Why are there more arboreal ant species in primary than in secondary tropical forests?

Petr Klimes\textsuperscript{1,2*}, Cliffson Idigel\textsuperscript{3}, Maling Rimandai\textsuperscript{3}, Tom M. Fayle\textsuperscript{1,4}, Milan Janda\textsuperscript{2,5}, George D. Weiblen\textsuperscript{6} and Vojtech Novotny\textsuperscript{1,2}

\textsuperscript{1}Faculty of Science, University of South Bohemia, Branisovska 31, CZ-370 05 Ceske Budejovice, Czech Republic; \textsuperscript{2}Institute of Entomology, Biology Centre of Czech Academy of Sciences, Branisovska 31, CZ-370 05 Ceske Budejovice, Czech Republic; \textsuperscript{3}New Guinea Binatang Research Center, P.O. Box 604, Madang, Papua New Guinea; \textsuperscript{4}Forest Ecology and Conservation Group, Imperial College London, Silwood Park Campus, Buckhurst Road, Ascot, Berkshire SL5 7PY, UK; \textsuperscript{5}Museum of Comparative Zoology, Harvard University, 26 Oxford Street, Cambridge, MA 02138, USA; and \textsuperscript{6}Department of Plant Biology, University of Minnesota, 220 Biological Sciences Center, 1445 Gortner Avenue, St. Paul, MN 55108, USA

Summary

1. Species diversity of arboreal arthropods tends to increase during rainforest succession so that primary forest communities comprise more species than those from secondary vegetation, but it is not well understood why. Primary forests differ from secondary forests in a wide array of factors whose relative impacts on arthropod diversity have not yet been quantified.

2. We assessed the effects of succession-related determinants on a keystone ecological group, arboreal ants, by conducting a complete census of 1332 ant nests from all trees with diameter at breast height $\geq 5$ cm occurring within two (unreplicated) $0.32$-ha plots, one in primary and one in secondary lowland forest in New Guinea. Specifically, we used a novel rarefaction-based approach to match number, size distribution and taxonomic structure of trees in primary forest communities to those in secondary forest and compared the resulting numbers of ant species.

3. In total, we recorded 80 nesting ant species from 389 trees in primary forest but only 42 species from 295 trees in secondary forest. The two habitats did not differ in the mean number of ant species per tree or in the relationship between ant diversity and tree size. However, the between-tree similarity of ant communities was higher in secondary forest than in primary forest, as was the between-tree nest site similarity, suggesting that secondary trees were more uniform in providing nesting microhabitats.

4. Using our rarefaction method, the difference in ant species richness between two forest types was partitioned according to the effects of higher tree density (22\%6\%), larger tree size (15\%5\%) and higher taxonomic diversity of trees (14\%3\%) in primary than in secondary forest. The remaining difference (47\%6\%) was because of higher beta diversity of ant communities between primary forest trees. In contrast, difference in nest density was explained solely by difference in tree density.

5. Our study shows that reduction in plant taxonomic diversity in secondary forests is not the main driver of the reduction in canopy ant species richness. We suggest that the majority of arboreal species losses in secondary tropical forests are attributable to simpler vegetation structure, combined with lower turnover of nesting microhabitats between trees.

Key-words: diversity partitioning, Formicidae, habitat disturbance, insects, species density, tree canopies, tropical ecosystems

Introduction

Most extant arthropod species live in tropical rainforests, with the forest canopy being particularly diverse (Basset \textit{et al.} 2003; Ozanne \textit{et al.} 2003). Despite this outstanding share of global biodiversity, we know relatively little about the processes that structure canopy species diversity and distribution (Hammond 1992; Basset \textit{et al.} 2003; Ellwood & Foster 2004; Tanaka, Yamane & Itioka 2010). Such knowledge is crucial as tropical forests are increasingly being converted into structurally simpler and less diverse habitats (Primack & Corlett 2005; Cayuela \textit{et al.} 2006), with poorly understood consequences for their arthropod biodiversity (Blüh \textit{et al.} 2008).

Ants are one of the most ecologically important animal groups in many terrestrial ecosystems because of their high
abundance and complex role as ecosystem engineers, predators, herbivores, decomposers and seed-dispersion agents (Hölldobler & Wilson 1990; Folgarait 1998). Consequently, they play crucial roles in the structuring of plant and animal interactions over both evolutionary and ecological time-scales (Moreau et al. 2006). These attributes, coupled with a solid taxonomic foundation and sensitivity to environmental changes, make ants ideal group for ecological studies (Folgarait 1998; Agosti et al. 2000).

In tropical rainforest canopies, ants represent 20–60% of the entire arthropod biomass and up to 90% of individuals (Floren & Linsenmair 1997; Davidson et al. 2003; Dejean et al. 2007). Despite their considerable importance in this habitat, there have been relatively few studies on the diversity and distribution of arboreal ant communities in natural systems (e.g. Floren & Linsenmair 2000; Schonberg et al. 2004; Fayle et al. 2010; Tanaka, Yamane & Itoika 2010; Yusah et al. 2012), with most research being focused on less diverse habitats, such as plantations and managed forests (e.g. Major 1993; Philpott & Foster 2005; Pfeiffer, Tuck & Lay 2008).

Arboreal ant assemblages have been traditionally considered to be highly structured by competition hierarchies that sometimes result in spatial ‘ant mosaics’ (Major 1993; review in Dejean et al. 2007). However, the role of neutral interactions between species and stochastic events such as colonisation and extinction has recently also been recognized as important (e.g. Floren & Linsenmair 2000; Ribas & Schoech 2002; Stuntz et al. 2003; Sanders et al. 2007). Furthermore, various abiotic and biotic factors influence the arboreal ant communities, for example, the distribution of nesting sites and food resources such as extrafloral nectaries and insect symbionts (e.g. Blühgen, Stork & Fiedler 2004; Tanaka, Yamane & Itoika 2010; Powell et al. 2011). Patterns observed among tropical forests are thus often incongruent, possibly due to the limited scale of sampling and methodological differences between studies (Basset et al. 2003; Floren 2005).

Ecological research of plants in tropical forests relies heavily on sampling of entire vegetation plots (Condit 1995), whereas comparable data sets for insects are lacking. To the best of our knowledge, no study has exhaustively surveyed the ant community from trees in a continuous tropical forest area. Consequently, how arboreal ant diversity responds to changes in plant species diversity, tree size and nesting resources remains poorly known at level of entire forest communities, although such factors have been suggested to play crucial roles (Ribas et al. 2003; Campos et al. 2006; Tanaka, Yamane & Itoika 2010; Powell et al. 2011).

The diversity of ants usually declines and species composition changes predictably along disturbance gradients (Watt et al. 1997; Schulz & Wagner 2002; Dunn 2004; Floren & Linsenmair 2005; Fayle et al. 2010). However, the causal mechanisms of this decline are not fully understood. Forest disturbance causes simultaneous loss of structural heterogeneity, plant diversity and resources (Primack & Corlett 2005). Although these succession-related factors have been thought to be responsible for lower ant diversity in disturbed forests (Schulz & Wagner 2002; Blühgen, Stork & Fiedler 2004; Widodo et al. 2004; Floren & Linsenmair 2005), their relative impacts on ant species richness have not been quantified. We argue that ‘whole-forest’ inventories, that is, the complete surveys of nests in continuous forest plots, are ideal tool for investigating patterns and determinants structuring the diversity and distribution of arboreal ant communities at local scales. For instance, a plot-based approach allows direct area-based quantification and comparison of ant communities, host trees and their interactions.

In this study, a novel ‘whole-forest’ survey of felled trees and ant nests was completed in a highly diverse rainforest ecosystem in New Guinea, contrasting old-growth (primary) and early successional (secondary) forest (Whitfeld et al. 2012b). We assess how ant diversity responds to the succession-related vegetation traits, including tree density, size and taxonomic diversity and how those factors contribute to the overall differences in species richness between the two forest types. In particular, the extensive data sets on ant communities from individual trees allowed us to construct simulated forests characterized by various tree densities, size structures and taxonomic compositions and to therefore partition the impacts of these traits on ant diversity. Our study focuses on a deceptively simple question: ‘Why are there more ant species in primary than in secondary forests in the tropics?’

Materials and methods

STUDY SITE

The study field site was located in a lowland evergreen rainforest on latosols near Wanang village in the Ramu river basin, Madang province, Papua New Guinea (100–200 m. a.s.l., 05°14’S, 145°11’E), partly used for slash-and-burn agriculture (Paljmans 1976). The climate is humid with a mean annual rainfall of 3600 mm, mean air temperature of 26.5 °C and a weak dry season from July to September (McAlpine, Keig & Falls 1983).

We sampled complete arboreal ant communities from two 0.32-ha plots (40 × 80 m), each of which formed part of a larger 1-ha study plot (Whitfeld et al. 2012b). One plot was located in an undisturbed primary forest with a canopy height up to c. 50 m and the other in a secondary forest with canopy reaching height of 25 m, which represented c. 10 years of successional vegetation growth on an abandoned food garden. The primary forest 1-ha plot included 1336 stems with diameter at breast height (DBH) ≥ 5 cm and a basal area of 299 m², representing 213 tree species from 130 genera and 54 families. The secondary forest 1-ha plot included 1206 stems with DBH ≥ 5 cm and a basal area of 136 m², representing 90 species from 67 genera and 29 families (Whitfeld et al. 2012b).

Both plots were chosen in cooperation with the indigenous landowners who practise subsistence swidden agriculture in their forests. This partnership allowed us to sample forest plots by felling all trees without contributing to further deforestation, as the plots were already scheduled to be felled to create food gardens. On the contrary, our research provided income and job opportunities for the village community and encouraged villagers to conserve their primary
rainforest, which is under threat from commercial logging, for further research (Novotny 2010).

**SAMPLING DESIGN AND MATERIAL COLLECTION**

Between February and November 2007, all trees with DBH ≥ 5 cm were felled, measured, identified to species and searched for ants in the primary (trees N = 389) and secondary (trees N = 295) plots. For each tree, we measured DBH, trunk height (to first branch), crown width, crown height and total fresh leaf weight.

Every felled tree was searched intensively for ant nests and foraging individuals by three collectors from its base to the top branches. Destructive sampling enabled the cutting of branches, attached lianas and leaves and the dissecting of parts of the trunk, bark and epiphytic aerial soil to record the complete ant fauna nesting on each tree. Each nest was classified according to its location and structure into the following categories (nest sites hereafter): in aerial soil, on bark, under bark, in trunk cavity, in dead twig (≤ 5 cm in diameter), in dead branch (≤ 5 cm in diameter), in live twig, in live branch, on leaves, on/in liana and in myrmecophytic plant.

Only nesting ants (with a queen and/or immature life stages) were included in the analysis in this study. Foraging ants were not considered as these came also from nests on the ground or in surrounding vegetation (P. Klimes, unpublished data), whilst our focus in this study is on the ant diversity closely associated with individual trees, tree species and nest sites.

**SPECIES IDENTIFICATION**

Several individuals from each nest, including all castes, were preserved in ethanol and later sorted to genus using Bolton (1994). Morphospecies were further determined using collections at the Institute of Entomology (the Czech Academy of Science) and the Museum of Comparative Zoology (MCZ), online databases (http://www. antweb.org; http://www.newguineants.org) and with the assistance of specialist taxonomists (see Acknowledgements). DNA barcodes (sequences of COI gene) were obtained for approximately half of the species and compared with c. 2000 sequences already available for New Guinea species (http://www.formicidaebol.org). The combination of morphological and molecular data was used to define species boundaries. Tree species were identified following Whitfield et al. (2012a).

**DATA ANALYSIS**

Our analysis focused on ant assemblages on individual trees, which constitute replicated data points within each plot. The logistical demands of the whole-forest survey, comprising all trees within 0.32-ha plots, did not allow for plot replication, that is, our data used for the comparison between primary and secondary forest are pseudoreplicated (Huribert 1984). This is a common problem for whole-ecosystem studies and studies on super-abundant and diverse taxa. It has been argued that when large scale has priority over replication (as in this study), cautious use of inferential statistic may be acceptable (Oksanen 2001; Chaves 2010).

**Predictors of ant species diversity**

Tree-based species accumulation curves were used to explore the relationships between ant species richness and the number of trees in each forest type (Mao Tau function). The total expected number of ant species for each community was estimated using the Chao2 index, in estimates v. 8.2 with 100 randomisations of sample order (Colwell 2009). As we conducted complete censuses of the ant nests within the plots, this index estimates here the richness of the local species pools, rather than the number of unsampled species within each plot.

The size-related tree traits (DBH, tree height, crown and trunk height and total fresh leaf weight) were first tested as predictors of ant diversity using multiple stepwise regression. Both dependent and explanatory variables were log-transformed prior to analysis to meet the assumption of normality and homoscedasticity of residuals and to improve the linearity because of the multiplicative effect of explanatory variables (allometric dependency). As DBH was found to be the best predictor of the number of ant species on trees (R² = 0.22, P < 0.001) and accounted for 85% of the total variability explained by all size traits, we used it hereafter as a surrogate of tree size (see Table S1, Supporting information for full model details).

Relationships between tree size and ant diversity, and nest density and diversity of nest sites were compared between the two forest types using a GLM model of homogeneity of slopes (effects: forest type, DBH). Mean ant species richness, nest number and richness of nest sites per tree were compared between plots using ANOVA. When needed, the variables were log-transformed prior to analyses for data normality and homoscedasticity. Mean species richness per tree was also calculated separately for different tree sizes. Trees were classified into size classes based on their log(DBH), starting from the minimum value of 0.70 (for DBH = 5 cm) and using class width of 0.1625 (i.e. one-eighth of entire range: 0.70–0.87, 0.87–1.02, etc.), except for the last class that included all trees with log(DBH) > 1.62. Analyses were performed in **STATISCA** software ver. 9.1 (StatSoft 2010).

The similarity of ant communities between trees within each forest plot was characterized by the Sørensen similarity index (SI), ranging from zero (no ant species shared by the compared communities) to one (all species shared). The Sørensen index was used also to estimate the similarity in nest sites between trees, using nest site category instead of species.

The correlation of taxonomic distance between trees and the dissimilarity of their ant assemblages (estimated as 1 – SO) was assessed using Mantel tests computed in **zt** software ver. 1.1 with 1000 runs per analysis (Bonnet & Van de Peer 2002). Trees without nests were excluded from the analysis because SO is not defined for two empty samples. Taxonomic distance between pairs of trees within each plot was coded at four levels of resolution, each of which was analysed separately: (i) full taxonomic distances where 0 = conspecific, 1 = congeneric, 2 = confamilial, 3 = allofamilial trees; (ii) familial distances, 0 = confamilial, 1 = allofamilial trees; (iii) generic distances, 0 = congeneric, 1 = allogeneric trees; and (iv) species distances, 0 = conspecific, 1 = allspecic trees. Finally, Mantel test was used in the same way to assess the correlation between ant species dissimilarity and dissimilarity of nest sites between trees in each plot.

**Partitioning of the differences in ant species richness**

Our data on ants and trees from plots in primary and secondary forest offered a unique opportunity to examine the effect of individual forest characteristics on ant species diversity. We selected subsets of trees from the primary forest in such a way that they matched those from secondary forest in terms of tree density, size distribution and taxonomic diversity and examined the effects of these matched parameters on ant diversity. Additive partitioning of diversity was used to explain the difference in ant species richness between the primary and secondary forests, where total ant species richness per plot (S_tot) is defined as the sum of species richness per tree (S_t), and beta diversity of species between trees (S_j) (Anderson et al. 2011).
First, we divided each study plot into three continuous 0.1-ha subplots (40 × 25 m each) and calculated the mean number of trees per 0.1 ha, as 96 (±22.2) trees in the secondary and 122 (±93) trees in the primary forest (Table 1). Further, we randomly selected 500 model sets of 96 trees from the entire 0.32 ha secondary forest and 500 sets of 122 trees from the 0.32 ha primary forest plots, matching thus the tree number in each set to the mean number of trees per 0.1 ha in each respective forest type. The 500 model sets of trees from secondary forest were then used as templates in the following rarefaction analyses.

The effect of tree density on ant diversity was examined by randomly reducing the number of selected trees in the primary forest data set (122 trees) to that found in the secondary forest template, that is, 96 trees. In the next step, we controlled also for tree size by selecting 96 primary forest trees such that their distribution among the DBH size classes matched the secondary forest template. Finally, we controlled for tree taxonomic diversity by selecting 96 trees whilst matching both tree size and plant diversity of secondary forest template. Specifically, we matched the relative abundance distributions of tree species within size classes, the numbers of species within each size class and the total number of tree species, genera and families across all size classes. Each of these three steps was repeated 500 times for different secondary forest template data sets. Although the identity of species, genera and families differed between primary and secondary data sets, the distribution of individual trees among taxonomic categories in the primary forest set matched very closely that in the model secondary forest set of trees (Table 1; Table S2, Supporting information). To match spatial scales for the primary and secondary data sets, and to make full use of the data available, we used randomly drawn trees (within the above constraints) from the entire 0.32-ha plots. For further details of the algorithms used and annotated R code, see Appendices S1–S2 (Supporting information).

The simulated primary forest data sets were then used to partition the difference in species diversity of ants between the model sets of trees from the primary and secondary forests into the effect of (i) tree density, (ii) tree size and (iii) tree taxonomic diversity (Table 2). The remaining difference in species richness, unexplained by these three factors, was partitioned into differences in alpha diversity of individual trees and beta diversity (i.e. species turnover) between trees.

### Results

**Predictors of ant species diversity**

We sampled a total of 1332 ant nests of 99 species from 684 trees in the two 0.32 ha rainforest plots combined. The primary forest plot was almost twice as rich in ant species as the secondary forest plot with 80 species nesting on 389 trees, compared with only 42 species on 295 trees in the secondary forest plot (Table 1). No nests were found on 30% and 23% of trees in the primary and secondary plot, respectively (usually trees with DBH ≤ 15 cm).

Accumulation curves of ant species observed in both primary and secondary forest plots did not reach asymptotes. This contrasted to the Chao2 estimates of total local species richness of 890 (SD = 5.56) species in primary forest and 526 (SD = 7.7) species in secondary forest (Fig. 1). Species nesting on only a single tree (uniques) or on two individual trees (duplicates) comprised together almost half of the species in each plot as there were 19 uniques and 18 duplicates in the primary forest plot and 13 uniques and 8 duplicates in the secondary forest plot. The most common species in primary forest, *Crematogaster polita* Smith F., nested on 96 trees (25%) and the most common secondary forest species, *Camponotus aff. macrocephalus* (Erichson), on 112 trees (38%).

### Table 1. Characteristics of primary and secondary forest vegetation and their associated ant communities in (i) the whole 0.32-ha plots, (ii) three continuous 0.1-ha subplots (mean ± SE), (iii) model sets of trees randomly selected from the 0.32-ha plots to match the mean number of trees in 0.1 ha (means for 500 selections) and (iv) randomly selected sets of primary forest trees matching the model sets from secondary forest in the number of trees, in the number and size [diameter at breast height (DBH)] of trees and in the number, size and taxonomic diversity of trees, respectively (means for 500 templates). Overall fit in tree taxonomy was achieved as follows (mean percentage match in number of taxa of simulated set compared with template over 500 replications): species = 100%; genera = 99%; families = 98.6% (See Methods and Appendix S1, Supporting information for details)

<table>
<thead>
<tr>
<th>Forest</th>
<th>Tree number</th>
<th>Tree sp. richness</th>
<th>Tree gen. richness</th>
<th>Tree fam. richness</th>
<th>DBH per tree</th>
<th>Nest number</th>
<th>Ant sp. richness</th>
<th>Ant sp. richness per tree</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Secondary forest</strong></td>
<td></td>
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<tr>
<td>Whole plot (0.32 ha)</td>
<td>295</td>
<td>47</td>
<td>33</td>
<td>19</td>
<td>11.3</td>
<td>580</td>
<td>42</td>
<td>1.48</td>
</tr>
<tr>
<td>Continuous 0.1-ha plot</td>
<td>96</td>
<td>26.9</td>
<td>19.3</td>
<td>13.5</td>
<td>11.3</td>
<td>188.6</td>
<td>27.5</td>
<td>1.58 ± 0.26</td>
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<tr>
<td>(mean ± SE)</td>
<td>960 ± 22.2</td>
<td>253 ± 30</td>
<td>177 ± 3.7</td>
<td>127 ± 22</td>
<td>116 ± 0.9</td>
<td>187 ± 20</td>
<td>25.3 ± 3.9 (a)</td>
<td></td>
</tr>
<tr>
<td>Model set of trees (0.1 ha)</td>
<td>96</td>
<td>26.9</td>
<td>19.3</td>
<td>13.5</td>
<td>11.3</td>
<td>188.6</td>
<td>27.5</td>
<td>1.48 (f)</td>
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<tr>
<td><strong>Primary forest</strong></td>
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<tr>
<td>Whole plot (0.32 ha)</td>
<td>389</td>
<td>115</td>
<td>78</td>
<td>41</td>
<td>12.7</td>
<td>752</td>
<td>80</td>
<td>1.52</td>
</tr>
<tr>
<td>Continuous 0.1-ha plot</td>
<td>122 ± 9.3</td>
<td>573 ± 3.4</td>
<td>460 ± 0.6</td>
<td>293 ± 0.7</td>
<td>128 ± 0.2</td>
<td>239 ± 53</td>
<td>47.3 ± 8.4 (a)</td>
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<tr>
<td>(mean ± SE)</td>
<td>122 ± 9.3</td>
<td>573 ± 3.4</td>
<td>460 ± 0.6</td>
<td>293 ± 0.7</td>
<td>128 ± 0.2</td>
<td>239 ± 53</td>
<td>47.3 ± 8.4 (a)</td>
<td></td>
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<td>Model set of trees (0.1 ha)</td>
<td>122</td>
<td>61.7</td>
<td>490</td>
<td>296</td>
<td>12.7</td>
<td>235.5</td>
<td>52.7 (b)</td>
<td>1.52</td>
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<tr>
<td>Selection of primary forest trees simulating the secondary model set of trees</td>
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<tr>
<td>Matching tree density</td>
<td>96</td>
<td>52.9</td>
<td>43.4</td>
<td>27.3</td>
<td>12.7</td>
<td>1849</td>
<td>47.0 (e)</td>
<td>1.52</td>
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<td>Matching tree density and size</td>
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<td>53.1</td>
<td>43.5</td>
<td>27.3</td>
<td>11.3</td>
<td>1739</td>
<td>43.3 (d)</td>
<td>1.45</td>
</tr>
<tr>
<td>Matching tree density, size and taxonomy</td>
<td>96</td>
<td>27.1</td>
<td>19.6</td>
<td>13.6</td>
<td>11.2</td>
<td>176.1</td>
<td>39.5 (e)</td>
<td>1.48 (g)</td>
</tr>
</tbody>
</table>

The distribution of trees in size classes differed between the plots ($\chi^2 = 27.3, \text{d.f.} = 6, P < 0.001$) with secondary forest having relatively fewer of the smallest trees, more of the middle size trees and no trees with DBH over 44 cm (Fig. S1, Supporting information). Other vegetation characteristics also differed considerably between the forest types (Table 1). However, despite these disparities, the number of ant species increased in the same way with tree size (DBH) in the two forest types (GLM: $F_1 = 1.27, P = 0.26$; Fig. 2a; Table S3, Supporting information). Similarly, there was no difference between the two forest types in the relationship between tree size and the number of nests (GLM: $F_1 = 0.65, P = 0.42$) and also between tree size and diversity of nest sites (GLM: $F_1 = 1.39, P = 0.24$) (Fig. S2 and Table S3, Supporting information).

Individual trees hosted 0–12 ant species (mean 1.52 ± 1.67) in the primary forest plot and 0–8 species (mean 1.48 ± 1.34) in the secondary forest plot. The mean number of ant species per tree did not differ between primary and secondary forest trees for all tree sizes combined ($F_{1,682} = 0.56, P = 0.45$). However, it varied considerably among tree size classes from one to six species per tree on average (Fig. 2b). The number of nests, 0–20 (mean 1.93 ± 2.37) per primary forest tree and 0–13 (mean 1.97 ± 2.11) per secondary forest tree, was also not significantly different between the two forest types ($F_{1,682} = 1.1, P = 0.29$). The number of nest site categories varied from one to five per tree in the both forest types.
(Fig. S2b, Supporting information) and also did not differ between primary forest (mean 1.23 ± 1.07) and secondary forest (mean 1.24 ± 0.99) ($F_{1,682} = 0.05, P = 0.83$).

The mean similarity of ant species composition between pairs of trees in secondary forest, $S\text{S}_\text{mean} = 0.10$, was approximately double of that in primary forest ($S\text{S}_\text{mean} = 0.04$). Moreover, the difference was evident in all size classes (Fig. S3, Supporting information), indicating that there is a robust pattern of higher ant species turnover between trees in primary than in the secondary forest. Similarity of ant species composition between pairs of conspecific trees was higher than between heterospecific trees in both forests (Fig. 3a). The highest mean species similarity was found between conspecific trees of the secondary forest ($S\text{S}_\text{mean} = 0.14$). The similarity patterns for nest sites were analogous to those for ant species (Fig. 3b), with the highest similarity again found between conspecific trees in the secondary forest ($S\text{S}_\text{mean} = 0.21$).

The taxonomic distance between trees and the dissimilarity of their ant communities were positively correlated with both forest types at all levels of taxonomic resolution: family, genus, species (Mantel test, $P = 0.001$) with exception of the familial level in primary forest (Mantel test, $P = 0.102$) (Table 3). The relationship was strongest at the tree species level in both forests, but the correlation was extremely low in all cases with weaker correlations in primary than in secondary forest ($R$: primary forest: range = 0.02–0.03; secondary forest: range = 0.04–0.09) (Table 3). The dissimilarity in ant species composition was positively correlated with the dissimilarity of their nest sites in both habitat types (Mantel test, $P = 0.001$). These correlations were much stronger than those between dissimilarities of ant communities and taxonomic distances between trees (primary forest: $r = 0.22$, $P = 0.001$; secondary forest: $r = 0.24$, $P = 0.001$).

**Partitioning of the differences in ant species richness**

The primary forest comprised on average 122 tree individuals and 473 ant species within a 0.1-ha plot, whilst the secondary forest included 96 tree individuals and 253 ant species in the same area (Table 1). The mean ant species richness of the sets of 122 primary forest trees randomly drawn from 0.32 ha of the forest comprised 527 ant species whilst for the sets of 96 secondary forest trees drawn from 0.32 ha the mean was 275 ant species. Our simulations showed that the difference of 252 ant species between the primary and secondary model sets could be explained by higher tree density (57 species), larger tree size (39 species) and higher tree taxonomic diversity (36 species), with the remaining difference being attributable to higher beta diversity between trees (120 species) in primary forest, as the simulated forests did not differ in the number of ant species per tree (alpha diversity) from the secondary forest data sets (Table 2, Fig. 4). The difference in the density of ant nests was explained completely by differences in tree density between the two forest types (see values for nest numbers in Table 1).

**Discussion**

With 684 trees sampled in total, this is currently the most extensive study of arboreal ants from a tropical rainforest. As expected, we found a much lower diversity of ants in the secondary than in the primary forest plot. The ant species richness in secondary forest was approximately half that of in primary forest, and this difference was proportionally similar at both 0.32 and 0.1 ha scales. Previous studies also usually found large differences in species richness between primary and secondary forests (Schonberg et al. 2004; Floren & Linsenmair 2005; Bihn et al. 2008; Klimes et al. 2011), although less dramatic effects of human disturbance on rainforest ant diversity have been also reported (e.g. Schulz & Wagner 2002; Woodcock et al. 2011). However, all these studies found considerable change in ant species composition during rainforest succession. In our study, the species composition was also very different as there were only 23 shared species between the two plots, and different species dominated each forest type at the site (Klimes et al. 2011).

Surprisingly, and in contrast to overall diversity in plots, mean ant species richness per tree did not differ between forest plots. We expected higher species richness on primary trees than secondary trees, because the higher structural complexity of vegetation and availability of epiphytes and climbers should provide more nesting opportunities for ant species (Ribas et al. 2003; Campos et al. 2006; Turner & Foster 2009; Tanaka, Yamane & Itioka 2010). Indeed,
Floren & Linsenmair (2005) found twice as many ant species per fogged tree in primary than in early successional rainforests. However, this pattern was not replicated in our study. As we specifically noted nesting species, we are confident that these results reflect the actual ant occupancy of studied trees. The relatively high proportion of unique species in our samples reflects genuinely low nest density of many species in trees, although some of these might be more commonly found at ground level.

The overall ant species diversity of 99 species sampled from 684 trees and the mean number of species per tree (1·5) were much lower compared with other studies of arboreal ant communities in tropical rainforests, for example, 231 spp. on 30 trees in Borneo (Floren & Linsenmair 2005) and 161 spp. on 61 trees in Africa (Schulz & Wagner 2002) and 4–40 ant species per tropical tree on average (Wilson 1987; Floren & Linsenmair 1997, 2000, 2005; Schulz & Wagner 2002; Schonberg et al. 2004). However, we argue that these estimates are not comparable to our study of arboreal nesting ants. First, these studies were based primarily on canopy fogging so that their diversity estimates include also numerous ant foragers from nearby vegetation and from the ground (see Hammond 1992; Schulz & Wagner 2002; Floren 2005). Second, previous research focused on small numbers of trees of particular size, selected over a larger area, whilst we sampled entire patches of vegetation within a rather small area (two 0·32-ha plots). It is thus not surprising that species richness in our study is lower. Although comparable data are scarce, several other studies found similar or even lower number of ant nesting species per tree with DBH over 40 cm in tropical forests (Tanaka, Yamane & Itooka 2010; Janda & Konecna 2011). Another recent study reported that arboreal species were present in only 56–74% of trees in transects within a continuous rainforest (Dejean et al. 2010), a similar rate of occupancy to that which we found. Hence, we believe that our rainforest site is representative for arboreal ant communities of a highly species-rich tropical region such as New Guinea.

A strong positive correlation of both nest density and species richness with tree size was observed in both forest types. This was expected as tree size has been shown previously to be an important predictor of ant richness and number of colonies (e.g. Campos et al. 2006; Powell et al. 2011). Recently, Powell et al. (2011) also demonstrated that there is a stronger correlation between ant species richness and tree size in forests of higher tree density than in less dense forests. However, this pattern is not universal as demonstrated by our surprising result that arboreal ant diversity, nest density and micro-habitat richness varied in the same way with tree size in both primary and secondary forest plots in our study, despite differences between the plots not only in tree density, but also in tree size-range distribution.

The diversity of plants is thought to be an important factor in maintaining the diversity of ant communities (Ribas et al. 2003; Moreau et al. 2006; Dejean et al. 2007) and insects in general (Novotny et al. 2006). It is also well known that in many myrmecophytic plants the composition of ant communities can differ between host tree species as a consequence of shared evolutionary plant-ant history or because of the variation in the provision of food and nesting resources (Blüthgen, Stork & Fiedler 2004; Dejean et al. 2007). However, myrmecophytic plants often represent only a...
fraction of local tree diversity and studies exploring ant fauna from locally common tree species usually find no significant effect of tree species identity on tropical ant communities (e.g. Floren & Linsenmair 1997; Schulz & Wagner 2002; Janda & Konecna 2011). Here, we demonstrate that the effect of plant taxonomic diversity on ants is significant at the ‘whole-forest’ level and that ant communities are more similar to each other on conspecific than on heterospecific trees. Our results also imply that arboreal ant communities are relatively more specialized to plant species (or genera) than families. Nevertheless, the overall effect of plant taxonomy was very small in both habitats suggesting that a higher taxonomic diversity of primary vegetation does not itself explain why ant diversity in primary forest is twice that of secondary forest. Our finding for arboreal ants is congruent with a recent analysis of leaf-litter fauna that also showed a low impact of plant diversity on rainforest ants (Donoso, Johnston & Kaspari 2010). Nevertheless, the future challenge is to explore also the affiliation between individual tree taxa and particular ant species relating to other environmental variables, for example, the presence of extrafloral nectaries, homopteran symbionts and spatial interactions (Davidson et al. 2003; Blüthgen, Stork & Fiedler 2004) and to expand our approach to include phylogeny (rather than taxonomy) of both ant and plant communities (Whitfeld et al. 2012b).

The differences in ant species diversity between the two forest plots were primarily due to variation in beta diversity (species turnover) of ant communities between trees as we did not found any difference in mean alpha diversity of ants per individual tree (Fig. 4). Indeed, the similarity of ant communities among primary forest trees was much lower than in secondary trees. Other studies have also noted high species turnover between rainforest trees, but they have not found any common mechanism contributing to these patterns (Floren & Linsenmair 1997, 2000; Stuntz et al. 2003; Yanoviak, Fisher & Alonso 2008). Here, we were able to explicitly assess the contribution of particular succession-related determinants on ant diversity by comparing the species richness between model data sets of primary and secondary forest trees. Intriguingly, our results demonstrate that even when both forest types are matched for tree abundance, tree size and taxonomic structure, there is still a much higher beta diversity of ant communities between primary forest trees than secondary forest trees, accounting for almost half (47.6%) of the difference in species richness. We suggest that this unexplained fraction probably relates to the markedly higher levels of microhabitat heterogeneity previously reported between trees in primary forests (Floren & Linsenmair 2005; Turner & Foster 2009) as we found a much higher similarity of ant nest sites among secondary trees than among primary trees (Fig. 3b). These results imply that secondary forest trees were more uniform in the nesting opportunities they provided compared with trees in primary forest, because numerous associated epiphytes and lianas increased tree-to-tree variability (microhabitat turnover) in primary forest. Indeed, large climbers (DBH ≥ 5 cm) were present only in the primary forest plot, and one of the most frequent nest sites in that habitat was aerial soil, which was not the case in the secondary forest plot (P. Klimes, unpublished data). The high importance of nesting resources for ants suggests also that the correlation in composition of ant communities with their nest site composition was much stronger than correlation with tree taxonomic composition. However, rather unexpectedly, we found the same nest density and nest site richness per tree (within-tree variability) in both primary and secondary forest. This may explain why we recorded the same alpha diversity of ants per tree in the two habitats, because canopy ant communities are thought to be limited by nest site variability and availability (Philpott & Foster 2005; Powell et al. 2011).

We thus did not find strong support for the conclusion of Ribas et al. (2003) that plant diversity and density are the most important factors for maintaining ant diversity at local scales. Analogous studies to our own from other tropical regions are needed to evaluate how the effects of these factors might vary between tropical ecosystems. Nevertheless, our results agree with those from structurally simpler environments such as savannas and coffee plantations, where nest site diversity has been shown to be important in sustaining high diversities of twig-dwelling ants (Philpott & Foster 2005; Powell et al. 2011).

Our results have potential implications for the conservation of rainforest ant communities. As vegetation structure was more important than tree diversity, current forest regeneration efforts (e.g. Hector et al. 2011) should focus specifically on recreating high canopy forest that emulates primary forest in terms of structure. For ants at least, high tree species diversity is not a priority. Indeed, selectively logged forests, in which particular tree species are removed, but the overall forest structure is left relatively unchanged, support rather similar diversity of ant communities as primary forests (Widodo et al. 2004; Woodcock et al. 2011). However, other arboreal arthropods such as herbivorous larvae show much greater host tree specificity (Novotny et al. 2006), and so conservation priorities are likely to be different for these groups.

The data presented are exceptional in that they represent full censuses of ant nests from all trees within continuous areas of tropical forest. We argue that this type of data, although difficult to obtain, is crucial for further studies of insect communities. Likewise, our understanding of the composition and dynamics of rainforest vegetation is based to a large extent on the analysis of census data from continuous plant plots (Condit 1995; Leigh et al. 2004). Insect data from continuous plots allow rigorous comparisons of various aspects of insect communities between different forest types, as demonstrated here for primary and secondary forests. Nevertheless, we acknowledge that our results should be interpreted with caution, as sampling units (trees) were replicated within plots, which were not themselves replicated. The challenge for further studies is to increase the size of each plot and also to replicate plots within each forest type. In particular, the effect of tree density on arboreal ants requires further research as data from replicated plots show that stem density...
varies both between individual plots of each forest type and among the secondary successional stages (Whitfield et al. 2012a).

In conclusion, we have used perhaps the most complete existing data sets of arboreal ant communities to show that their species and nest density per tree scale universally with tree size in both primary and early successional tropical forests, despite large differences in their overall ant species diversity. Furthermore, we have demonstrated that approximately half of the impact of habitat conversion on ant species richness relates to decreases in tree abundance, a shift in tree size distribution and a simplification of tree taxonomic structure. The remaining effect is entirely at the level of ant beta diversity between trees and probably relates to a reduction in the turnover of nesting sites between trees. These findings suggest that plant taxonomic diversity is not the main driver of canopy ant diversity.

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Supporting Information

Additional Supporting Information may be found in the online version of this article.

Fig. S1. Distribution of trees in individual size classes in primary and secondary forest.

Fig. S2. Relationships between tree size, number of ant nests and diversity of nest sites.

Fig. S3. Similarity in ant species composition between pairs of trees in primary and secondary forest for all trees together and across the seven tree size classes.

Table S1. Summary of stepwise regression of ant species richness on tree size traits.

Table S2. Distribution of trees among size classes and taxonomic categories in simulated 0.1 ha forest plots.

Table S3. Summary of GLM model of effects of the forest type, tree size and their interaction on ant species richness, nest number and nest site richness.

Appendix S1. Supplementary methods for the rarefaction algorithm for partitioning of difference in species richness.

Appendix S2. Text file providing the R code for the rarefaction algorithm.

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