the high plant diversity at
563	these two locations, but it also may be related to the relative occurrence of fleshy vs. dry
564	fruits (Basset et al. 2018). It is obvious that more surveys of insects feeding on seeds and
565	fruits are required at different rainforest locations to discuss adequately this issue.
566	
567	
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732 SUPPORTING INFORMATION

733	Additional Supporting Information may be found online in the Supporting Information
734	section at the end of the article.
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736	Appendix S1. Details of insects reared from seeds and fruits and Khao Chong.
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738	Appendix S2. Seed and pulp eaters reared from Dipterocarpaceae at Khao Chong.
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742 Figure legends

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744 Figure 1 Source of pest species recorded at Khao Chong, detailed for species and 745 individuals and by (a) insect families, (b) insect guilds, (c) main host family and (d) main 746 host seed syndromes. Curculionoidea do not include Scolytinae, which are indicated 747 748 separately. 749 750 Figure 2 Plant species free of seed attack. (a) Inset: full distribution of the number of seeds free of attack for each species not attacked; main figure: first quartile of the distribution 751 752 with name of species detailed and plant families abbreviated and colored similarly. (b) Proportion of seeds free of attack (black) detailed by plant family (when no. of plant 753 species surveyed \geq 3). (c) Same, detailed by seed syndrome. Abbreviations of plant 754 families: An=Anacardiaceae; Ao=Annonaceae; Ap=Apocynaceae; Ar=Arecaceae; 755 756 Cl=Clusiaceae; Er=Erythroxylaceae; Eu=Euphorbiaceae; Ge=Gentianaceae;

758 Ph=Phyllanthaceae; Po=Poaceae; Ru=Rubiaceae; Un=Unknown.

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760 Figure 3 Heavily attacked plant species. (a) Number of insect species (white = pulp eaters, black = seed eaters) reared from main hosts (as defined in methods), detailed by plant 761 families. Inset: same presentation, detailed by seed syndrome. (b) Number of insects reared 762 per seed for the 25 most attacked plant species. Black = seed eaters, white or different 763 colour = pulp eaters (same colour denotes same plant family). (c) Average number of 764 765 insects reared per seed and plant species, detailed by plant family (white = pulp eaters, 766 black = seed eaters). (d) Same presentation, detailed by seed syndrome. Abbreviations of 767 families for (b): An=Anacardiaceae; Ao=Annonaceae; Ar=Arecaceae; 768 Ch=Chrysobalanaceae; Di=Dilleniaceae; Di=Dipterocarpaceae; Eb=Ebenaceae; Eu=Euphorbiaceae; Fa=Fabaceae; La=Lauraceae; Me=Meliaceae; Mo=Moraceae; 769 My=Myristicaceae; Ru=Rubiaceae; Sa=Sapindaceae. 770

Mo=Moraceae;

772	Figure 4 Insect species ranked by their stability index. (a) Species reared from
773	non-dipterocarp hosts (grey bars) vs. species reared from dipterocarp hosts (black bars). (b)
774	Species reared from fleshy fruits (grey bars) vs. dry fruits (black bars).
775	
776	Figure 5 Maximum clade credibility consensus trees depicting the phylogenetic
777	relationships between 622 host plant species, with for each consensus tree, taxa marked in
778	red indicate (a) all species attacked, (b) species rarely attacked, (c) species heavily attacked
779	and (d) species from which Ichneumonoidea were reared.





(c) main host family and (d) main host seed syndromes. Curculionoidea do not include Scolytinae, which are indicated separately.



Figure 2 Plant species free of seed attack. (a) Inset: full distribution of the number of seeds free of attack for each species not attacked; main figure: first quartile of the distribution with name of species detailed and plant families abbreviated and colored similarly. (b)

- Proportion of seeds free of attack (black) detailed by plant family (when no. of plant species surveyed \geq 3). (c) Same, detailed by seed
- 794 syndrome. Abbreviations of plant families: An=Anacardiaceae; Ao=Annonaceae; Ap=Apocynaceae; Ar=Arecaceae; Cl=Clusiaceae;
- 795 Er=Erythroxylaceae; Eu=Euphorbiaceae; Ge=Gentianaceae; La=Lauraceae; Lo=Loganiaceae; Ly=Lythraceae; Me=Meliaceae;
- 796 Mo=Moraceae; Ph=Phyllanthaceae; Po=Poaceae; Ru=Rubiaceae; Un=Unknown.
- 797



Figure 3 Heavily attacked plant species. (a) Number of insect species (white = pulp eaters, black = seed eaters) reared from main hosts (as defined in methods), detailed by plant families. Inset: same presentation, detailed by seed syndrome. (b) Number of insects reared per seed for the 25 most attacked plant species. Black = seed eaters, white or different colour = pulp eaters (same colour denotes same plant family). (c) Average number of insects reared per seed and plant species, detailed by plant family (white = pulp eaters, black = seed eaters). (d) Same presentation, detailed by seed syndrome. Abbreviations of families for (b): An=Anacardiaceae; Ao=Annonaceae; Ar=Arecaceae; Ch=Chrysobalanaceae; Di=Dilleniaceae; Di=Dipterocarpaceae; Eb=Ebenaceae; Eu=Euphorbiaceae; Fa=Fabaceae; La=Lauraceae;

804 Me=Meliaceae; Mo=Moraceae; My=Myristicaceae; Ru=Rubiaceae; Sa=Sapindaceae.



Figure 4 Insect species ranked by their stability index. (a) Species reared from non-dipterocarp hosts (grey bars) vs. species reared from

807 dipterocarp hosts (black bars). (b) Species reared from fleshy fruits (grey bars) vs. dry fruits (black bars).





Figure 5 Maximum clade credibility consensus trees depicting the phylogenetic relationships between 622 host plant species, with for each consensus tree, taxa marked in red indicate (a) all species attacked, (b) species rarely attacked, (c) species heavily attacked and (d) species from which Ichneumonoidea were reared.

817	from seeds and fruits	at Khao Chong	, presented in	decreasing %	of species	parasited.
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Taxa	No. spp.	No. spp.	% species	% individuals
	reared	parasitized	parasitized	parasitized
Tephritidae	26	7	26.9	7.0
Anthribidae	8	1	12.5	0.3
Curculionidae *	26	3	11.5	0.8
Tortricidae	13	1	7.7	4.6
Stratiomyidae	8	0	0	0
Pyralidae	8	0	0	0
Crambidae	15	0	0	0
Scolytinae	22	0	0	0
All pulp eaters	113	8	7.1	6.3
All seed eaters	34	4	11.8	0.7
All	147	12	8.2	2.9

819 * Without Scolytinae