

1 **Mangroves are an overlooked hotspot of insect diversity despite low plant diversity**

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20

21 **Abstract**

22 We here compare the tropical arthropod fauna across a freshwater swamp and six different
23 forest types (rain-, swamp, dry-coastal, urban, freshwater swamp, mangroves) based on
24 140,000 specimens belonging to ca. 8,500 species. Surprisingly, we find that mangroves, a
25 globally imperiled habitat that had been expected to be species-poor for insects, are an
26 overlooked hotspot for insect diversity despite having low plant diversity. Mangroves are very
27 species-rich (>3,000 species) and distinct (>50% of species are mangrove-specific) with high
28 species turnover across Southeast and East Asia. Overall, plant diversity is a good predictor
29 for insect diversity for most habitats, but mangroves compensate for the low number of
30 phytophagous and fungivorous species by supporting an unusually rich community of
31 predators whose larvae feed in the productive mudflats. For the remaining habitats, the
32 insect communities have diversity patterns that are largely congruent across guilds. The
33 discovery of such a sizeable and distinct insect fauna in a globally threatened habitat
34 underlines how little is known about global insect biodiversity.

35 **Keywords**

36 Insect biodiversity, Mangroves, NGS barcoding, species discovery, beta-diversity, global
37 insect decline, Southeast Asia

38

39 **Introduction**

40 Insects are currently experiencing anthropogenic biodiversity meltdowns with
41 declines having attracted much attention[1–4] and controversy[5–10]. The controversy is
42 largely due to the paucity of high-quality data for arthropods, which is also responsible for
43 imprecise estimates of global animal species richness[11,12] and understanding species
44 turnovers[13–15]. These knowledge gaps are also likely to threaten the health of whole
45 ecosystems given that arthropods provide a large number of important ecosystem
46 services[3,16–19], contribute much of the animal biomass[20] and are yet frequently ignored
47 in habitat assessments. The lack of baseline data is particularly worrisome at a time when
48 tropical ecosystems are heavily impacted by habitat conversion and global change[21].

49 The situation is particularly dire for the species-rich tropics, for which so few
50 comprehensive surveys have been conducted[22–24] that only three of the 73 studies in a
51 recent review of insect declines involved tropical sites[8]. Furthermore, tropical insect
52 surveys have traditionally focused on tropical rainforests[24], with other tropical habitats
53 being largely neglected. Mangrove forests are a prime example of a tropical habitat for which
54 the insect fauna is poorly characterized. Mangroves used to cover more than 200,000 km² of
55 the global coastline[25], but have been experiencing an annual area loss of 1-2%[25,26].
56 Indeed, the losses of mangroves far exceed those of more high-profile ecosystems such as
57 rainforests and coral reefs[26]. Unfortunately, these losses are further exacerbated by
58 climate change[27], with some simulations predicting a further reduction by 46–59% for all
59 global coastal wetlands by the year 2100[28]. This is a particularly worrying trend as
60 mangrove ecosystems have been found to be sequester more carbon per hectare than
61 tropical dryland forests[29]. These changes will not only endanger entire ecosystems that
62 provide essential ecosystem services[30–32], but also threaten the survival of numerous
63 mangrove species with unique adaptations. Mangrove specialists with such adaptations are
64 well known for vertebrates and vascular plants[33,34], but the invertebrate diversity is largely
65 unknown.

66 One reason why the mangrove insect fauna is likely to have received little attention is
67 the low plant diversity in mangroves. Tropical arthropod diversity is usually positively
68 correlated with plant diversity[23,24,35] which implied that mangroves would provide few
69 insights into understanding whether insect herbivores drive high plant diversity in the tropics
70 [36–38] or high plant diversity was responsible for high insect diversity [22,39]. Arguably, the
71 traditional focus on addressing this question had the undesirable side-effect that the insect
72 fauna of habitats with low plant diversity received comparatively little interest. Yet, many of
73 these habitats are threatened with destruction, with mangroves being a good example. The
74 few existing studies of mangrove insects focused on specific taxa[40–42], only identified
75 specimens to higher taxonomic levels[43–45], and/or lacked quantitative comparison with
76 the insect fauna of adjacent habitats. Given these shortcomings, these studies yielded
77 conflicting results[44,46,47] with some arguing that high salinity and/or low plant
78 diversity[33,44,46] were responsible for a comparatively poor insect fauna, while others
79 found high levels of species diversity and specialization[47].

80 Here, we present the results of a comprehensive study of species richness and
81 turnover of arthropods across multiple tropical habitats. The assessment is based
82 on >140,000 specimens collected over >4 years from mangroves, rainforests, swamp forests,
83 disturbed secondary urban forests, dry coastal forests, and freshwater swamps in Singapore
84 (Fig. S1). In addition, we assess the species richness and turnover of mangrove insects
85 across East and Southeast Asia by including samples from Brunei, Thailand, and Hong
86 Kong. Specifically, our study (1) estimates mangrove insect diversity, (2) evaluates the
87 distinctness in reference to five different forest habitats, (3) analyzes the biodiversity patterns
88 by ecological guild, and (4) determines species turnover across larger geographic scales.
89 Most of the work was carried out in Singapore because it has a large variety of different
90 habitats that occur within 40km on a small island (724 km²) that lacks major physical barriers.
91 In addition, all habitats have experienced similar levels of habitat degradation or loss (>95%

92 overall loss of original vegetation cover[48]; ca. 90% loss of rainforest[49]; ca. 93% loss of
93 swamp forest[50]; 91% loss for mangroves[51]).

94 A thorough assessment of insect biodiversity requires dense sampling over an
95 extended period of time[52–54]. We sampled 107 sites using Malaise traps and
96 subsequently processed specimens for 16 arthropod orders (Fig. S2) typically found in
97 Malaise traps. The samples were typical in that Diptera and Hymenoptera comprised >75%
98 of all specimens (Fig. S2) and these orders were therefore subsampled by taxon and
99 ecological guild (Table S2). More than 140,000 specimens were NGS-barcoded[55] and
100 grouped into putative species, which allowed for species richness and abundance
101 estimates[56–58]. Contrary to expectations, we demonstrate that mangrove forests have a
102 very distinct and rich insect fauna. In addition, the species turnover for all habitats in
103 Singapore and the different mangrove sites in Asia is very high.

104 **Results**

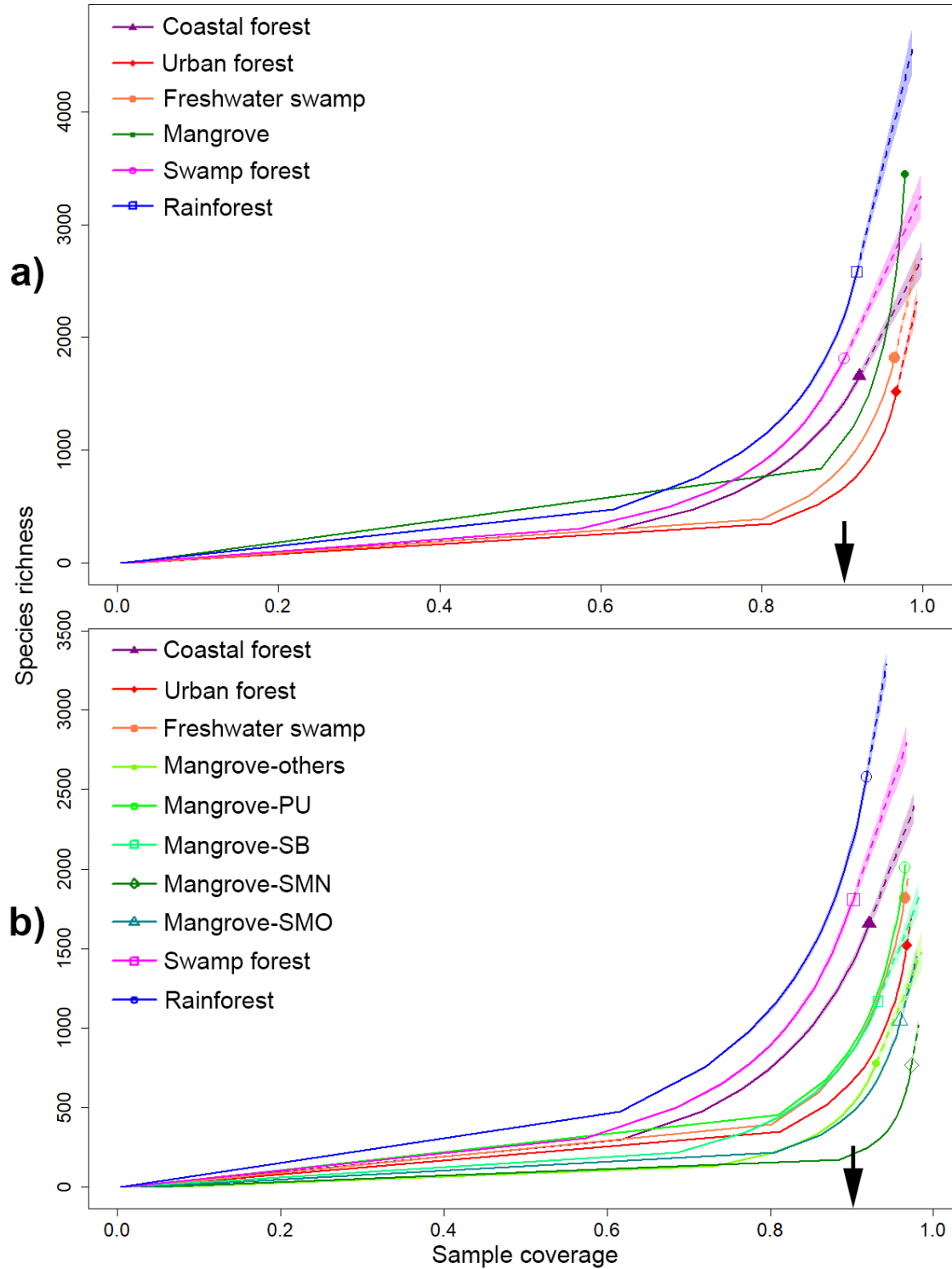
105 *Species delimitation based on NGS barcodes*

106 We obtained 143,807 313-bp *cox1* barcodes, which were grouped into 8256–8903
107 molecular operationally taxonomic units (mOTUs, henceforth referred to as species) using
108 objective clustering[59] at different p-distance thresholds (2–4%; Table S5). An alternative
109 species delimitation algorithm, USEARCH[60], yielded similar species richness estimates of
110 8520–9315 species using the identity (--id) parameters 0.96–0.98. Most species boundaries
111 were stable, with species numbers only varying by <12% across species delimitation
112 techniques and parameters. We hence used the species generated via objective clustering
113 at 3% p-distance for the analyses (see supplementary data Fig. S3 for results obtained with
114 2% and 4%).

115 *Alpha-diversity across habitats*

116 We rarefied the species richness curves by sample coverage[61] (Fig. 1) for each
117 habitat, as well as by the number of specimens processed (Fig. S3). In addition, we only

118 included trapping sites that had at least 100 barcoded individuals to prevent poorly-sampled
119 sites from artificially inflating site dissimilarity. Alpha-diversity comparisons were made at the
120 rarefaction point with the lowest coverage/number of specimens (i.e., swamp forest in Fig. 1,
121 *top*). Our initial analysis compared the Alpha-diversity of rainforest, swamp forest, urban
122 forest, freshwater swamp and coastal forest habitats and mangroves with all sites being
123 grouped as a single habitat type. The species diversity of mangroves (1102.5 ± 10.8 species)
124 is ca. 50-60% of the rarefied species richness of adjacent tropical primary/secondary forest
125 (2188.4 ± 42.6 species) and swamp forest sites (1809 species) (Fig. 1a), but a site-specific
126 analysis also revealed that two of the major mangrove sites in the study (PU & SB) have
127 similar species richness as the freshwater swamp site after rarefaction (Fig. 1b). The species
128 richness of a third mangrove site (SMO) was lower and more similar to the richness of an
129 urban forest site. A newly regenerated mangrove (SMN), adjacent to an old-growth
130 mangrove (SMO) had much lower species richness.



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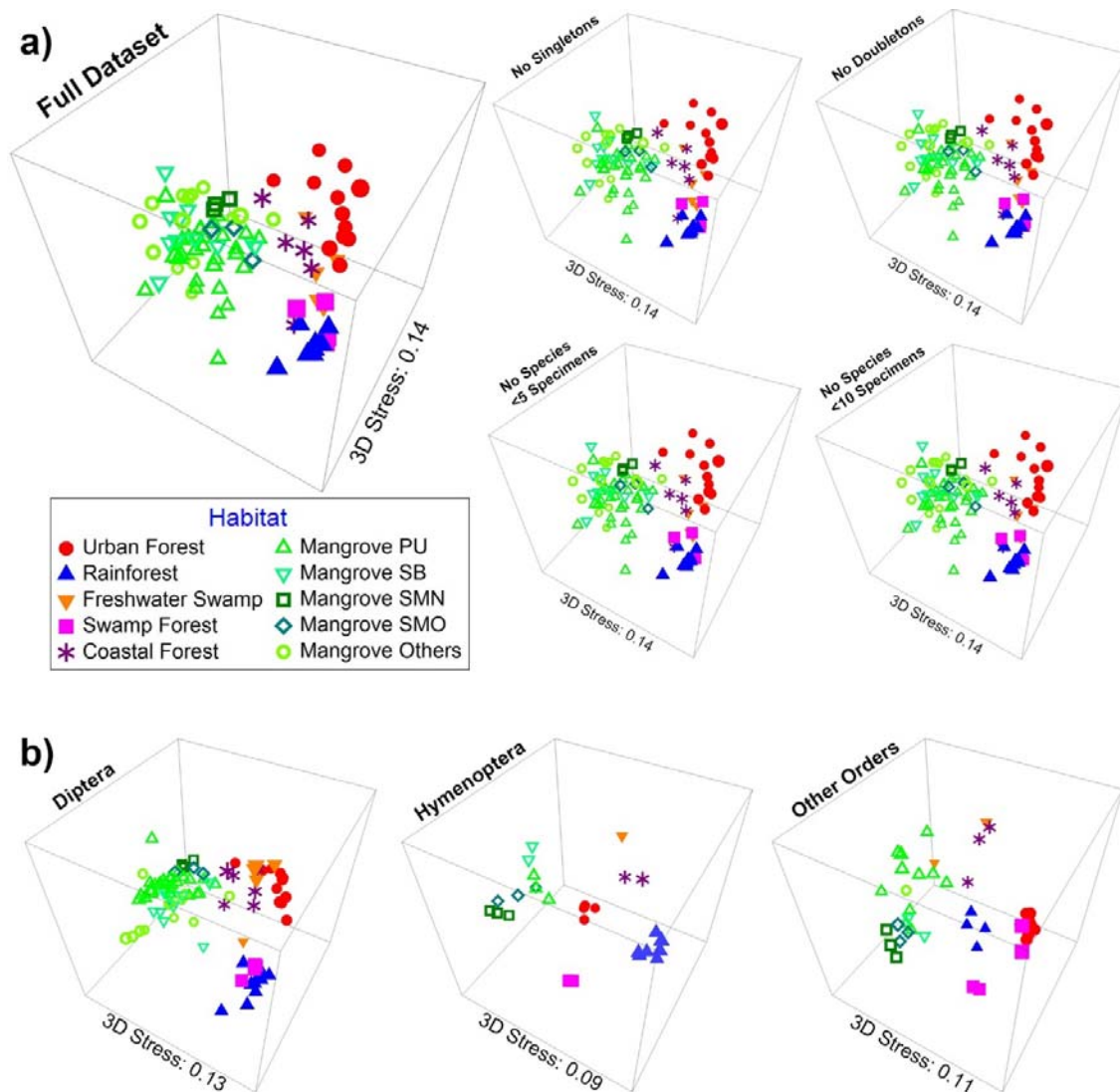
132 **Figure 1.** Insect alpha-diversity across tropical forest habitats. (a) Mangroves treated as one
133 habitat; (b) Comparison of mangrove sites: Pulau Ubin (PU), Sungei Buloh (SB), Pulau
134 Semakau old-growth (SMO), Pulau Semakau new-growth (SMN); solid lines = rarefaction;
135 dotted = extrapolations. The arrow on the x-axis indicates the point of rarefaction at which
136 species richness comparisons were made.

137 *Species turnover across habitats*

138 Mangrove arthropod communities are very distinct from those of the other habitats,
139 with the communities from most habitats being well separated on NMDS plots (Fig. 2) even
140 though several mangrove sites (PU, SB, SM) are geographically further from each other
141 (>30 km) than from the other habitat types (Fig. S1). These patterns are also observed when
142 the data are split into three taxon sets: (1) Diptera, (2) Hymenoptera, (3) remaining arthropod
143 taxa (Fig. 2b). These results are also robust to the removal of rare species (Fig. 2a). Only 48
144 (0.6%) of the 8572 putative species in the species turnover analysis are found in all habitat
145 types while 5989 (69.9%) are only in a single type (Table S6); within the mangroves, 50.2%
146 of the 3557 species are only known from the mangrove habitat. The habitat type the
147 mangroves share the most species with is the coastal forest (873 of 3557 species, 24.5%).
148 When rare species are removed (<10 specimens), 481 of the remaining 1773 species
149 (27.1%) are found in a single habitat while only 48 (2.7%) are found in all (Table S6); i.e.,
150 even after excluding rare species, a large proportion of the insect communities are putative
151 habitat specialists.

152 Dissimilarity of the habitat-specific communities was confirmed with ANOSIM tests (Table
153 1A), which find significant differences between communities in both global ($P = 0.001$, $R =$
154 0.784) and pairwise habitat comparisons ($P = 0.001 - 0.019$, $R = 0.341 - 0.983$). The only
155 exception are the coastal and urban forests ($P = 0.079$, $R = 0.172$) which may be due to the
156 close proximity of Pulau Ubin coastal forest sites to urban settlements (Fig. S1). Note that a
157 SIMPER analysis (Table 1B) finds a substantial number of shared species between the
158 rainforest and swamp forest sites (13.88%). Both sites are in close geographic proximity
159 (<5km; Fig. S1) and the within-habitat values for both sites are fairly high (rainforest =
160 29.59%, swamp forest = 31.10%). ANOSIM and SIMPER results are again robust to the
161 removal of rare species (Tables S7 & S8) and the ANOSIM p-values for most comparisons
162 are significant even according to re-defined statistical criteria for unexpected or new results
163 ($p < 0.005$)[62]. The observed dissimilarity was largely due to species turnover with the

164 turnover component (0.898) greatly outweighing nestedness (0.048; Table 1C & S9). This
165 was similarly observed in most pairwise comparisons of habitats (turnover = 0.704 – 0.956,
166 nestedness = 0.001 – 0.102). The only exception was mangroves and coastal forests
167 (turnover = 0.658, nestedness = 0.254) which are in close geographic proximity on Pulau
168 Ubin (Fig. 1).
169



170
171 **Figure 2.** Insect communities across tropical forest habitats are distinct based on Bray-
172 Curtis distances illustrated on 3D NMDS plots, regardless of whether (a) rare species are
173 removed or (b) the data are split into different taxonomic groups.

174

175

176 **Table 1.** Species turnover across habitats. (A) Distinctness of communities in each habitat
 177 type as assessed with ANOSIM (pairwise p-value below and R-statistics above diagonal. (B)
 178 Distinctness of communities in each habitat type as assessed with SIMPER. (C) Species
 179 turnover and nestedness analysis (pairwise turnover values below and nestedness above
 180 diagonal).

181 **A)**

Overall P: 0.001

Overall R: 0.784

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.817	0.983	0.953	0.973	0.955
Urban forest	0.001		0.759	0.815	0.575	0.172
Swamp forest	0.001	0.001		0.934	0.769	0.893
Mangrove	0.001	0.001	0.001		0.856	0.546
Freshwater swamp	0.001	0.001	0.008	0.001		0.341
Coastal forest	0.001	0.079	0.005	0.001	0.017	

182

183 **B)**

	Within habitat (%)	Between habitats (%)					
		Rain-forest	Urban forest	Swamp forest	Mangrove	Fresh-water swamp	Coastal forest
Rainforest	29.59						
Urban forest	12.91	3.20					
Swamp forest	31.10	13.88	2.94				
Mangrove	12.25	1.62	3.09	1.98			
Freshwater swamp	17.29	2.13	4.69	4.10	2.74		
Coastal forest	12.09	3.82	9.41	4.00	6.08	9.05	

184

185 **C)**

Overall Dissimilarity: 0.946

Overall Turnover: 0.898

Overall Nestedness: 0.048

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.011	0.072	0.054	0.007	0.021
Urban forest	0.916		0.028	0.097	0.005	0.102
Swamp forest	0.710	0.922		0.092	0.027	0.001
Mangrove	0.914	0.819	0.878		0.062	0.254
Freshwater swamp	0.956	0.891	0.932	0.878		0.093
Coastal forest	0.908	0.704	0.940	0.658	0.756	

186

187 *Relationship between insect and plant richness*

188 Compared to mangroves (ca. 250 plant species), rainforest and swamp forest sites
189 have 4.6 or 7.6 times the number of recorded plant species based on checklists for the sites
190 (Table S4). This higher species richness is also confirmed by plot data for the rainforest[63]
191 (839 species in 52 plots of 100m²) and swamp forest[64] (671 species in 40 plots of 400m²).
192 However, the insect biodiversity of the rainforest and swamp forest sites is only 1.64 – 1.98
193 times higher than in the mangroves after rarefaction.

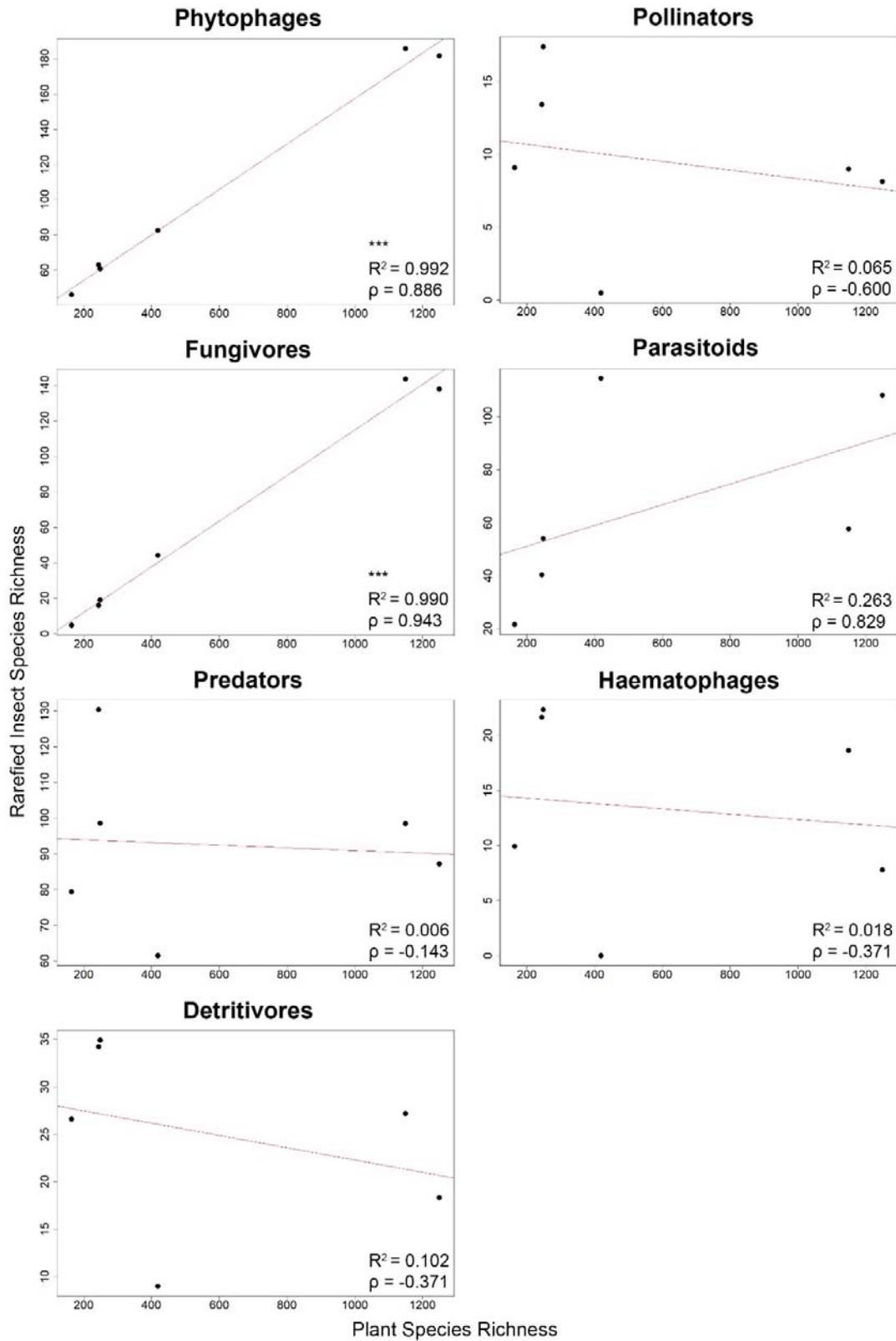
194 *Analysis of ecological guilds and correlation between insect and plant diversity*

195 For this analysis, we focused primarily on the Diptera and Hymenoptera which
196 occupy a broad range of ecological guilds and dominate Malaise trap samples (see Brown
197 2005[65] & Hebert et al. 2016[66]). We also excluded trapping sites that were sampled for
198 fewer than 6 months. We assigned insect species with known family/genus identities to
199 ecological guilds (42,092 specimens belonging to 2,230 putative species) in order to
200 understand how different habitats maintain insect diversity. After stepwise refinement of a
201 multivariate ANCOVA model, the final model was defined as: *insectdiv* ~ *habitat* + *guild* +
202 *plantdiv* + *guild:plantdiv* (*insectdiv*: rarefied insect species richness, *plantdiv*: plant species
203 richness). The type-II sum of squares test reveal that guild and the interaction term between
204 guild and plant diversity are highly significant factors ($p < 0.001$), while plant diversity ($p =$
205 0.063) and habitat ($p = 0.468$) are not. This suggests guild and plant diversity together have
206 an important role in determining insect diversity but the precise relationship warranted further
207 testing. Single variable linear regressions (*insectdiv* ~ *plantdiv*) were performed on each
208 guild separately (Fig. 3) and plant diversity was found to only be highly significantly and
209 positively correlated with the alpha-diversity of phytophagous and fungivorous insects ($p <$
210 0.001 , $R^2 = 0.992$ and 0.990 , $p = 0.886$ and 0.943 respectively).

211 After rarefaction, the different habitat types vary in composition (Fig. 4, see Table
212 S10). Rainforest and freshwater swamp forest sites have higher numbers and proportions of

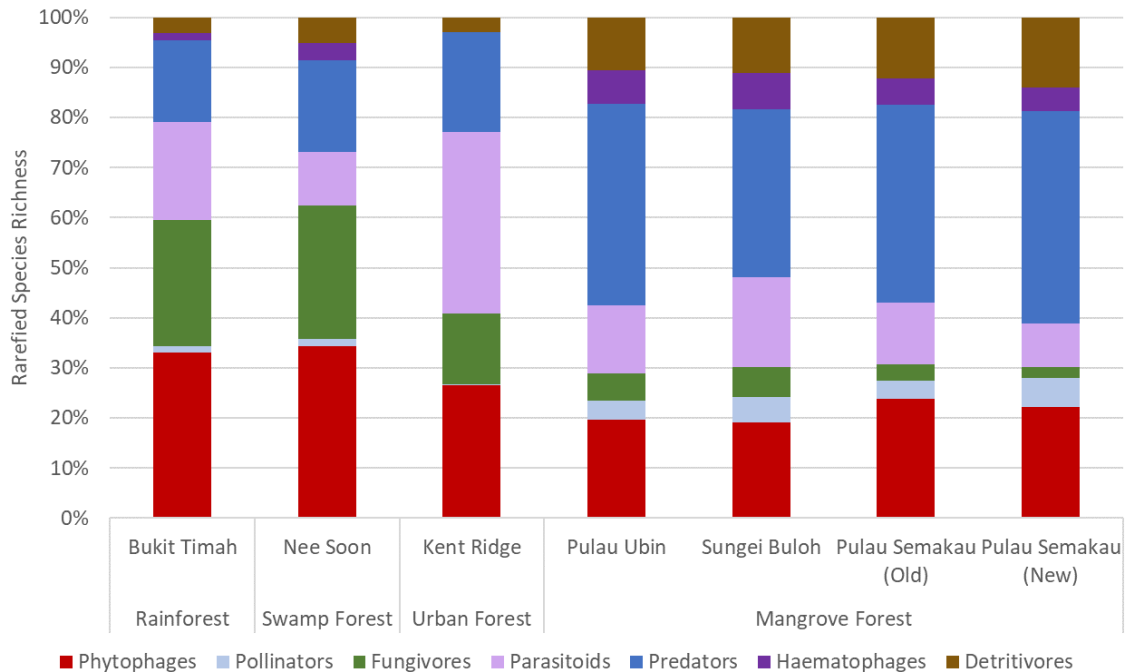
213 phytophagous and fungivorous insect species (see also Figs S4 & S5). The insect
214 communities of mangroves, however, are characterized by an unusually high proportion of
215 predatory species while the urban forest sites are dominated by parasitoids. With regard to
216 species turnover, communities are separated by habitat for most guilds and pairwise
217 comparisons (Fig. 5, Tables S11 & S12).

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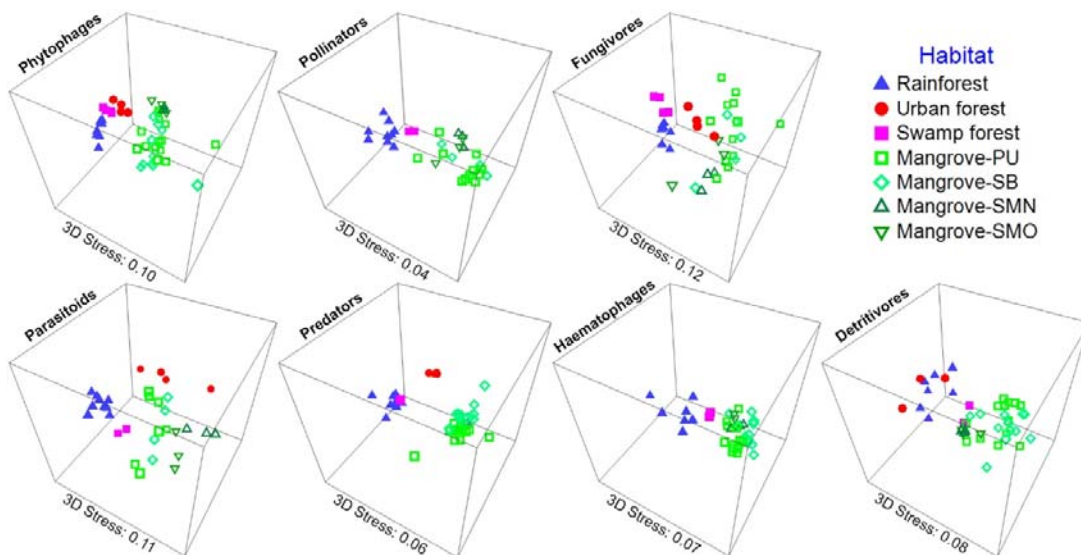
220 **Figure 3.** Only the diversity of phytophagous and fungivorous insects is correlated with plant
221 diversity based on a linear regression model using rarefied insect species richness (*: ≤ 0.05 ,
222 **: ≤ 0.01 , ***: ≤ 0.001).



223

224 **Figure 4.** Distribution of insect guilds across habitats. Phytophages and fungivores dominate
 225 in rain and swamp forest, predators in mangroves and parasitoids in the urban forest
 226 (rarefied samples).

227



228

229 **Figure 5.** Habitat differentiation by insect guilds (3D NMDS plot of Bray-Curtis distances for
 230 habitats with >2 sites).

231 *Species turnover across Asian mangroves*

232 The specimens from Hong Kong belonged to 109 dolichopodid, 129 phorid, and 25
233 mycetophilid species. The corresponding number for Brunei were 96 and 76 species for
234 dolichopodids and phorids, with too few mycetophilids being available for evaluation (Table
235 S3). The southern Thai dolichopodids belonged to 74 species. We find high species turnover
236 between Hong Kong, Brunei and Singapore, even after rarefying the specimen sample sizes
237 (Fig. S6). Approximately 90% of all dolichopodid and phorid species are unique to each
238 region and <1% are shared across all regions. Species turnover is even higher for the
239 mycetophilids of Hong Kong and Singapore (>95%). Species turnover for the dolichopodids
240 of Southern Thailand and Singapore is again high with only 11.5% of all species shared
241 between both countries.

242

243 **Discussion**

244 *Discovery of a largely overlooked, predator-enriched insect community in mangroves*

245 It is often assumed that the insect diversity in mangroves is low because high salinity
246 and low plant diversity are thought to interfere with insect diversification[23,67,68]. However,
247 we here show that mangroves are species-rich despite low plant diversity (<250 species:
248 [69–71]). In addition, the fauna of mangroves is very unique. More than half of its species
249 are not found in other habitats, even though coastal forests are adjacent to mangroves.
250 Indeed, after adjusting for sampling effort, the species diversity in Singapore's premier
251 rainforest reserve (Bukit Timah Nature Reserve: 1.64 km²) and largest swamp forest
252 remnant (Nee Soon: 5 km²) is only 50% higher than the diversity of major mangrove sites
253 (PU: 0.904 km², SB: 1.168 km², SM: 0.174 km²). The high diversity encountered in the
254 mangrove sites was particularly unexpected because the rainforests of Bukit Timah Nature
255 Reserve have been protected for more than 50 years[72,73] and have very high plant
256 diversity (e.g., 1,250 species of vascular plants[63] including 341 species of trees[74] in a 2
257 ha plot of the Centre for Tropical Forest Science). Moreover, we extensively sampled the

258 insect diversity in the reserve by placing multiple Malaise traps in primary, maturing
259 secondary, and old secondary forests. Similarly, we expected the insect diversity of
260 Singapore's largest swamp forest (Nee Soon) to greatly exceed the number of species found
261 in the mangrove sites because the swamp forest is also known for its high species richness
262 (e.g., 1,150 species of vascular plant species[75]).

263 A guild-level analysis reveals how mangroves maintain high species diversity. They
264 are impoverished with regard to phytophagous and fungivorous species, but are home to a
265 disproportionately large number of predatory species (Fig. 4) whose larvae develop in
266 sediments (Empidoidea and Tabanidae). This suggests that the high insect diversity in
267 different tropical habitats may be achieved by having larger proportions of species
268 developing in the biologically most productive microhabitats – plants and fungi for many
269 forest habitats and the rich and productive mud flats for mangroves.

270 In addition to finding high alpha-diversity in mangroves, we also document that the
271 mangrove insect communities are very distinct. This conclusion is supported by a multitude
272 of analyses (NMDS, ANOSIM & SIMPER). It is furthermore insensitive to the removal of rare
273 species (Fig. 2) and driven by high species turnover rather than nestedness (see Table 1C).
274 This stratification by habitat is still evident even when the two dominant insect orders in
275 Malaise trap samples (Diptera and Hymenoptera) are removed (Fig. 2). Comparatively high
276 overlap is only observed between mangroves and coastal forests (860 shared species)
277 which is likely due to close proximity of the habitats on Pulau Ubin (Fig. S1) where back
278 mangroves and coastal forests are contiguous. The uniqueness of the mangrove insect
279 community is likely due to the unusual environmental conditions characterized by extreme
280 daily fluctuations in salinity, temperature, and inundation. These extreme conditions likely
281 require physiological and behavioural adaptations that encourage the emergence of an
282 evolutionarily distinct fauna. What is surprising, however, is that we find no evidence for an
283 adaptive radiation of particular clades. Instead, a large number of independent colonization
284 events seems more likely given that the mangrove species usually belong to genera that are

285 also known from other habitats (e.g., Dolichopodidae). This challenges the view that high
286 salinity is a potent colonization barrier for invertebrates[67,68].

287 Mangrove regeneration is pursued in many countries, with mixed success in restoring
288 the original plant diversity[76,77], but it remains poorly understood whether the regenerated
289 mangroves harbour the original arthropod biodiversity. Our preliminary data based on 311
290 Malaise trap samples from one regenerated site suggests that this may not be the case. The
291 regenerated mangrove (SMN) was replanted with a monoculture of *Rhizophora stylosa*[71]
292 which replaced old-growth mangroves that had been cleared during reclamation work
293 (1994–1999[51]). The restored site (SMN) has markedly lower insect species richness than
294 all other mangrove sites, including a neighbouring old-growth mangrove (SMO; Fig. 1). This
295 highlights once more the need for holistic habitat assessments that goes beyond plants and
296 vertebrates[78].

297 Mangrove insect communities are not only rich and distinct in Singapore. Within Asia,
298 we reveal a 92% species turnover between Singapore and Hong Kong (2,500 km north; Fig.
299 S1) for taxa representing different guilds (Dolichopodidae–predators: 483 species,
300 Mycetophilidae–fungivores: 67 species, Phoridae–mostly saprophagous: 591 species).
301 While climatic differences could be advanced as a potential explanation, comparisons with
302 the mangroves in the geographically close and tropical Borneo (Brunei) confirm a high
303 species turnover of 85% (see also Grootaert 2019[79]). Further evidence for high regional
304 species turnover in mangroves emerges when the dolichopodid fauna of Singapore’s and
305 Brunei’s mangroves are compared with the fauna of Southern Thailand (coasts of South
306 China and Andaman seas). Only 34 and 10 of the 74 known Thai species are shared with
307 Singapore and Brunei respectively; These data suggest that a significant proportion of the
308 global insect diversity may reside in mangroves. Based on the data from Singapore, it
309 appears that much of the diversity may still be intact, given that we find no evidence that the
310 insect diversity in Singapore’s mangroves is depressed relative to what is found in the more
311 pristine sites in Brunei or Hong Kong. This suggests that the loss of species diversity for

312 small, flying insects in Singapore may not have been as dramatic as what has been
313 documented for vertebrates and larger invertebrates[48,80,81].

314 *Discovering a new insect hotspot with NGS barcoding*

315 Global insect declines have recently received much attention by the scientific
316 community[2] and public[82]. Obtaining relevant data is very difficult since quantifying insect
317 diversity using conventional techniques is slow and expensive. This is because too many
318 specimens have to be sorted into too many species before a holistic habitat assessment can
319 be carried out[83]. In our study, this problem is overcome via sorting based on NGS
320 barcodes which differ from traditional barcodes by costing only a fraction of barcodes
321 obtained with Sanger sequencing. Based on previous tests, we find that species delimited
322 with NGS barcodes have >90% congruence with species-level units delimited with
323 morphological data[56,57,84,85]. This suggests that large-scale species discovery with NGS
324 barcoding yields sufficiently accurate information on species abundance and distribution for
325 habitat assessments[55,56]; i.e., NGS barcodes can be used for quickly revealing hidden
326 hotspots of insect diversity in countries with high diversity and limited funding. We estimate
327 that the ~140,000 specimens in our study could today be sequenced for <USD25,000 using
328 350 manpower days whereas a similar study based on morphology would require >150
329 manpower years[86]; i.e. some of the traditional obstacles to understanding insect
330 biodiversity caused by the taxonomic impediment are finally disappearing.

331 *Concluding remarks*

332 We here document that the insect fauna inhabiting mangroves is not only rich, but
333 also distinct when compared to many other tropical forest habitats. The discovery of such an
334 unexpectedly rich and distinct insect community highlights how little we know about insect
335 diversity. We predict that advances in sequencing technology will facilitate the discovery of
336 numerous additional insect diversity hotspots in tropical and temperate habitats. Mangroves
337 will likely be only one of many future additions to the growing list of habitats that have only

338 recently been recognized as containing a large proportion of the global biodiversity (e.g., dry
339 forests[87,88], forest savannahs[89,90]). Our study highlights that accelerating species
340 discovery is a pressing task given that many of these habitats are disappearing at a much
341 faster rate than tropical rainforests.

342

343 **Methods**

344 *Sampling site, sample collection, and processing*

345 Singapore has a large number of tropical habitat types that are all within 40 km of
346 each other without being separated by major physical barriers. This allowed us to sample
347 rainforests (from early secondary to mature secondary forest), urban-edge forests,
348 mangroves, swamp forests, freshwater swamps and dry coastal forests. The freshwater
349 swamp habitat differs from swamp forests by largely lacking tree-cover, while the dry coastal
350 forests are distinct from the mangroves by lacking typical mangrove tree species. Note that
351 the habitats had experienced similar levels of habitat degradation or loss due to urbanization
352 (>95% loss of original vegetation cover[48]; ca. 90% loss for rainforests[49]; ca. 93% loss of
353 swamp forest[50]; 91% loss for mangroves[51]). We sampled these habitat types using 107
354 trapping sites (Fig. S1). The mangrove sites were located primarily along the North-western
355 and Southern coasts of the mainland, as well as on offshore islands in the south and
356 northeast. The major mangrove sites were on Pulau Ubin (PU), Sungei Buloh (SB) and
357 Pulau Semakau (SM), the last of which is represented by an old-growth (SMO) and a newly
358 regenerated mangrove fragment (SMN). The swamp forest site (Nee Soon) was Singapore's
359 largest remaining freshwater swamp remnant which is known for a rich insect fauna[91],
360 overall high species richness, and level of endemism[92,93]. Bukit Timah Nature Reserve
361 was selected as the tropical rainforest site given its high species diversity and protected
362 status[72]. This reserve consists of forests in various stages of succession and hence we
363 sampled different forest types with three sites each being in primary forest, old secondary

364 forest, and maturing secondary forest. The “urban secondary forest” sites were located along
365 a disturbance gradient ranging from the campus of the National University of Singapore
366 (NUS) through several urban parks and forest edges in Central and South Singapore. The
367 freshwater swamp site is located primarily in Kranji, a freshwater marsh at the flooded edge
368 of a reservoir. The “coastal forest” sites were dry secondary forests adjacent to the coast at
369 Labrador Park and Pulau Ubin, which are also close to urban settlements.

370 All specimens were collected between 2012–2019 (Table S1) using Malaise traps.
371 These traps are widely used for insect surveys because they are effective sampling tools for
372 flying insects and allow for standardized, long-term sampling. Note that the use of Malaise
373 traps in our study was appropriate because the canopy height was comparable for most
374 habitats given that we compared mature mangroves (PU, SB and SMO) with a wet swamp
375 forest site, and different kinds of secondary forests (pers. obs.). Only the canopy height of
376 some sites in Bukit Timah Nature Reserve (BTNR) was higher, but for BTNR we also
377 included secondary forests and several traps were placed on steep slopes that would be
378 able to sample canopy-active fauna from a lower elevation. With regard to the habitat
379 patches, the fragments were larger for the rainforest and swamp forest than for any of the
380 mangrove sites (tropical rainforest: 1.64 km²; swamp forest: 5 km², mangrove forest
381 fragments: 0.904 km² [PU], 1.168 km² [SB], 0.174 km² [SM][51]). Malaise traps in the
382 mangroves were set up in the intertidal zone. Each Malaise trap sample consisted of one-
383 week’s worth of insects preserved in molecular grade ethanol. After an ethanol change, the
384 specimens were sorted to order/family level by para-taxonomists, and specimens from 16
385 arthropod orders were extracted for barcoding (Fig. S2): Araneae, Blattodea, Coleoptera,
386 Diptera, Hemiptera, Hymenoptera, Lepidoptera, Mantodea, Megaloptera, Neuroptera,
387 Orthoptera, Phasmida, Plecoptera, Psocodea, Strepsiptera and Trichoptera. Diptera and
388 Hymenoptera were the dominant orders in the Malaise traps (Fig. S2: >75% of specimens)
389 and sorted further to family and genus-level where possible (Table S2), either based on
390 morphology or based on DNA barcodes identified using the Global Biodiversity Information

391 Facility (GBIF: www.gbif.org) or the Barcode of Life Data (BOLD: www.boldsystems.org)
392 databases. Only matches above 95% and 97% similarity were considered sufficiently precise
393 for family- and genus-level matches respectively. The mangrove specimens from Hong Kong
394 were collected by 24 Malaise traps installed between October 2017 to October 2018, while
395 those from Brunei were collected by six Malaise traps from July to November 2014. Note
396 that the mangrove forests in Brunei were less affected by urbanization than those in
397 Singapore. The dolichopodid specimens from Thailand were obtained by different
398 techniques including sweep-netting from 42 mangrove sites over a period of 15 months from
399 Mar 2014 – Dec 2015.

400 *Putative species sorting with NGS barcoding*

401 NGS barcoding combines the advantages of cost-effective sequencing with Illumina
402 with the approximate species-level resolution provided by DNA barcodes. The molecular
403 procedures can be learned in hours and several hundred specimens can be processed per
404 person and day. The overall barcode costs are now <10 cents per specimen if Illumina
405 Novaseq is used for sequencing (2 cents/barcode based on USD 6,900 per 250-bp PE flow
406 cell yielding 800 million reads: <https://research.ncsu.edu/gsl/pricing>). We used NGS
407 barcoding to amplify and sequence a 313-bp fragment of the cytochrome oxidase I gene
408 (*cox1*) using a protocol described in Meier et al.[55]. Direct-PCR[94] was conducted for
409 specimens collected early in the study; during this phase, we used 1-2 legs of the specimen
410 as template for obtaining the amplicon with the primer pair mICO1intF: 5'-
411 GGWACWGGWTGAACWGTWTAYCCYCC-3'[95] and jgHCO2198: 5'-
412 TANACYTCNGGRTGNCCRAARAAYCA-3'[96]. For samples processed later, the whole
413 specimen was immersed in Lucigen QuickExtract solution or HotSHOT buffer[97] and gDNA
414 extraction was conducted non-destructively. The gDNA extract was then used as a PCR
415 template with the afore-mentioned reagents and protocol. The primers used were labelled
416 with 9-bp long barcodes that differed by at least three base pairs. Every specimen in each
417 sequencing library was assigned a unique combination of labelled forward and reverse

418 primers, which allowed the Illumina reads to be binned according to specimen. A negative
419 control was prepared and sequenced for each 96-well PCR plate. Amplification success
420 rates for each plate were assessed via gel electrophoresis for eight random wells per plate.

421 The amplicons were pooled at equal volumes within each plate and later pooled
422 across plates. Equimolarity was estimated by the presence and intensity of bands on gels.
423 The pooled samples were cleaned with Bionline SureClean Plus and/or via gel cuts before
424 outsourcing library preparation to AITbiotech using TruSeq Nano DNA Library Preparation
425 Kits (Illumina) or the Genome Institute of Singapore (GIS) using NEBNext DNA Library
426 Preparation Kits (NEB). Paired-end sequencing was performed on Illumina Miseq (2x300-bp
427 or 2x250-bp) or HiSeq 2500 platforms (2x250-bp) over multiple runs, thereby allowing
428 troubleshooting and re-sequencing for specimens which initially failed to yield a sufficiently
429 large number of reads. Some of the specimens were also sequenced on the MinION (Oxford
430 Nanopore) platform using primers with a slightly longer tags (13-bp) and following the
431 protocol described in Srivathsan et al.[98,57]. Raw Illumina reads were processed with the
432 bioinformatics pipeline and quality-control filters described in Meier et al.[55]. A BLAST
433 search to GenBank's nucleotide (nt) database was also conducted to identify and discard
434 contaminants by parsing the BLAST output through *readsidentifier*[99] and removing
435 barcodes with incorrect matches at >97% identity.

436 To obtain putative species units, the *cox1* barcodes were clustered over a range of
437 uncorrected p-distance thresholds (2–4%) typically used for species delimitation in the
438 literature[100]. The clustering was performed with a python script that implements the
439 objective clustering algorithm of Meier et al. 2006[59] and allows for large scale processing.
440 USEARCH[60] (*cluster_fast*) was used to confirm the results by setting *-id* at 0.96, 0.97 and
441 0.98. To gauge how many of our species/specimens matched barcodes in public databases,
442 we used the "Sequence ID" search of the Global Biodiversity Information Facility (GBIF). We
443 then determined the number of matches with identity scores <97. We then counted the
444 number of matches to barcodes with species-level identifications.

445 *Diversity analyses*

446 For analysis of species richness and turnover, we excluded 11 trapping sites which
447 had <100 specimens per site in order to prevent poor sampling from inflating site
448 distinctness. To assess the species richness of the six major habitat types, samples were
449 rarefied with the *iNEXT*[101] R package (R Development Core Team) using 1,000 bootstrap
450 replicates in order to account for unequal sampling completeness. The rarefaction was
451 performed by coverage[61] in the main analysis (Fig. 1) and by specimen count in the
452 supplementary (Fig. S3). Site comparisons were carried out by comparing species diversity
453 post-rarefaction to the lowest coverage/smallest number of specimens. The habitat type
454 “mangrove” was treated both as a single habitat as well as separate sites (PU, SB, SMN,
455 SMO, others) in separate analyses.

456 In order to study species turnover, we determined the distinctness of the
457 communities across habitats using non-metric multidimensional scaling (NMDS) plots that
458 were prepared with PRIMER v7[102] using Bray-Curtis dissimilarity. Plots were generated
459 for each habitat type and the different mangrove sites; Bray-Curtis was chosen because it is
460 a preferred choice for datasets that include abundance information. The dataset was split
461 into three groups: the dominant orders (Diptera and Hymenoptera) and all others combined,
462 in order to test if the results were driven by the dominant orders. Analysis of similarities
463 (ANOSIM) and similarity percentages (SIMPER) were performed in PRIMER under default
464 parameters in order to obtain ANOSIM p-values and R-statistics for both the entire dataset
465 and the pairwise comparisons between habitat types. The SIMPER values were calculated
466 for within and between-habitat types. The ANOSIM p-values can be used to assess
467 significant differences while the R-statistic allows for determining the degree of similarity,
468 with values closer to 1 indicating greater distinctness. We also used the *betapart*[103] R
469 package to examine if the observed dissimilarity (Bray-Curtis) was due to species turnover
470 or nestedness. The *beta.multi.abund* and *beta.pair.abund* functions were used to split the
471 global and pairwise dissimilarity scores into turnover and nestedness components. Lastly,

472 the robustness of the results was tested by removing singleton, doubleton and rare species
473 (<5 and <10 individuals) from the datasets. The pruned datasets were subjected to the same
474 analyses as the full dataset. For the guild-specific datasets, traps with fewer than three
475 species were excluded in the species turnover analyses because large distances driven by
476 undersampling can obscure signal.

477 To examine species turnover across larger geographic scales, dolichopodid, phorid,
478 and mycetophilid specimens from Singapore were compared with those from Hong Kong
479 (Dolichopodidae: 2,601; Phoridae: 562, Mycetophilidae: 186), and Brunei (Dolichopodidae:
480 2,800; Phoridae: 272), and data for the dolichopodids of Southern Thai mangroves (942
481 specimens). Since Singapore was more extensively sampled, the Singaporean dataset was
482 randomly subsampled (10 iterations in Microsoft Excel with the RAND() function) to the
483 number of specimens available for the other two countries (Table S3). The species diversity
484 after rarefaction was then compared (with 95% confidence intervals for the rarefied data).

485 *Ecological guild and plant diversity analyses*

486 For the guild-level analysis, we focused primarily on the two dominant orders Diptera
487 and Hymenoptera, which comprised of species from a large variety of ecological guilds. As
488 splitting the dataset into smaller guild-level partitions would create low-abundance subsets,
489 we excluded trapping sites that were sampled for <6 months, resulting in a dataset
490 consisting of 62,066 specimens from 9 rainforest, 4 swamp forest, 4 urban forest, and 32
491 mangrove sites (Fig. S1). In order to test for an overall correlation between plant and insect
492 diversity, we obtained data for the plant diversity in the respective sites from checklists and
493 survey plots (Table S4). In order to further examine the correlation between plant and insect
494 diversity across multiple ecological guilds, we assigned the identified Diptera and
495 Hymenoptera families and genera non-exclusively to ecological guilds (phytophages,
496 pollinators, fungivores, parasitoids, predators, haematophages and detritivores) based on
497 known adult and larval natural history traits for the group (Table S2). Taxa with different adult
498 and larval natural histories are placed in both guilds. Taxa lacking sufficient information or

499 with highly variable life-history strategies were assigned to the “Others/Unknown” category
500 and excluded from analysis.

501 Barcodes from each guild were separately aligned and clustered at 3% p-distance.
502 These subsets were used for further analysis by randomly subsampling (10 iterations in
503 Microsoft Excel with the RAND() function) the same number of specimens at the site with the
504 smallest number of specimens (urban forest site, 2,543 specimens). For taxa that have
505 adults and immatures with different natural histories (i.e., belong to two distinct ecological
506 guilds), the species counts were halved and placed into both guilds when calculating rarefied
507 species abundance and richness. Species turnover for the guild-specific subsets were
508 analysed with PRIMER to generate NMDS plots, as well as ANOSIM and SIMPER values.
509 The rarefied species richness values were also used for a multivariate model analysis. An
510 ANCOVA model was constructed in R[104] with the *lm* function: *insectdiv ~ site * habitat *
511 guild * plantdiv*, with *insectdiv* representing rarefied insect alpha-diversity and *plantdiv*
512 representing plant species counts. The “site” factor was excluded due to collinearity and the
513 model was refined via stepwise removal of factors starting with the most complex (interaction
514 terms) and least significant ones. At each stage, the *anova* function was used to assess loss
515 of informational content and the final model was derived when the reported p-value was
516 significant ($p < 0.05$). The model’s residuals were examined to ensure the data were normal.
517 Subsequently, the *Anova* function from the *car* package[105] was used to obtain type-II test
518 statistics. Finally, single-variable linear regression was performed in R with the *lm* function:
519 *insectdiv ~ plantdiv* for each guild separately to obtain significance, multiple R-squared and
520 Spearman’s rho values.

521

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Competing interests

None

Table S1. Collection periods and trap localities; M = mangroves, SF = swamp forest, UF = urban forest, TF = tropical rainforest, CF = coastal forest, FS = freshwater swamp.

Sampling period	Location	Habitat type	GPS coordinates	No. of traps	Total no. of weekly samples	Used for guild-level analyses
Singapore						
Apr 2012 – Mar 2014	Pulau Ubin	M	1°24'36.3"N 103°59'25.5"E	3	72	Y
	Pulau Semakau original	M	1°12'17.6"N 103°45'37.7"E	3	72	Y
	Pulau Semakau replanted	M	1°12'03.1"N 103°45'45.4"E	3	72	Y
	Sungei Buloh Wetland Reserve	M	1°26'46.3"N 103°43'49.9"E	2	48	Y
	Nee Soon freshwater swamp	SF	1°23'00.3"N 103°48'46.5"E	2	48	Y
May 2014 – Jun 2014	Mandai Nature Park	M	1°26'18.3"N 103°45'49.7"E	3	6	N
	Pulau Tekong	M	1°25'47.3"N 104°03'46.3"E	3	6	N
	Sarimbun	M	1°25'59.1"N 103°41'21.8"E	3	6	N
Nov 2014 – May 2015	Nee Soon freshwater swamp	SF	1°23'00.3"N 103°48'46.5"E	2	14	Y
Apr 2015 – Sep 2015	NUS	UF	1°17'49.6"N 103°46'35.7"E	4	24	Y
Mar 2016 – Aug 2016	Pulau Ubin	M	1°25'11.64"N 103°56'6.25"E	10	60	Y
	Sungei Buloh Wetland Reserve	M	1°26'43.20"N 103°43'5.10"E	10	60	Y
	Labrador Park	M	1°16'13.3"N 103°48'10.1"E	3	18	N
	Labrador Park	CF	1°16'05.4"N 103°48'16.2"E	2	18	N
Aug 2016 – Oct 2017	Bukit Timah Nature Reserve primary forest	TF	1°21'13.90"N 103°46'47.57"E	3	45	Y
	Bukit Timah Nature Reserve old secondary forest	TF	1°21'17.96"N 103°46'54.01"E	3	45	Y
	Bukit Timah Nature	TF	1°21'4.57"N 103°46'53.80"E	3	45	Y

	Reserve maturing secondary forest						
Apr 2017 – 20 Sep 2017	NUS	UF	1°17'49.6"N 103°46'35.7"E	4	18	N	
Sep 2017 – Dec 2017	NUS	UF	1°17'45.3"N 103°46'13.8"E	3	16	N	
May 2018	Bishan-Ang Moh Kio Park	UF	1°21'35.7"N 103°50'49.9"E	2	2	N	
	Enabling Village	UF	1°17'13.6"N 103°48'53.3"E	1	1	N	
	Esplanade Theatre	UF	1°17'26.4"N 103°51'17.9"E	1	1	N	
Mar 2018 – Jun 2018	Sungei Buloh Wetland Reserve	M	1°26'52.45"N 103°43'24.16"E	4	16	N	
	Kranji Marshes	FS	1°25'0.56"N 103°43'43.50"E	3	12	N	
	Lim Chu Kang	M	1°26'48.80"N 103°42'35.71"E	2	8	N	
	Mandai Nature Park	M	1°26'37.96"N 103°45'59.70"E	4	16	N	
	Pulau Ubin	CF	1°24'26.3"N 103°57'16.3"E	3	12	N	
	Pulau Ubin	M	1°24'32.2"N 103°57'12.1"E	8	32	N	
	Labrador Park	M	1°16'13.3"N 103°48'10.1"E	5	20	N	
	Labrador Park	CF	1°16'05.4"N 103°48'16.2"E	4	20	N	
Mar 2019 – Jun 2019	Coney Island	M	1°24'37.3"N 103°55'23.1"E	5	15	N	
	Kranji Marshes	FS	1°25'11.0"N 103°43'54.3"E	4	15	N	
	Pulau Ubin	CF	1°25'34.7"N 103°56'29.2"E	1	15	N	
	Pulau Ubin	M	1°25'05.3"N 103°56'06.5"E	7	15	N	
Hong Kong							
Nov 17 – Dec 17, May 18 – Jul 18	Ha Pak Nai	M	22°25'31.48"N 113°56'20.11"E	6	30	Y	
	Hang Mei	M	22°15'9.83"N 113°52'5.84"E	5	25	Y	
	Ho Chung	M	22°21'13.18"N 114°15'7.45"E	6	30	Y	
	Lai Chi Wo	M	22°31'37.63"N 114°15'43.63"E	5	25	Y	
	Nam Chung	M	22°31'31.62"N 114°12'28.94"E	5	25	Y	
	Sai Keng	M	22°25'13.48"N 114°16'4.66"E	5	25	Y	
	Sam A Chung	M	22°30'29.84"N 114°16'20.93"E	5	25	Y	

	Sam A Tsuen	M	22°30'55.22"N 114°16'16.36"E	5	25	Y
	Sha Tau Kok	M	22°32'4.34"N 114°12'39.78"E	10	50	Y
	Sheung Pak Nai	M	22°27'7.09"N 113°57'45.11"E	5	25	Y
	Shui Hau	M	22°13'9.70"N 113°55'8.33"E	5	25	Y
	So Lo Pun	M	22°32'17.20"N 114°15'21.49"E	5	25	Y
	Tai O	M	22°15'28.44"N 113°51'48.96"E	6	30	Y
	Tai Tam	M	22°14'46.10"N 114°13'24.02"E	3	15	Y
	Tai Tan	M	22°26'18.85"N 114°19'59.77"E	1	5	Y
	To Kwa Peng	M	22°25'43.07"N 114°19'59.30"E	5	25	Y
	Tsim Bei Tsui	M	22°29'20.47"N 113°59'53.95"E	5	25	Y
	Tung Chung	M	22°16'52.50"N 113°55'44.04"E	6	30	Y
	Wong Chuk Wan	M	22°23'44.27"N 114°17'10.21"E	5	25	Y
	Yim Tin Tsai	M	22°22'32.74"N 114°18'5.76"E	5	25	Y
Brunei						
Jul 14 – Nov 14	Pulau Berembang	M	4°54'7.44"N 115°1'17.94"E	2	10	Y
	Labu Forest Reserve	M	4°51'41.75"N 115°6'59.69"E	2	10	Y
	Tutong Forest	M	4°46'9.54"N 114°36'20.64"E	2	10	Y

Chloropidae	<i>Tricimba</i>				✓	
Clusiidae		✓				✓
Coelopidae		✓				
Cryptochetidae			✓			
Culicidae					✓	✓
Diastatidae						✓
Diopsidae		✓				✓
Dolichopodidae				✓		
Drosophilidae	<i>Apenthecia</i>	✓				
Drosophilidae	<i>Chymomyza</i>	✓				
Drosophilidae	<i>Colocasiomyia</i>	✓				
Drosophilidae	<i>Dichaetophora</i>		✓			
Drosophilidae	<i>Drosophila</i>					✓
Drosophilidae	<i>Gitona</i>	✓				
Drosophilidae	<i>Hirtodrosophila</i>		✓			
Drosophilidae	<i>Hypselothyrea</i>	✓				
Drosophilidae	<i>Leucophenga</i>	✓				
Drosophilidae	<i>Liodrosophila</i>	✓				
Drosophilidae	<i>Luzonimyia</i>					✓
Drosophilidae	<i>Microdrosophila</i>	✓				
Drosophilidae	<i>Mycodrosophila</i>		✓			
Drosophilidae	<i>Paramycodrosophila</i>		✓			
Drosophilidae	<i>Scaptodrosophila</i>	✓				
Drosophilidae	<i>Scaptomyza</i>	✓				
Drosophilidae	<i>Stegana</i>	✓				
Drosophilidae	<i>Zaprionus</i>	✓				
Empididae				✓		
Ephydriidae	<i>Allotrichoma</i>					✓
Ephydriidae	<i>Atissa</i>					✓
Ephydriidae	<i>Brachydeutera</i>	✓				
Ephydriidae	<i>Cerobothrium</i>					✓
Ephydriidae	<i>Ceropsilopa</i>	✓				
Ephydriidae	<i>Discocerina</i>	✓				
Ephydriidae	<i>Donaceus</i>					✓

Ephydridae	<i>Glenanthe</i>				✓
Ephydridae	<i>Hecamedoides</i>				✓
Ephydridae	<i>Hydrellia</i>	✓			
Ephydridae	<i>Limnellia</i>				✓
Ephydridae	<i>Nostima</i>	✓			
Ephydridae	<i>Notiphila</i>				✓
Ephydridae	<i>Ochthera</i>			✓	
Ephydridae	<i>Orasiopa</i>				✓
Ephydridae	<i>Paralimna</i>	✓			
Ephydridae	<i>Placopsidella</i>			✓	
Ephydridae	<i>Polytrichophora</i>				✓
Ephydridae	<i>Ptilomyia</i>				✓
Ephydridae	<i>Rhynchopsilopa</i>				✓
Ephydridae	<i>Trimerogastra</i>				✓
Ephydridae	<i>Trypetomima</i>				✓
Ephydridae	<i>Zeros</i>				✓
Hybotidae				✓	
Keroplastidae		✓	✓		
Lauxaniidae		✓			
Lonchaeidae		✓			
Lygistorrhinidae		✓	✓		
Megamerinidae				✓	
Micropezidae					✓
Milichiidae	<i>Aldrichiomyza</i>				✓
Milichiidae	<i>Leptometopa</i>			✓	
Milichiidae	<i>Milichia</i>				✓
Milichiidae	<i>Milichiella</i>			✓	
Milichiidae	<i>Neophyllomyza</i>	✓			
Milichiidae	<i>Paramyia</i>				✓
Milichiidae	<i>Phyllomyza</i>			✓	
Muscidae					✓
Mycetophilidae		✓	✓		
Neriidae		✓			✓
Odiniidae					✓

Periscolidae		✓					
Phoridae							✓
Pipunculidae					✓		
Platypozidae			✓			✓	
Platystomatidae							✓
Psilidae		✓					
Pyrgotidae					✓		
Rhagionidae					✓	✓	
Rhiniidae					✓		
Sarcophagidae							✓
Sciaridae		✓	✓				
Sphaeroceridae							✓
Stratiomyiidae		✓				✓	
Syrphidae	<i>Allobaccha</i>		✓		✓		
Syrphidae	<i>Allograpta</i>		✓		✓		
Syrphidae	<i>Asarkina</i>		✓		✓		
Syrphidae	<i>Ceriana</i>		✓			✓	
Syrphidae	<i>Eosmallota</i>		✓			✓	
Syrphidae	<i>Eristalinus</i>		✓			✓	
Syrphidae	<i>Eristalis</i>		✓			✓	
Syrphidae	<i>Eumerus</i>		✓			✓	
Syrphidae	<i>Graptomyza</i>		✓			✓	
Syrphidae	<i>Ischiodon</i>		✓		✓		
Syrphidae	<i>Microdon</i>		✓		✓		
Syrphidae	<i>Paragus</i>		✓		✓		
Syrphidae	<i>Psilota</i>		✓			✓	
Syrphidae	<i>Spheginobaccha</i>		✓				✓
Syrphidae	<i>Syrpita</i>		✓			✓	
Syrphidae	<i>Volucella</i>		✓			✓	
Tabanidae					✓	✓	
Tachinidae					✓		✓
Tephritidae		✓					
Ulidiidae							✓
Xenasteiidae							✓

Xylomyidae				✓
Hymenoptera				
Aphelinidae			✓	
Apidae	✓			
Bethylidae			✓	✓
Braconidae			✓	
Ceraphronidae			✓	
Chalcidae			✓	
Chrysididae			✓	
Colletidae	✓			
Crabronidae				✓
Diapriidae			✓	
Dryinidae			✓	
Eulophidae			✓	
Eupelmidae			✓	
Evaniidae			✓	
Figitidae			✓	
Formicidae	<i>Acropyga</i>			✓
Formicidae	<i>Anochetus</i>			✓
Formicidae	<i>Anoplolepis</i>			✓
Formicidae	<i>Aphaenogaster</i>			✓
Formicidae	<i>Brachyponera</i>			✓
Formicidae	<i>Camponotus</i>			✓
Formicidae	<i>Cardiocondyla</i>			✓
Formicidae	<i>Carebara</i>			✓
Formicidae	<i>Cataulacus</i>			✓
Formicidae	<i>Chronoxenus</i>			✓
Formicidae	<i>Colobopsis</i>			✓
Formicidae	<i>Crematogaster</i>			✓
Formicidae	<i>Cryptopone</i>			✓
Formicidae	<i>Diacamma</i>			✓
Formicidae	<i>Discothyrea</i>			✓
Formicidae	<i>Dolichoderus</i>			✓
Formicidae	<i>Echinopla</i>			✓

Formicidae	<i>Ectomomyrmex</i>		✓	
Formicidae	<i>Euponera</i>		✓	
Formicidae	<i>Euprenolepis</i>			✓
Formicidae	<i>Gauromyrmex</i>			✓
Formicidae	<i>Hypoponera</i>		✓	
Formicidae	<i>Iridomyrmex</i>			✓
Formicidae	<i>Leptogenys</i>			✓
Formicidae	<i>Lioponera</i>		✓	
Formicidae	<i>Mayriella</i>			✓
Formicidae	<i>Meranoplus</i>			✓
Formicidae	<i>Mesoponera</i>			✓
Formicidae	<i>Monomorium</i>			✓
Formicidae	<i>Myrmecina</i>		✓	
Formicidae	<i>Nylanderia</i>			✓
Formicidae	<i>Odontomachus</i>			✓
Formicidae	<i>Odontoponera</i>			✓
Formicidae	<i>Oecophylla</i>			✓
Formicidae	<i>Paraparatrechina</i>			✓
Formicidae	<i>Paratopula</i>			✓
Formicidae	<i>Paratrechina</i>			✓
Formicidae	<i>Pheidole</i>			✓
Formicidae	<i>Philidris</i>	✓		
Formicidae	<i>Platythyrea</i>		✓	
Formicidae	<i>Polyrhachis</i>	✓		
Formicidae	<i>Ponera</i>			✓
Formicidae	<i>Prenolepis</i>			✓
Formicidae	<i>Prionopelta</i>			✓
Formicidae	<i>Proatta</i>		✓	
Formicidae	<i>Probolomyrmex</i>			✓
Formicidae	<i>Pseudoneoponera</i>		✓	
Formicidae	<i>Strumigenys</i>		✓	
Formicidae	<i>Rhopalomastix</i>		✓	
Formicidae	<i>Solenopsis</i>			✓
Formicidae	<i>Stigmatomma</i>		✓	

Formicidae	<i>Strumigenys</i>		✓	
Formicidae	<i>Tapinoma</i>			✓
Formicidae	<i>Technomyrmex</i>			✓
Formicidae	<i>Tetramorium</i>		✓	
Formicidae	<i>Tetraoponera</i>	✓		
Formicidae	<i>Vollenhovia</i>			✓
Halictidae		✓		
Ichneumonidae			✓	
Megachilidae		✓		
Mymaridae			✓	
Platygastridae			✓	
Pompilidae			✓	
Pteromalidae			✓	
Scoliidae			✓	
Sphecidae			✓	
Sphecidae				✓
Tiphiidae			✓	
Trichogrammatidae			✓	
Vespidae		✓		✓

Table S3. Number of specimens from Singapore, Hong Kong and Brunei, as well as the size of the randomized subsample from Singapore.

Taxon	No. of Specimens				
	Singapore	Singapore (Rarefied)	Hong Kong	Brunei	Thailand
Dolichopodidae	17860	2800	2563	2798	924
Phoridae	2134	560	562	272	-
Mycetophilidae	223	180	186	-	-
Total	20217	3540	3311	3070	924

Table S4. Number of species of vascular plants for each sampling site in Singapore from checklist data.

Sampling Site	Habitat	No. of Plant Species	Reference
Nee Soon freshwater swamp	Freshwater swamp forest	1150	Wong et al., 2013 ¹
Bukit Timah Nature Reserve	Rainforest	1250	Ho et al., 2019 ²
Kent Ridge	Urban-edge/disturbed forest	420	Tan et al., 2019 ³
Pulau Ubin	Mangrove	245	Lee et al., 2003 ⁴
Sungei Buloh Wetland Reserve	Mangrove	249	Tan et al., 1997 ⁵
Pulau Semakau	Mangrove	165	Teo et al., 2011 ⁶

Table S5. Number and distribution of mOTUs delimited using different thresholds (144,865 barcoded specimens)

Habitat/Country	No. of Barcodes	No. of mOTUs from Objective Clustering			No. of mOTUs from USEARCH		
		2%	3%	4%	id=0.98	id=0.97	id=0.96
Singapore full dataset							
Mangroves	67239	3557	3437	3320	3710	3524	3436
Rainforest	15669	2625	2573	2539	2669	2603	2570
Swamp forest	9464	1843	1804	1753	1895	1828	1795
Urban forest	20323	1552	1515	1478	1616	1549	1510
Freshwater swamp	21994	1881	1812	1744	1988	1878	1805
Coastal forest	9118	1707	1667	1627	1755	1691	1664
Total	143807	8903	8572	8256	9315	8821	8520
Subset used for guild-level analysis							
Mangroves	37641	1778	1720	1673	1828	1744	1702
Rainforest	9212	1525	1490	1474	1545	1503	1483
Swamp forest	5893	1090	1052	1030	1105	1070	1048
Urban forest	9320	919	898	885	941	908	893
Total	62066	4169	4002	3917	4298	4098	3994
Southeast and East Asian datasets							
<i>Dolichopodidae</i>							
Singapore	17860	263	254	248	280	259	249
Hong Kong	2601	111	109	104	115	110	106
Brunei	2800	98	96	95	107	98	95
Thailand	924	80	74	72	93	80	73
Total	24185	480	453	426	543	482	447
<i>Phoridae</i>							
Singapore	2134	293	281	278	300	285	280
Hong Kong	562	137	129	125	138	130	129
Brunei	272	76	76	75	77	76	75
Total	2968	453	429	417	467	437	431
<i>Mycetophilidae</i>							
Singapore	223	45	44	43	45	44	44
Hong Kong	186	26	25	25	26	25	25
Total	409	69	67	67	70	67	67

Table S6. Common and rare species found in only 1, 2, 3, 4, 5 or all habitats.

	No. of species				
	Full dataset	No singletons	No doubletons	No species with <5 specimens	No species with <10 specimens
Species in mangroves only	1788	880	638	441	256
Species in rainforests only	1569	638	415	243	91
Species in swamp forests only	875	342	200	102	39
Species in urban forests only	509	166	101	58	25
Species in freshwater swamps only	794	360	237	127	56
Species in coastal forests only	454	153	71	33	14
Species in two habitats	1580	1580	1253	887	555
Species in three habitats	565	565	565	494	350
Species in four habitats	274	274	274	265	230
Species in five habitats	116	116	116	116	109
Species in all habitats	48	48	48	48	48
Total	8572	5122	3918	2814	1773

Table S7. Species turnover ANOSIM analysis results indicate distinct communities in each habitat type, whether with singletons and doubletons removed, or species with less than 5 and 10 specimens. Pairwise p-value outputs are displayed in the bottom-left of the pairwise matrix while the R-statistics are displayed at the top-right.

No Singletons

Overall P: 0.001

Overall R: 0.777

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.809	0.981	0.948	0.972	0.951
Urban forest	0.001		0.747	0.815	0.571	0.173
Swamp forest	0.001	0.001		0.927	0.756	0.893
Mangrove	0.001	0.001	0.001		0.852	0.541
Freshwater swamp	0.001	0.001	0.008	0.001		0.347
Coastal forest	0.001	0.083	0.005	0.001	0.017	

No Doubletons

Overall P: 0.001

Overall R: 0.774

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.803	0.980	0.946	0.972	0.954
Urban forest	0.001		0.735	0.816	0.563	0.179
Swamp forest	0.001	0.001		0.922	0.750	0.889
Mangrove	0.001	0.001	0.001		0.849	0.538
Freshwater swamp	0.001	0.001	0.008	0.001		0.331
Coastal forest	0.002	0.072	0.005	0.001	0.019	

No Species <5 Specimens

Overall P: 0.001

Overall R: 0.767

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.795	0.970	0.941	0.971	0.954
Urban forest	0.001		0.720	0.817	0.559	0.180
Swamp forest	0.001	0.001		0.913	0.750	0.885
Mangrove	0.001	0.001	0.001		0.843	0.533
Freshwater swamp	0.002	0.001	0.008	0.001		0.331
Coastal forest	0.002	0.061	0.005	0.001	0.017	

No Species <10 Specimens

Overall P: 0.001

Overall R: 0.759

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.779	0.959	0.934	0.967	0.952
Urban forest	0.001		0.701	0.819	0.548	0.178
Swamp forest	0.001	0.002		0.904	0.738	0.877
Mangrove	0.001	0.001	0.001		0.837	0.526
Freshwater swamp	0.002	0.001	0.008	0.001		0.331
Coastal forest	0.001	0.062	0.005	0.001	0.017	

Table S8. Species turnover SIMPER analysis results indicate distinct communities in each habitat type, whether with singletons and doubletons removed, or species with less than 5 and 10 specimens.

No Singletons

	Within habitat (%)	Between habitats (%)					
		Rain-forest	Urban forest	Swamp forest	Mangrove	Fresh-water swamp	Coastal forest
Rainforest	33.65						
Urban forest	13.70	3.57					
Swamp forest	35.74	15.86	3.31				
Mangrove	12.78	1.80	3.26	2.22			
Freshwater swamp	18.80	2.36	5.06	4.57	2.93		
Coastal forest	12.98	4.27	10.04	4.50	6.44	9.82	

No Doubletons

	Within habitat (%)	Between habitats (%)					
		Rain-forest	Urban forest	Swamp forest	Mangrove	Fresh-water swamp	Coastal forest
Rainforest	35.83						
Urban forest	14.15	3.79					
Swamp forest	38.05	17.12	3.55				
Mangrove	13.14	1.91	3.36	2.39			
Freshwater swamp	19.61	2.52	5.30	4.90	3.07		
Coastal forest	13.62	4.57	10.44	4.86	6.68	10.37	

No Species <5 Specimens

	Within habitat (%)	Between habitats (%)					
		Rain-forest	Urban forest	Swamp forest	Mangrove	Fresh-water swamp	Coastal forest
Rainforest	38.68						
Urban forest	14.86	4.13					
Swamp forest	40.06	18.89	3.93				
Mangrove	13.65	2.08	3.50	2.65			
Freshwater swamp	20.84	2.76	5.68	5.46	3.29		
Coastal forest	14.49	4.99	11.08	5.39	7.03	11.15	

No Species <10 Specimens

	Within habitat (%)	Between habitats (%)					
		Rain-forest	Urban forest	Swamp forest	Mangrove	Fresh-water swamp	Coastal forest
Rainforest	42.79						
Urban forest	15.91	4.79					
Swamp forest	42.93	21.56	4.49				
Mangrove	14.55	2.41	3.75	3.04			
Freshwater swamp	22.41	3.25	6.28	6.25	3.63		
Coastal forest	15.95	5.83	12.09	6.19	7.65	12.16	

Table S9. Species turnover and nestedness analysis reveal that the high dissimilarity is due more to turnover rather than nestedness, whether with singletons and doubletons removed, or species with less than 5 and 10 specimens. Pairwise turnover values are displayed in the bottom-left of the pairwise matrix while the nestedness values are in the top-right.

No Singletons

Overall Dissimilarity: 0.944 Overall Turnover: 0.894 Overall Nestedness: 0.051

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.013	0.075	0.058	0.009	0.020
Urban forest	0.911		0.031	0.099	0.005	0.107
Swamp forest	0.693	0.918		0.098	0.030	0.002
Mangrove	0.908	0.816	0.871		0.063	0.263
Freshwater swamp	0.953	0.889	0.928	0.876		0.098
Coastal forest	0.905	0.695	0.936	0.648	0.748	

No Doubletons

Overall Dissimilarity: 0.944 Overall Turnover: 0.892 Overall Nestedness: 0.052

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.015	0.078	0.060	0.010	0.020
Urban forest	0.909		0.033	0.099	0.004	0.112
Swamp forest	0.685	0.915		0.102	0.032	0.003
Mangrove	0.906	0.816	0.868		0.064	0.268
Freshwater swamp	0.952	0.889	0.926	0.875		0.101
Coastal forest	0.903	0.687	0.934	0.643	0.744	

No Species <5 Specimens

Overall Dissimilarity: 0.944 Overall Turnover: 0.891 Overall Nestedness: 0.054

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.017	0.081	0.063	0.011	0.020
Urban forest	0.905		0.037	0.099	0.004	0.118
Swamp forest	0.677	0.912		0.107	0.035	0.004
Mangrove	0.904	0.817	0.862		0.064	0.274
Freshwater swamp	0.950	0.889	0.922	0.875		0.103
Coastal forest	0.902	0.679	0.931	0.638	0.741	

No Species <10 Specimens

Overall Dissimilarity: 0.945
0.057

Overall Turnover: 0.888

Overall Nestedness:

	Rainforest	Urban forest	Swamp forest	Mangrove	Freshwater swamp	Coastal forest
Rainforest		0.023	0.082	0.069	0.014	0.018
Urban forest	0.897		0.041	0.099	0.003	0.128
Swamp forest	0.665	0.907		0.115	0.038	0.005
Mangrove	0.898	0.818	0.856		0.065	0.282
Freshwater swamp	0.944	0.890	0.917	0.876		0.105
Coastal forest	0.899	0.665	0.927	0.632	0.742	

Table S10. Rarefied species richness values and confidence intervals of data subset that could be assigned to ecological guilds.

Habitat Type	Rainforest		Freshwater Swamp Forest		Urban Forest		Mangrove Forest							
Site	Bukit Timah		Nee Soon		Kent Ridge		Pulau Ubin		Sungei Buloh		Pulau Semakau (Old)		Pulau Semakau (New)	
Ecological Guild	Mean	C.I.	Mean	C.I.	Mean	C.I.	Mean	C.I.	Mean	C.I.	Mean	C.I.	Mean	C.I.
Phytophages	181.80	±2.64	185.95	±3.25	82.50	NA	62.75	±2.23	60.60	±1.43	51.15	±1.89	31.15	±1.63
Pollinators	8.15	±0.71	9.00	±0.87	0.50	NA	13.45	±1.28	17.40	±0.76	7.40	±1.41	9.10	±1.12
Fungivores	137.95	±2.58	143.55	±2.09	44.00	NA	16.00	±1.62	19.05	±0.73	6.20	±1.05	3.10	±0.72
Parasitoids	108.15	±2.22	57.60	±2.2	114.50	NA	40.25	±2.31	53.90	±1.52	29.90	±2.68	13.75	±1.62
Predators	87.15	±2.71	98.50	±1.05	61.50	NA	130.40	±3.23	98.60	±1.63	85.95	±1.94	59.80	±2.64
Haematophages	7.80	±0.76	18.65	±0.76	0.00	NA	21.65	±0.67	22.35	±0.83	12.05	±0.86	5.95	±0.43
Detritivores	18.30	±1.37	27.15	±1.17	9.00	NA	34.20	±1.29	34.90	±0.87	26.65	±1.24	19.05	±1.01

Table S11. Species turnover ANOSIM analysis results indicate distinct communities in each habitat type for each ecological guild. Pairwise p-value outputs are displayed in the bottom-left of the pairwise matrix while the R-statistics are displayed at the top-right.

Phytophages

Overall P: 0.001

Overall R: 0.588

	Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest		1.000	0.706	0.721
Urban forest	0.001		1.000	0.468
Swamp forest	0.001	0.029		0.665
Mangrove	0.001	0.003	0.001	

Pollinators

Overall P: 0.001

Overall R: 0.836

	Rainforest	Swamp forest	Mangrove
Rainforest		0.387	0.915
Swamp forest	0.127		0.853
Mangrove	0.001	0.004	

Fungivores

Overall P: 0.001

Overall R: 0.351

	Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest		1.000	0.726	0.435
Urban forest	0.001		1.000	0.088
Swamp forest	0.001	0.029		0.432
Mangrove	0.001	0.206	0.001	

Parasitoids

Overall P: 0.001

Overall R: 0.758

	Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest		0.962	0.925	0.793
Urban forest	0.001		1.000	0.736
Swamp forest	0.018	0.067		0.711
Mangrove	0.001	0.001	0.006	

Predators

Overall P: 0.001

Overall R: 0.906

	Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest		1.000	0.414	0.954
Urban forest	0.001		1.000	0.916
Swamp forest	0.109	0.067		0.913
Mangrove	0.001	0.001	0.002	

Haematophages

Overall P: 0.001

Overall R: 0.905

	Rainforest	Swamp forest	Mangrove
Rainforest		0.435	0.957
Swamp forest	0.139		0.791
Mangrove	0.001	0.002	

Detritivores

Overall P: 0.001

Overall R: 0.853

	Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest		0.613	0.487	0.949
Urban forest	0.008		1.000	0.904
Swamp forest	0.056	0.100		0.614
Mangrove	0.001	0.001	0.002	

Table S12. Species turnover SIMPER analysis results indicate distinct communities in each habitat type for each ecological guild.

Phytophages

	Within habitat (%)	Between habitats (%)			
		Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest	30.46				
Urban forest	27.72	4.93			
Swamp forest	34.21	18.38	5.18		
Mangrove	12.37	1.29	4.13	1.67	

Pollinators

	Within habitat (%)	Between habitats (%)		
		Rainforest	Swamp forest	Mangrove
Rainforest	41.27			
Swamp forest	48.30	28.15		
Mangrove	26.01	0.88	3.57	

Fungivores

	Within habitat (%)	Between habitats (%)			
		Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest	32.01				
Urban forest	31.71	4.48			
Swamp forest	36.88	19.40	4.61		
Mangrove	10.58	1.87	8.26	1.34	

Parasitoids

	Within habitat (%)	Between habitats (%)			
		Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest	27.47				
Urban forest	10.13	3.18			
Swamp forest	59.26	10.76	1.14		
Mangrove	12.00	2.40	2.43	2.84	

Predators

	Within habitat (%)	Between habitats (%)			
		Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest	29.22				
Urban forest	34.03	4.60			
Swamp forest	64.88	19.72	3.62		
Mangrove	22.78	0.28	1.35	1.20	

Haematophages

	Within habitat (%)	Between habitats (%)		
		Rainforest	Swamp forest	Mangrove
Rainforest	18.86			
Swamp forest	56.42	10.55		
Mangrove	27.40	0.61	9.27	

Detritivores

	Within habitat (%)	Between habitats (%)			
		Rainforest	Urban forest	Swamp forest	Mangrove
Rainforest	14.99				
Urban forest	20.18	7.12			
Swamp forest	52.77	9.49	1.67		
Mangrove	18.87	0.37	1.33	6.91	

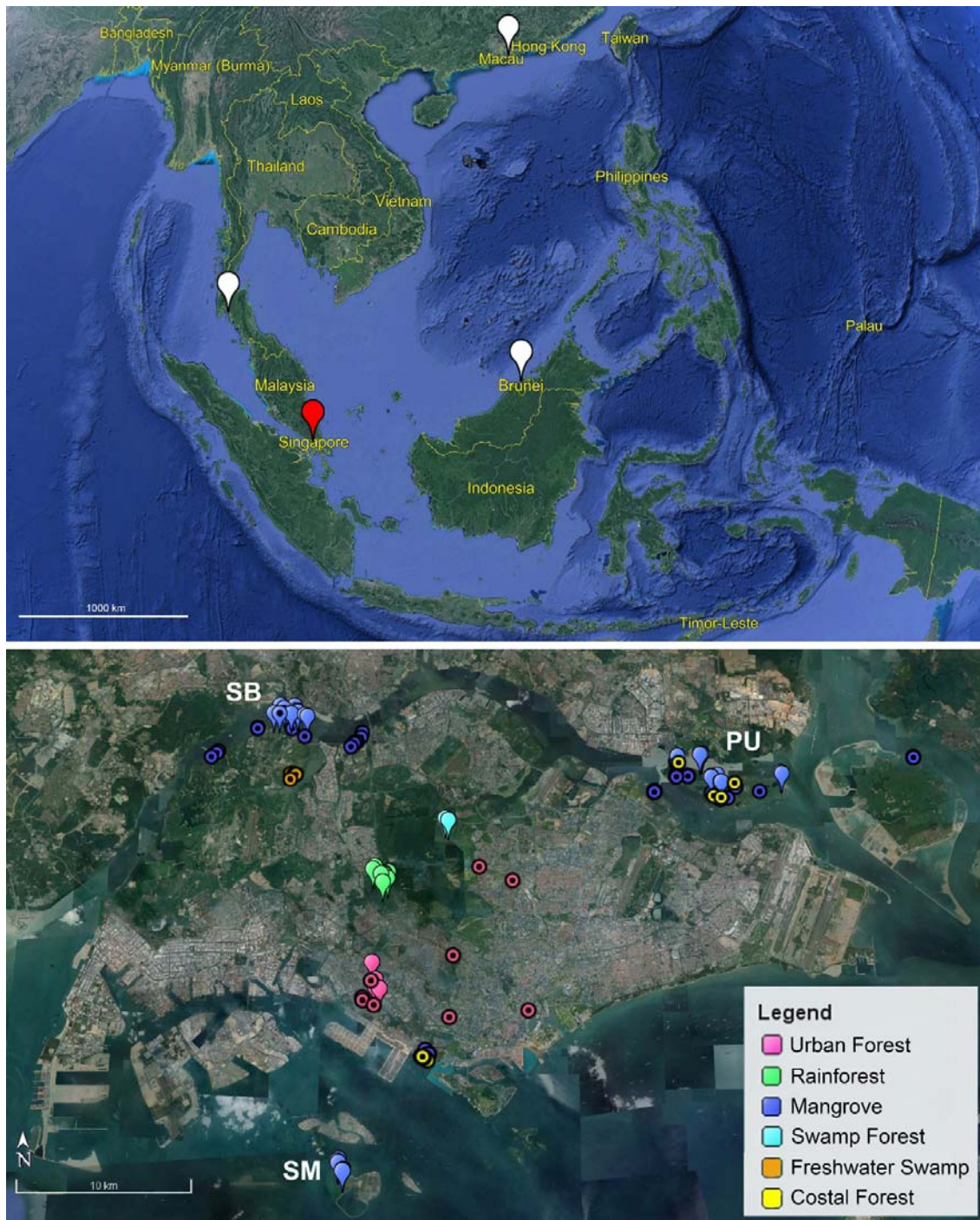
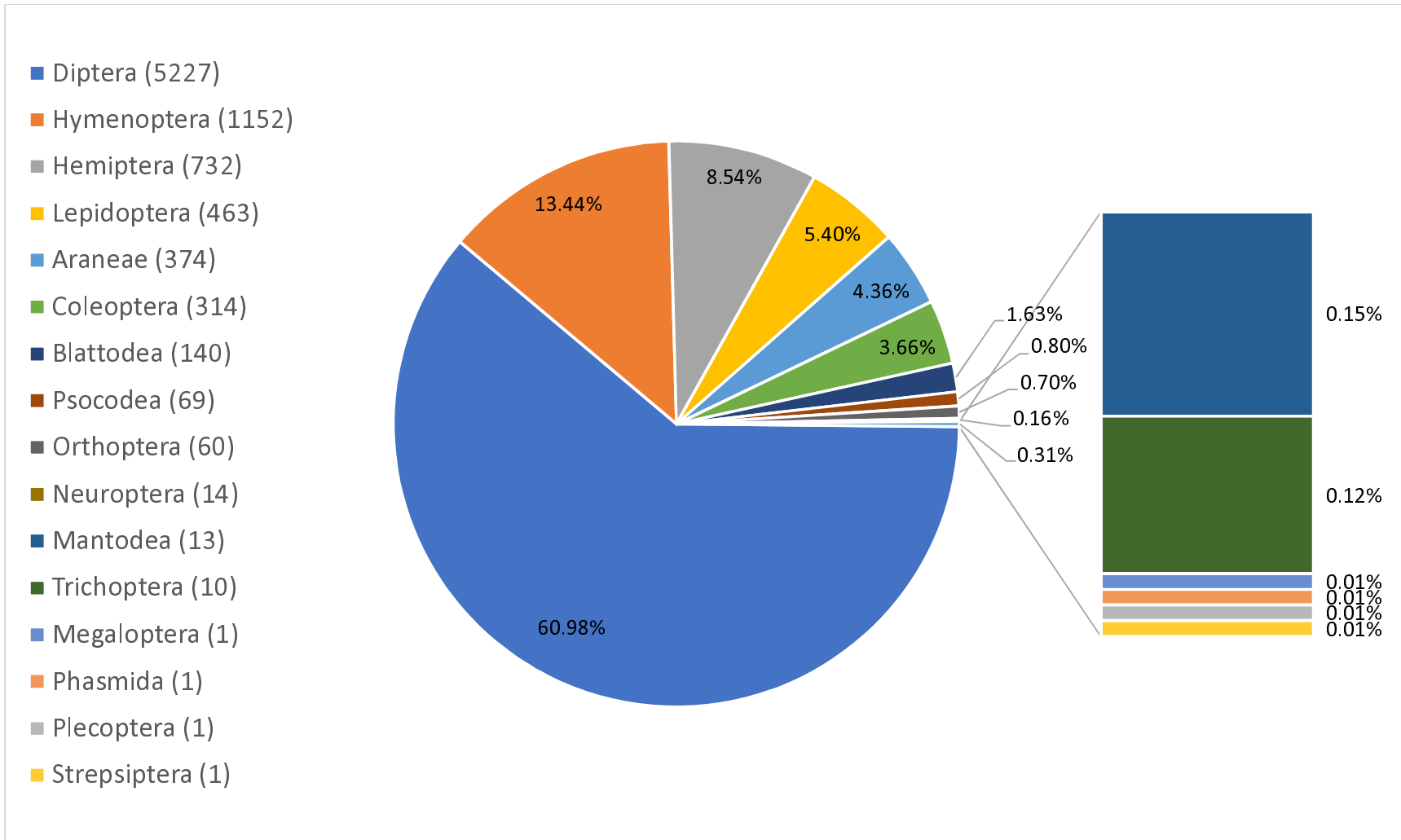


Figure S1. Sampling localities in the Oriental Realm (*top*: Singapore, red; other countries, white) and within Singapore (*bottom*: circular markers indicate trapping sites excluded from the species turnover analyses; pin markers with dot indicate traps excluded from guild-level analyses).

Figure S2. Arthropod orders sampled with Malaise traps in this study and their species proportions. The number beside each order indicates the number of species sampled based on 3% p-distance objective clustering.



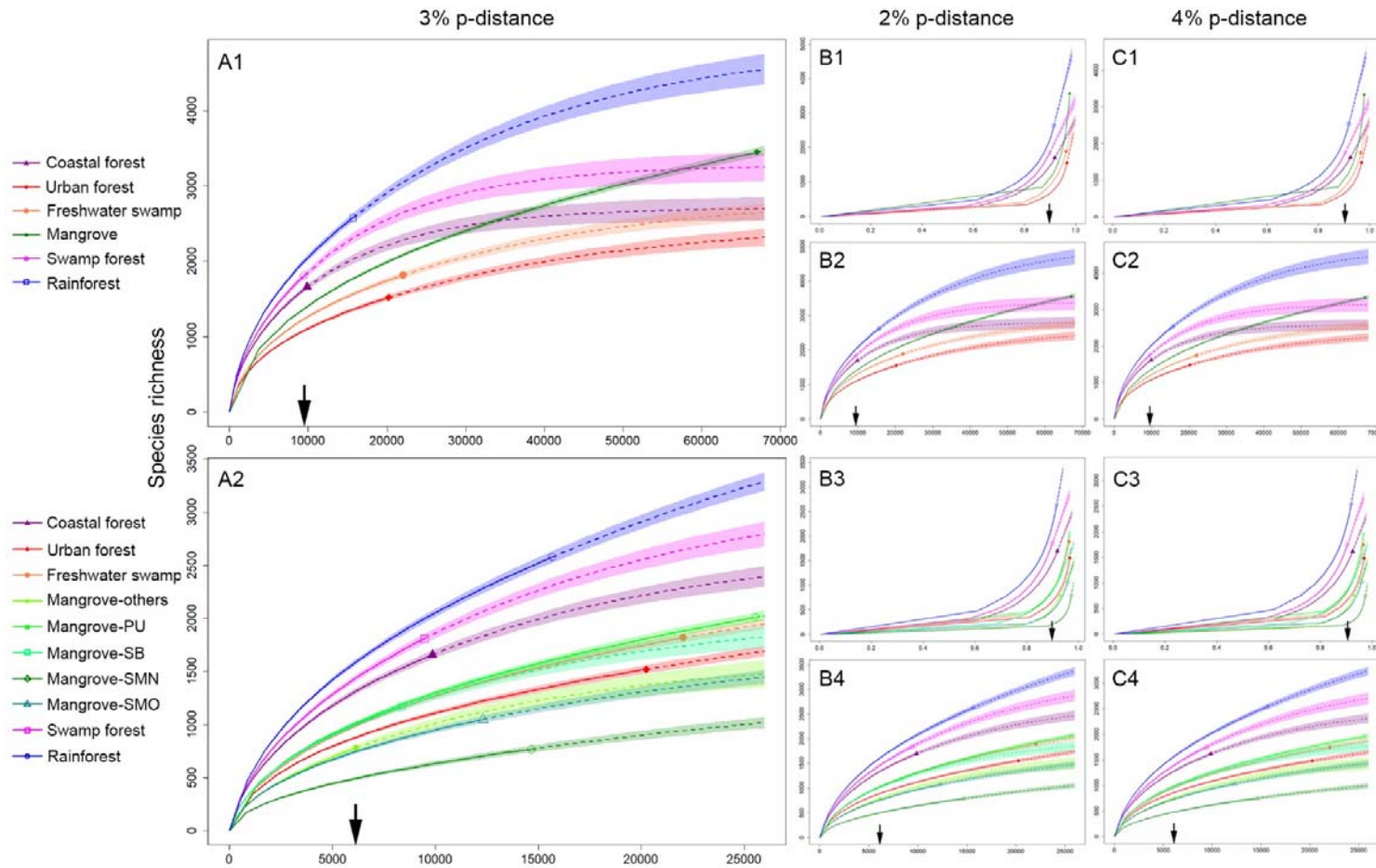


Figure S3. Insect alpha-diversity across tropical forest habitats rarefied by specimens (A1 & 2, B2 & 4, C2 & 4) and coverage (B1 & 3, C1 & 3), for 2% (B1 – 4), 3% (A1 – 2) and 4% (C1 – 4) p-distances mOTUs. Mangroves are treated as a single habitat (top) and split by site in a separate analysis (bottom): Pulau Ubin (PU), Sungei Buloh (SB), Pulau Semakau old grove (SMO), Pulau Semakau new grove (SMN); solid lines = rarefaction; dotted = extrapolations. The arrow on the x-axis indicate the point of rarefaction at which species richness comparisons were made.

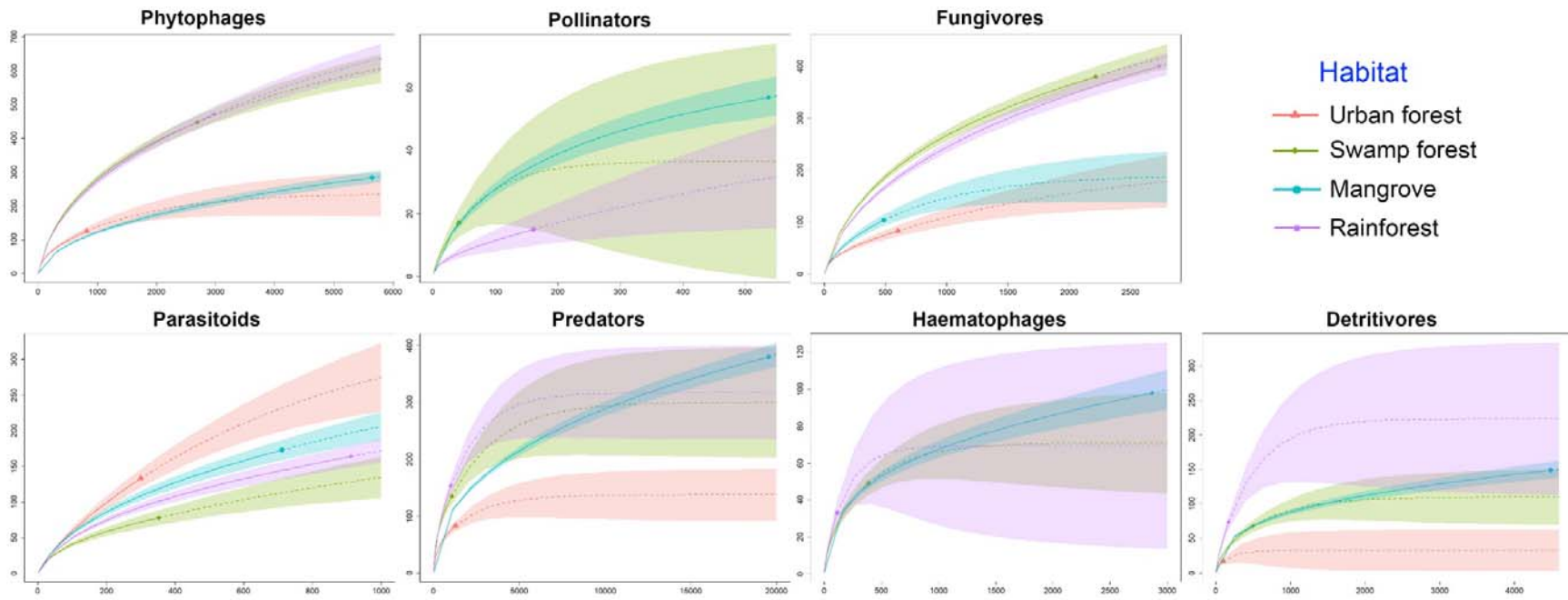


Figure S4. Comparison of species diversity across habitats (3% p-distance mOTUs) split by ecological guild. Curves were plotted for the mangrove sites as a single habitat type. The full lines represent rarefactions, while the dotted lines extrapolations and the point between the lines as actual observed values.

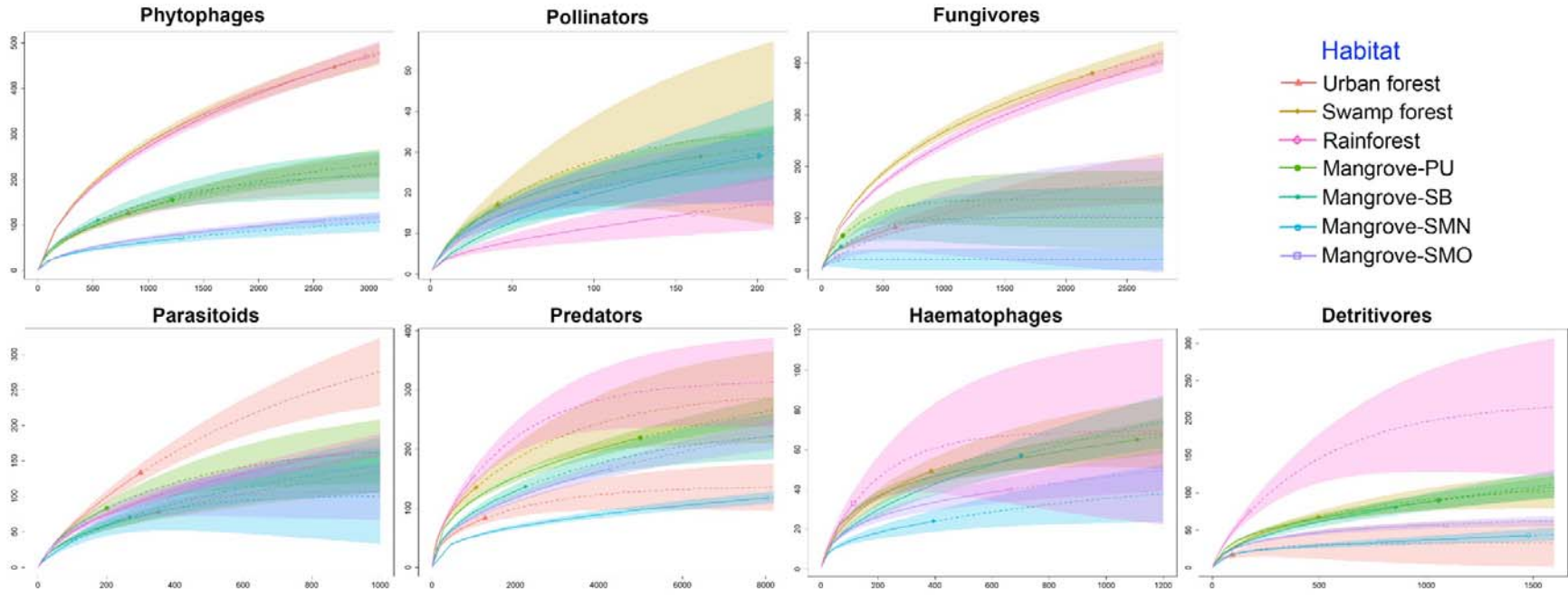


Figure S5. Comparison of species diversity across habitats (3% p-distance mOTUs) split by ecological guild. Mangrove sites are represented by Pulau Ubin (PU), Sungei Buloh (SB), Pulau Semakau old grove (SMO), Pulau Semakau new grove (SMN). Curves were plotted for the mangrove sites as separate sites. The full lines represent rarefactions, while the dotted lines extrapolations and the point between the lines as actual observed values.

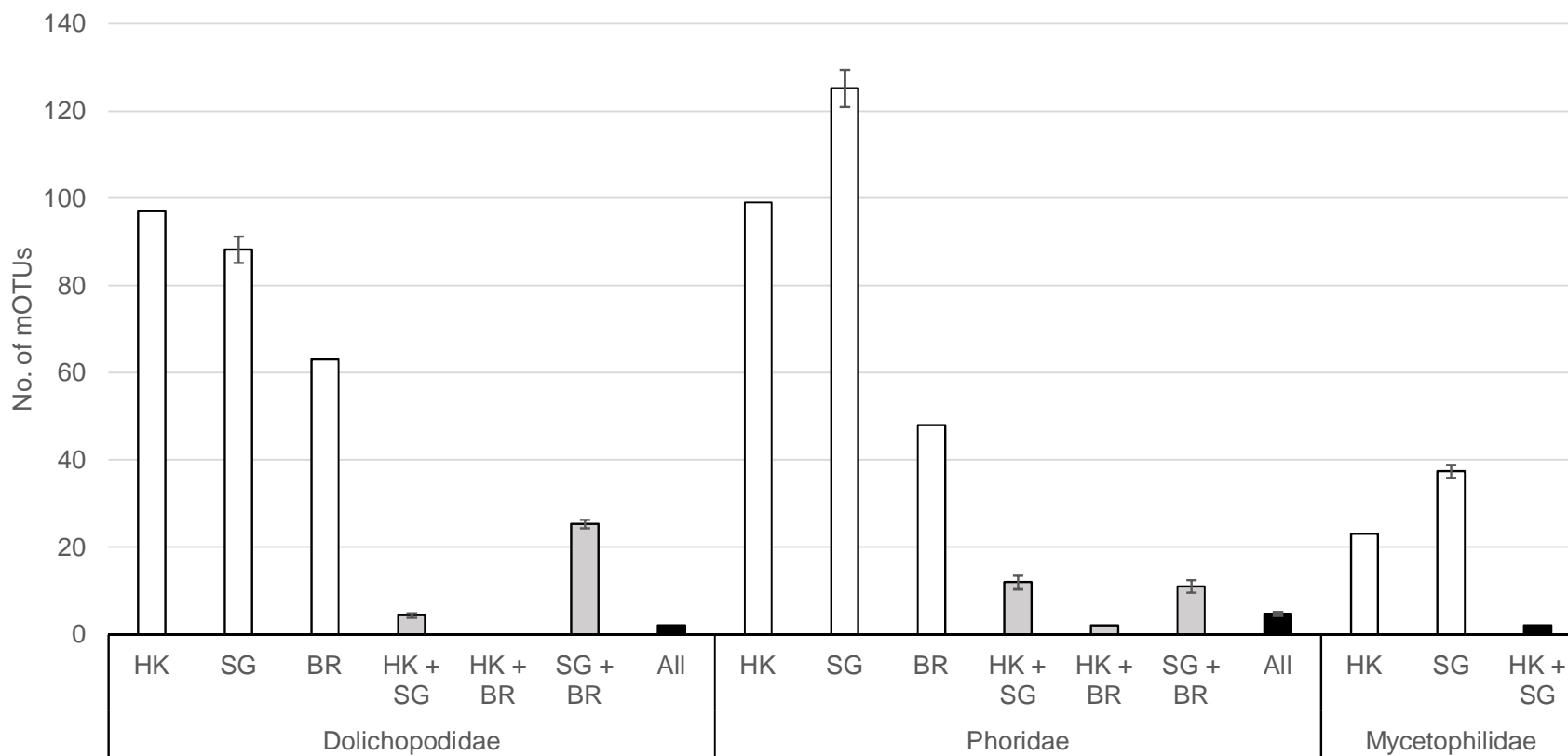


Figure S6. High species diversity and turnover for mangroves from Singapore, Brunei, and Hong Kong based on three Diptera families. Singapore data are rarefied to specimen numbers from Brunei and HK (error bars = 95% confidence intervals).

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