The biodiversity of arthropods from Australian rainforest canopies: General introduction, methods, sites and ordinal results

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Abstract The arthropod assemblages occurring in the canopies of tropical, subtropical and cool temperate sites have been sampled using a pyrethrum knockdown technique. Details of the techniques used and the climate and vegetation of the areas studied are presented together with an analysis of the distribution of individual arthropods across Orders. An approach using generalized linear modelling partitioned the variance in numbers among sites within forest types and across the three forest types. The effects of both these components were significant. The differences between the ordinal signatures of each forest type are discussed and a number of hypotheses proposed to account for these differences, based on knowledge of the biology of the groups concerned. For the tropical and subtropical sites a comparison was made between samples collected in the low to mid-canopy with ones collected in the high canopy. Numbers of both insects and non-insects collected differed significantly with height in the subtropical forest and the distribution of insects across Orders was also significantly different in this forest type. In the tropical forests numbers of insects differed significantly between the two strata but neither the numbers of non-insects nor the ordinal profiles of either insects or non-insects were shown to be significantly different.

INTRODUCTION

There has been considerable interest in the arthropod fauna that occurs in rainforest canopies recently. Erwin’s (1982) hypotheses concerning the total number of species of tropical forest arthropods, which were based on his studies of Coleoptera in Panamanian canopies, have been followed by further studies in the Neotropics (Adis et al. 1984, Erwin 1983, 1990), Borneo (Stork 1987a, b), Seram (Stork 1988) and Sulawesi (Stork & Brendell 1990, Hammond 1990). Considerable debate has ensued over estimates of the size of the global arthropod fauna (May 1988, Wilson 1988, Stork 1988, Monteith 1990, Gaston 1991, Erwin 1991, Hodkinson & Casson 1991) with participants arguing fiercely over the various multipliers involved in each estimate. The actual database on which these generalizations are based is small, perhaps reflecting the difficulty in generating appropriate field data.

While the actual number of species in canopy faunas remains ambiguous, their speciosity has become apparent with the application of techniques involving insecticidal knockdown of canopy faunas. This method, pioneered by workers such as Roberts (1973) and Southwood (e.g. Southwood et al. 1982), enables large mixed samples of arthropods to be obtained from forest canopies using either fogging devices or a variety of spraying machines, from aerosol cans to orchardists’ backpack dispensers. Samples can then be returned to the laboratory and sorted both taxonomically (usually to ‘morphospecies’) and ecologically (to guilds). The recent interest in the canopy fauna of rainforests stems from the studies of Erwin (1983, 1990), who applied fogging techniques in selected Neotropical forests.

The arthropods from Australian rainforest cano-
pies have received little attention to date, although the knockdown technique has been used in dry Australian sclerophyll woodlands (Majer & Recher 1988) and Tasmanian temperate rainforests (Yen & Lillywhite 1990). Nevertheless, Australia presents to the rainforest ecologist a unique opportunity to relate arthropod numbers and diversity to the gradient of climatic conditions and floral complexity evident between tracts of rainforest from the tropical north of Queensland to the cool, temperate regions of Tasmania.

This paper is the first in a projected series which describes the results of knockdown sampling of arthropods from canopies of cool temperate, tropical and subtropical rainforests. General methods and descriptions of study sites are presented together with results and analyses at the level of the Order. In particular, the role of within and between forest type variation is examined, and upper and lower canopy layers are compared. More detailed results on particular Orders and/or forest types will be presented in subsequent papers.

**METHODS**

Subtropical and tropical sites

In subtropical and tropical forests, 10 × 10 m plots were established in forest and a 'column' of canopy sampled. Sites were selected for study on the basis of canopy structure, tree species composition and accessibility. The need to transport considerable amounts of equipment and to find appropriate, high horizontal branches (see below) precluded randomization of sites within the study areas. However, a spread of locations within each forest type was achieved and the locations and intersite distances of each were recorded.

At each site a rope was attached to a pulley over a high central branch after initial access had been gained by catapulting a lead weight and fishing line over the branch. A cats-cradle of lighter ropes was then erected at head height over the area of the plot and a number of circular, cotton collecting funnels, each 0.5 m², were hung throughout the plot. Generally, 10 funnels were used, with minor variation in this number resulting from individual situations and logistic vicissitudes. A collecting vial containing ethanol was inserted into the apex of each funnel.

A pyrethroid insecticide, Pyrethrins 2EL™, delivered using a Stihl SG-17™ backpack mister, was used in all cases. The insecticide, in concentrated form, comprised natural pyrethrins at 20 g L⁻¹ with piperonyl butoxide at 80 g L⁻¹. This mixture was diluted in water at the rate of 1.25 L to 9 L of water. The mister has a range of 6.2 m in still conditions. Two separate samples were taken at each site; the first, delivered from the ground, focused on the understorey from 2 to 6 m in height, and the second, delivered after the sprayer had been hauled into the canopy using the suspended rope, focused on the high canopy. We refer to these as low and high sprays. The exact height of the high sample reflected the height of the canopy at particular sites, being some 8–10 m below the top of the canopy in most cases. There was no physical overlap between the target areas of the two sprays. The two sprays at each site were separated by at least 24 h, each occurred early in the morning in windless conditions and lasted for approximately 5 min. Arthropods were brushed into the collecting vials in each funnel between 1 and 3 h after each spraying event. In a pilot study in which trays were exposed after spraying for 4 h, we found that 63% of arthropods collected fell from the canopy within 1 h, 78% within 2 h and 90% after 3 h. Vials were collected and sorted to the ordinal level in a field laboratory.

A vegetation transect, 30 × 4 m, was run through all nine sites in the subtropical forest and five of the six in the tropical situation. All plants greater than 1 m in height were identified, and their positions and heights measured. A detailed profile diagram was then constructed. For the tropical sites, a complete survey of all woody plants greater than 1.5 m in height in the 10 × 10 m plot was also made.

All samples were taken in the wet season. Results reported here are from nine sites located in the subtropical rainforest adjacent to Lamington National Park, southeast Queensland, collected in December 1989 and January 1990; and, an additional six sites in the tropical forests around Cape Tribulation, north Queensland, collected in January 1991. Descriptions of the sites are presented below.

Cool temperate sites

Five trees were sampled in the Styx River State Forest, east of Armidale, in February 1990. Nothofagus moorei is by far the most abundant tree species in this forest. Individual trees of this species were...
sampled by suspending 10 collecting funnels within the canopy of each tree. Other details of the spraying events were as for the subtropical and tropical forests.

STUDY SITES AND VEGETATION

Tropical forest

All tropical sites are located close to Cape Tribulation (16°04'S, 145°28'E) in north Queensland. The first two sites sampled are located on the property 'Pilgrim Sands' within 0.5 km of the sea, the remaining four sites are approximately 4 km to the southwest and 1 km from the sea. All are at less than 100 m altitude. All sites are in undisturbed areas, although some clearing for settlement has occurred in areas adjacent to each site.

Figure 1a presents a climatic summary for 'Pilgrim Sands' based on data from 1987 and 1988. Although there may be considerable falls of rain throughout the year, the area has a distinct wet season centred in February and March, with monthly falls of over 600 mm. There is a small annual temperature range with mean maxima ranging from 27°C in July to 34°C in January, and mean minima ranging from 21°C in July to 26°C in January.

The vegetation of the region has been described by Tracey (1982) and species lists of the trees and climbing plants of Cape Tribulation have been published by Jessup and Guymer (1985) and McKenzie (1985). The vegetation of the actual sites sampled has been described and details will be presented in a separate paper. The forest is a complex mesophyll vine forest following the structural-physiognomic classification of Webb (1959) and Webb et al. (1976). Palms were an important element of the vegetation with *Calamus* spp. being abundant on all sites; *Licuala ramseyi*, *Linospadix* spp. and *Normanbya normanbyi* were also present although in lesser numbers. Numbers of tree and shrub species in each 10 X 10 m plot ranged from about 12 to 22. There was no overlap of tree and shrub species with those recorded in subtropical sites and little overlap among sites within the tropical forest. The majority of tree and shrub species occurred in only one or two plots making floristic generalizations difficult.

Subtropical forest

All the subtropical sites are located in the region known as 'Green Mountains' adjacent to O'Reillys' Guesthouse, Lamington National Park, southeast Queensland (28°13'S; 153°07'E). All sites are within 1 km² of forest. The forest has been disturbed by cyclones and minor clearing in patches over the last forty years, and some sites have substantial understorey growth of secondary species. Current use is exclusively for tourism.

Figure 1b presents a climatic summary for a nearby area (redrawn from Kitching 1983). Rainfall is distributed throughout the year with a sum-
mer peak in February and March (ca 500 mm per month) and a winter minimum in August (ca 100 mm per month). Mean maximum temperatures range from 16°C in July to 25°C in January, and mean minima from 8°C in July to 16°C in January.

The vegetation of the region has been described by McDonald and Whiteman (1979) and McDonald and Thomas (1990). The forest is a complex notophyll vine forest following the classification of Webb (1959), and a 'Subtropical Rainforest, Agyrodendron actinophyllum alliance, Suballiance 11 (Caldcluvia-Cryptocarya erythroxylon-Ortites-Melicope octandra-Acmena ingens)' of Floyd (1990). Floyd (1990) gives detailed floristic information on the same vegetation type from the adjacent Border Ranges National Park, in New South Wales. A total of 74 woody species was recorded from the transects taken through the sites. Their canopies were varied but were dominated by species such as Geossois benthamii, Lophostemon conferta, Agyrodendron actinophyllum, Ficus watkinsiana, Baloghia lucida and Pseudowienmannia lachnocarpa. The commonest understorey species were Acradenia euodiiformis, Synoum glandulosum, Dysoxylon rubrum, Wilkeia spp. and Triunia youngiana.

Cool temperate forest

The sites are located in the Styx River State Forest (30°3’S, 152°17’E) of northern New South Wales, 53 km east of the city of Armidale. Their altitude is approximately 1100 m. All sites are in undisturbed areas.

Figure 1c presents a climatic summary for the 1988–91 period based on data collected at the Ebor Trout Hatchery, a few kilometres west of the study site. Rainfall is, overall, somewhat less than at the other sites, with no marked peaks. Average monthly totals vary from 34.8 mm in August to 342.5 mm in April. In general, the late summer months are the wettest. Mean maximum temperatures range from 9.1°C in July to 22.8°C in December, and mean minima from −1.2°C in July to 11.4°C in January.

The vegetation of the region has been described by Floyd (1990). The forest is a microphyll fern forest following the structural-physiognomic classification of Webb et al. (1976). Floyd (1990) described it as 'cool temperate rainforest of the Nothofagus moorei alliance, Nothofagus-Ceratopetalum suballiance'. The canopy is a virtual monoculture of Nothofagus moorei with Ceratopetalum apetalum in gaps and in the understorey. Other common trees include Doryphora sassafras, Calcluvia paniculosa and Quintinia sieberi (Floyd 1990).

Analysis

The original data sets for each site contained 32 categories comprising 22 orders of insects and 10 orders of non-insectan arthropods. Many of these categories occurred very rarely and, accordingly, some were combined for statistical analysis. The resulting 14 categories, based for the most part on combining closely related groups, were as follows: Collembola; Blattoid-Orthopteroids (Orthoptera, Phasmatodea, Blattodea, Isoptera, Embioptera, Plecoptera, Dermaptera, Mantodea — referred to in the figures as Orthopteroids); Psocoptera; other Hemipteroids (Homoptera, Heteroptera, Thyasanoptera); Coleoptera; Diptera; Lepidoptera; Hymenoptera; other Insects (Megaloptera, Neuroptera, Strepsiptera, Mecoptera and Trichoptera); Araneae; Acarina; 'other Arachnids' (Opiliones, Scorpiones, Pseudoscorpiones, Onychophora); Myriapoda (Chilopoda and Diplopoda); and Crustacea (Isopoda and Amphipoda).

Structured multinomial logit models were fitted separately to the insect data and to the non-insect data using the GLIM™ package for generalized linear modelling. The purpose of this analysis was to investigate differences in the frequency distribution of individuals among orders (henceforth, 'ordinal profile'). For these analyses, the response variable was the number of individuals in each order at each sampling site. For comparisons of one forest type with another, this required summing the samples from the high and low canopy in the tropical and subtropical sites. A separate analysis using the same approach was carried out in order to compare the arthropod assemblages from the high canopy with that from the low canopy for the tropical and subtropical sites. Because unequal numbers of funnels were used in the different forest types and in different sites within each forest, the counts were standardized, in each case, as the number of individuals per funnel.

The models that were fitted considered that counts fell on a logarithmic scale (log link function
in GLIM\textsuperscript{TM}) and that errors followed a Poisson distribution (Aitken et al. 1989). Assessment of normal probability plots verified that these assumptions were appropriate. For the first analysis, categories designating the orders, the forest types and the sampling sites were used as the predictor variables. Sampling sites were nested within forest. In order to assess whether the ordinal profile varied across forests and/or across sampling sites within forests, the relevant terms were the interactions between order and forest or order and site, respectively.

As indicated above, for both the tropical and subtropical forests, separate samples were collected in the low and the high canopy at each site. For each of these forest types, statistical analysis of the pattern of occurrence of orders between the two strata were carried out by fitting additional multinomial logit models to the insect and non-insect results. The effects of order, height and the height by order interaction were added sequentially. Again, the number of individuals per order per funnel was used as the response variable. The counts were assessed on a logarithmic scale (log link function in GLIM\textsuperscript{TM}) and errors followed a Poisson distribution. The predictor variables were categories designating the orders and the height of the collecting hoop. Heights were nested in the sampling sites so that pairs were considered jointly. When determining whether the ordinal profile differed between the two height categories, the relevant term was the interaction between order and height.

RESULTS

Figure 2 presents a summary of the results from the tropical and the subtropical sites. Results from the cool temperate forest at the Styx River site are presented in Figure 3.

The tropical samples (Fig. 2a), summed across all six sites, comprised 9967 individual arthropods of which 89% were insects. Overall, there were 18 insect orders and 8 non-insect Orders present. Five Orders made up 88% of the insect total, within which the Diptera (38.6%), Hymenoptera (19.1%) and Coleoptera (16.8%) dominated. The Hemiptera (8.1%) and the Collembola (5.4%) were the other two relatively abundant Orders. Seven Orders were represented by 10 or fewer individuals. Among the non-insects the spiders (55.7%) and the mites (34.3%) comprised a large majority of the total of 1116 individuals. In particular contrast with samples from the other forests, it was noted that the Psocoptera comprised a mere 2.02% of the insects sampled.

The nine subtropical samples (Fig. 2b) comprised 22 984 individual arthropods of which 84.6% were insects. Overall, there were 17 insect Orders and 7 non-insect Orders present. As in the tropical samples, five Orders made up a substantial majority (81.5%) of the individuals sampled. However, the five Orders involved were a different set than in the tropical results. The Collembola (28.7%) were the most abundant followed by the Psocoptera (20.3%) and the Diptera (19.8%). The Coleoptera (12.8%) and the Hymenoptera (7.2%) complete the sum. Five Orders were represented by 10 or fewer individuals. As in the tropical samples, among the non-insects the spiders (43.8%) and the mites (42.5%) comprised a large majority of the total of 3537 individuals.
The arthropods collected from the five trees of *Nothofagus moorei* (Fig. 3) comprised 5360 individuals of which 79% were insects. Overall, there were 14 insect and 4 non-insect Orders present. Four Orders made up 85.3% of the insect total, within which the Collembola (44.9%) were by far the most abundant. The other three relatively abundant Orders were the Coleoptera (22.9%), the Diptera (8.47%) and the Hymenoptera (9.1%). Five Orders were represented by 10 or fewer individuals. Among the non-insects the spiders (23.5%) and the mites (72.8%) comprised a large majority of the total of 1108 individuals. Again, in contrast with the subtropical sites, there were very few Psocoptera (2.42%).

Results from the multinomial logit models used to analyse these ordinal profiles are presented in Tables 1a and b where the changes in scaled deviance associated with each term are indicated. The models were fitted sequentially in the order presented in the tables. Since the number of counts was large, these differences in scaled deviance were approximately Chi-square distributed (Aitken *et al.* 1989). In all cases, the Chi-square values were highly significant (*P* < 0.01). This indicates that Order, forest and site within forest all contribute significantly towards explaining variation in the number of individuals collected. Furthermore, there was significant variation in the ordinal profiles among the forests and among the sites within forests. Other than the first, these are all important results for the present study. That major amounts of variation were due to Order is neither surprising nor very interesting from an ecological point of view.

The tables also indicate the percentage of variation that is explained by each factor. Over half of the variation in the number of insects sampled was attributed to variation in natural abundances associated with the different Orders. This variation was considerably lower for non-insects (19.52%) where, instead, most of the variation was associated with local differences between sites within forests.

In comparing the ordinal signatures from low and high canopy for the tropical sites, a total of 3381 (38.2%) insects from the lower strata and 5470 (61.8%) from the higher strata were recorded. Comparable numbers for non-insects were 585 (52.4%) and 531 (47.6%), respectively. Among the insects 16 Orders were recorded from the lower canopy, 15 from the upper; the difference reflected the detection of a single strepsipteran, ectoparasitic upon an auchenorrynch homopteran, and is not, in our view, significant. Of the major Orders only the Collembola showed markedly higher numbers in the lower canopy samples as compared with the higher canopy ones (8.2% of insects in the lower canopy vs 3.7% in the higher canopy ones). Two major Orders showed the reverse pattern: the Hemiptera (7.4% of insects, lower canopy, *vs* 22.6%, higher canopy) and the Hymenoptera (16.3% of insects, lower canopy, *vs* 21% higher canopy). The proportions of Diptera and Coleoptera in samples of insects from the two strata were more or less the same. For the non-insects the proportion of spiders was comparable across the lower and upper samples but mites were better represented in the lower samples (43.1% of non-insects) than in the higher ones (24.5%).
For the subtropical sites, a total of 10 017 (51.5%) insects was recorded from the lower strata, 9430 (48.5%) from the upper; and, for non-insects, 2076 (58.6%) and 1461 (41.3%), respectively. Sixteen insect orders were recorded from both the lower and upper canopies. As in the case of the tropical samples only the Collembola among the major orders showed markedly higher numbers in the lower canopy samples as compared with the higher canopy ones (41.7% of insects in the lower canopy vs 14.8% of the higher canopy ones). Four major orders showed the reverse pattern: the Hemiptera (3.9% of insects, lower canopy, vs 8.4%, higher canopy), Coleoptera (10.4% of insects, lower canopy, vs 15.3%, higher canopy), Diptera (15.5% of insects, lower canopy, vs 24.3%, higher canopy) and the Hymenoptera (5.1% of insects, lower canopy, vs 9.5%, higher canopy). The proportions of Psocoptera in samples of insects from the two strata were more or less the same. For the non-insects the proportion of spiders was comparable across the lower and upper samples but, in contrast with the tropical samples, mites were better represented in the higher samples (53.6% of non-insects) than in the lower ones (34.6%).

No such analysis of the results from the cool temperate forest was possible as the arthropod samples were not stratified into upper and lower fractions.

In both forests, there was a significant effect of order on the numbers of insects ($P<0.001$ in each case) and non-insects ($P<0.001$ in each case) collected. Height was also important in determining the numbers of insects and non-insects collected in the subtropical forest ($P<0.001$ in each case), but only the number of insects collected in the tropical sites. More interestingly, whether ordinal profiles differed in the high and low canopy was tested by examining the height by order interaction. We found evidence that the ordinal profile of insects differed in the high and low canopy of the subtropical ($\chi^2 = 316.4$, d.f. = 168, $P<0.001$), but not in the tropical forest ($\chi^2 = 37.9$, d.f. = 88, $P>0.9$). In neither forest did the ordinal profile of non-insects vary with height (tropical forest: $\chi^2 = 9.1$, d.f. = 44, $P>0.9$; subtropical forest: $\chi^2 = 65.2$, d.f. = 84, $P>0.9$).

**DISCUSSION**

The statistical analyses show unequivocally that each forest type has a characteristic ordinal profile and that these differ from each other in a highly significant fashion. The reasons for these significant differences are apparent on close examination of the profiles themselves. Particularly noteworthy differences are the gradual decrease in numbers of Collembola from the cool temperate sites northwards. The reverse trend is seen clearly in the levels of abundance of Diptera. Among the non-insects an increasing tropical to temperate trend is seen in the numbers of Acarina, the reverse for the spiders. Lastly, of special note, is the relative prominence of the Psocoptera in the subtropical samples compared with the other two forest types. The biological reasons for these trends cannot be ascertained with confidence at this time. A number of hypotheses are available to explain the differences observed for particular orders. In particular, consider the Collembola, Diptera, Psocoptera and Araneida. Collembola are 'found predominantly in leaf litter and other decomposition habitats' (Greenslade 1991) and their increased abundance in the more southern forest types may well reflect the greater occurrence of aerial leaf litter in the canopy associated with the slower rates of decomposition that must occur in the cooler and drier conditions. In general, tropical forests are supposed to have higher rates of litter decomposition and, in consequence, smaller amounts of 'standing' litter at any time (Anderson & Swift 1983; Takeda et al. 1984). The reverse trend in the Diptera, which for the most part require moist larval habitats (Colless & McAlpine 1991), may well reflect the consistently moister conditions of the tropical forest compared with more southern ones. Similar arguments may be mounted for the temperate to tropical increase in proportions of the Araneida, many of which live in exposed situations which must put them more in danger of desiccation. The Psocoptera are an interesting case, being the Order that seems to mark out the subtropical samples from both the tropical and cool temperate ones. T. New (pers. comm.) observed that Psocoptera are not generally abundant within tropical forests in southeast Asia and that direct heavy rainfall may be inimical, especially to the bark-dwelling forms. The higher rainfall regime in the tropical sites and the more open canopy in the cool temperate forests may well disadvantage Psocoptera in this regard, producing the pattern observed in the results. This remains a hypothesis to be tested.
Site to site differences within a particular forest type, shown by the analysis to account for over 10% of the overall variance in numbers of arthropods, may reflect a number of factors. Intersite distance, site to site differences in vegetation and/or microclimate come to mind. An analysis of these site to site differences within each forest type for species of Coleoptera and Psocoptera, using measures of β-diversity, is in preparation.

We are aware that a different sampling design was employed in the cool temperate forest, involving the sampling of individual trees rather than the mixed blocks of canopy that were sampled in the other two forest types. The reasons for this were two-fold. First, the structure and vegetational composition of the cool temperate forest does not lend itself to the mixed block approach. The more open structure and consequent sub-canopy air movement reduce the efficiency of methods which involve collecting arthropods following a drop of several metres to the collecting trays. Further, the near monocultural nature of the cool temperate canopy reduces the need to incorporate mixed blocks of canopy in a sampling strategy. Second, the method used in the cool temperate forest canopy was also part of a sampling design in use for a different purpose, the results of which will be presented in due course. From the point of view of the GLIM™ analysis this difference in sampling methods, although perhaps unfortunate, was in no way disabling. As long as the proportional representation of the different Orders across the three forest types was well sampled, by whatever method (and the variances involved suggest they were), then the results of the analyses are valid.

Differences reflected in the ordinal profiles between the high and low canopy samples in the tropical and subtropical canopies are significant for the subtropical forest. The upper canopy of the subtropical forest is characterized by a sharp decline in the numbers of Collembola compared with the lower canopy. This difference, given the dominance of Collembola, numerically, in the samples may well be largely responsible for the statistical result. Exactly why this difference might arise is unknown but it could be a result of an association of the Collembola with the tree stems and their epiphytes rather than the leaves themselves. The alternative idea, that the canopy Collembola are merely an upward extension of the fauna of the leaf litter of the forest floor (which diminishes numerically with distance from the ground), is not borne out by species level analyses, which identify several canopy specialists among the Collembola from the samples. Similarly, Coy and Greenslade (in press) showed that the Collembola of tree trunks in Tasmanian cool rainforests are not merely extensions of the ground fauna.

The profiles obtained bear comparison with similar results available in the literature. In particular it is of interest to consider Australian results and those from other Old World rainforests.

Majer and Recher's (1988) results from three species of Western Australian Eucalyptus (marginata, wandoow and accedens) show dramatic differences from the rainforest profiles presented here. The key differences are in the relative abundances of Hemiptera and, in particular, Hymenoptera. Proportions of Hemiptera in the results of these authors vary from 9.4 to 12% for most samples; a figure comparable with our tropical results. However, in one set of samples this proportion reached 71.3% as a result of an outbreak of psyllids in the plot of E. wandoow concerned. Hymenoptera, in general, formed a much higher proportion of the insect assemblages than in any of our rainforest results (21.9–62.8% except for the psyllid outbreak situation when the number of Hymenoptera was only 2.6% of the overall insect fauna — an artifact of the extreme skewness of the counts in this case). This higher proportion of Hymenoptera was due, almost exclusively, to the very large proportions of ants involved (18.4–61% around a mean of 37%, again excluding the psyllid outbreak case). In our samples the highest proportion of the insects ever represented by ants alone was 26% (tropical forest) with much lower figures in the other two forest types. Psocoptera and Collembola were virtually absent from the Eucalyptus results but Blattodea were relatively abundant comprising up to 9.3% of the samples (cf. our highest representation of 1.4% in the tropical samples).

Yen and Lillywhite (1990) sampled eight trees of Nothofagus cunninghamii and one of Nothofagus gunnii in Tasmania in February 1990. Their results form a useful comparison with our Styx River results. The key differences among the insects between our results and those of Yen and Lillywhite is in Collembola and Psocoptera. Our samples had many more Collembola (44.9%) compared with
the Tasmanian data (3.4% for *N. cunninghamii*, 19.9% for *N. gunnii*) and many fewer Psocoptera (2.4% compared with 24% for *N. cunninghamii* and 27.8% for *N. gunnii*). In other respects our insect results were not greatly dissimilar to those for *N. cunninghamii*. The *N. gunnii* comparison is less useful, being based on only a single Tasmanian sample. Whether the different profiles represent a seasonal and/or locational difference cannot be differentiated at this stage. A January sample taken in another northern New South Wales site (the results of which are not presented here) showed reduced Collembola proportions (2.2%) but no corresponding increase in Psocoptera proportions (3.8%), suggesting that although there obviously are important seasonal changes taking place this is unlikely to account for the differences observed between our insect results and those of Yen and Lillywhite (1990). For non-insects, our results and the Tasmanian ones for *N. cunninghamii* are remarkably similar (Araneida: our results 23.8%, Tasmanian results 23.8%; Acarina: our results 72.8%, Tasmanian results 73.5%).

Basset's (1990) results from individuals of the tree *Argyrodendron actinophyllum*, from subtropical rainforest near Brisbane, on first sight seem to provide a better comparison, at least with our subtropical results. However, he was using restricted canopy fogging (RCF), a method which involves fumigating enclosed branches with carbon dioxide, rather than pyrethrum knockdown. Further, he did not count Collembola or Acari, two of the most abundant groups in our subtropical samples. He obtained high proportions (41.9%) of usually sessile psyllids (Homoptera), which are almost absent from our samples. This may well be a result of the difference between his sampling procedure and ours. However, even excluding his psyllid counts and recalculating proportions for our samples after removing the Collembola counts, his proportions of other orders still differ markedly from ours. In terms of the proportions represented by each order, we encountered many more Diptera than he did (27.6% compared with 2.7%), more Psocoptera (28.4% compared with 14.5%) and fewer Coleoptera (17.9% compared with 26.8%). We are confident that these differences are also methodological. The manoeuvring of plastic sheeting involved before RCF would undoubtedly disturb many of the highly mobile Diptera; the method does not sample tree trunks, habitat for most of the Psocoptera; and the higher proportion of Coleoptera is adequately accounted for given the undersampling of these other two major groups.

Lastly, the results of Stork and Brendell (1990) for tropical canopies at a variety of altitudes in northern Sulawesi provide a useful comparison, especially for our tropical results. These authors used pyrethrum knockdown methods as we did, although they used a fogging device in contrast to our mister. Their results for the Diptera are most instructive. The range of proportions for this order in their samples ranged from 43% to 70.3% with the proportion diminishing with altitude. Their lowland results show an even larger proportion of Diptera (70.3%) than our tropical samples (39.8%) and it seems that the diminution with increasing altitude at the one latitude in Sulawesi is recapitulated with increasing latitude in our results. Our tropical results also show a consistently higher proportion of Coleoptera than do Stork and Brendell (1990; 15.2% compared with their highest figure of 9.1%), but again this may be an artifact resulting from the higher proportions of Diptera in their samples which necessarily must diminish the proportions of other orders.

Many further questions are engendered by our results. Two in particular will bear investigation. First, all the results presented here were collected during the summer wet seasons in each forest type. Further, in all three cases, the seasons concerned were actually wet in contrast to the most recent (1991–92) season, for example, in which widespread drought produced a substantial impact on the vegetation of the tropical and subtropical canopies at our study sites. Further, we have no knowledge of the way in which these profiles might differ between seasons even in a 'typical' year. In addition, we cannot assume that the results from each forest type presented here are representative of the type in each case. We know from other results (Yen & Lillywhite 1991, Kitching unpublished) that ordinal profiles can differ substantially from one patch of *Nothofagus* forest to another, even over short distances. We have no information on this aspect for tropical or subtropical forests.

Study of the patterns of biodiversity and the mechanisms that underlie them at the ordinal, familial and specific levels across Australian forest canopies remains in its infancy and yet such results
are potentially of significance for making informed conservation decisions on a variety of issues. The absence of invertebrate diversity data, particularly from Australian tropical ecosystems, both in general (Haynes et al. 1991) and with respect to particular conservation debates (Stewart et al. 1991), has been commented upon recently. Our present contribution is to be regarded as a first necessary step in redressing these omissions.

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