

RESEARCH ARTICLE

Growth variations of edaphic specialist species in a reciprocal pot experiment in Sri Lanka

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Abstract: Edaphic factors may determine the habitat associations shown by some tropical trees. In Sinharaja, a tropical rainforest in Sri Lanka, about 79% of tree species show habitat preferences. The present study attempted to investigate the role of edaphic factors in determining habitat specialization, using a reciprocal pot experiment with four edaphic specialist species (*Agrostistachys hookeri*, *Semecarpus gardneri*, *Mesua nagassarium* and *Agrostistachys intramarginalis*). Both *Agrostistachys* species showed increased growth rates and total biomass when grown in their native soils compared to that of non-native soils, while *M. nagassarium* showed a better growth in non-native soils. Higher biomass allocation to roots was evident in all species when they were grown in sand-rich valley soil irrespective of their specialized soil type. No significant trend was observed in foliar nutrient concentrations when grown in native or non-native soils. The results suggest that edaphic factors play a vital role in the spatial distribution of *Agrostistachys* species. This is perhaps due to plants showing more acclimatory shifts when grown in their native soils compared to non-native soils. The results also suggest the importance of other factors such as plasticity, water-use efficiency and herbivory in the spatial distribution of plants.

Keywords: *Agrostistachys hookerii*, *Agrostistachys intramarginalis*, edaphic factors, habitat specialization, *Mesua nagassarium*, Sri Lanka.

INTRODUCTION

Early workers assumed that the high diversity associated with tropical forests is always accompanied by an even spatial distribution of species. However, later studies report that tropical tree species always show a

certain degree of clumping rather than even distribution across the landscape¹. Many factors contribute to the maintenance of species diversity in tropical rainforests²⁻⁴. Niche differentiation with respect to resource availability remains a prominent hypothesis to account for the maintenance of tree species diversity in tropical forests⁵. One way of expressing resource-based niche differentiation is habitat specialization; different tree species are best suited to different habitats, where they are competitively dominant and relatively more abundant⁶. The importance of edaphic variables determining habitat associations of tree species has been stressed in previous studies⁷. To establish the relative contribution of habitat specialization to the maintenance of diversity in tropical forests, a rigorous quantification of vegetation and environmental parameters is required⁸.

It has been proposed that there may be sufficient heterogeneity in light and other physical factors in and around canopy gaps to explain the coexistence of tree species⁹. This is an example of the 'regeneration niche' hypothesis. In many plant communities, habitat partitioning is possible mostly at the seedling level, where habitats are more heterogeneous than those experienced by an adult plant⁹. There is substantial evidence to favour the habitat partitioning theory though its underlying mechanisms and processes are not yet identified precisely. Scientists have also focused on alternative hypotheses, many of which invoke the role of natural enemies in distance and density dependent mortality of juveniles and the disproportionate mortality of more abundant species.

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In contrast to the early thinking that tropical forest trees are evenly distributed throughout the landscape, later studies suggest that the distribution of trees may be influenced strongly by edaphic heterogeneity⁹⁻¹². A study⁶ found that 64% of species in a 50-ha plot on Barro Colorado Island, Panama show significant habitat associations driven by edaphic factors. A comparative study carried out on three diverse neotropical forests (in Colombia, Ecuador and Panama) also concluded that belowground resource availability plays an important role in the assembly of 36–51% of species in tropical tree communities at a local scale¹³. Different approaches (mainly plot-based studies) have been undertaken by researchers to find out whether species distributions are significantly correlated with edaphic factors^{6,14,15}. In most instances, where spatial variation in the abundance of species of trees has been studied, associations have been found with different habitat variables and their interactions with the biotic environment^{1,16}. Studies have shown that species distribution was strongly correlated with variations in topography and other edaphic factors such as soil moisture and nutrient status^{9, 12, 13, 17-19}. In support of these findings, a study²⁰ showed that 60% of the tree species in a 52-hectare plot at Lambir, Malaysia, have spatial distributions biased across the edaphic and topographic gradient. Some of the most plausible edaphic factors driving this non-random distribution of trees in the tropics may include soil drainage, nutrient availability and soil texture²¹⁻²⁴. A study carried out in the Peruvian Amazon²⁵ has shown that herbivore pressure also could promote habitat specialization. However, in spite of all these studies, there are many unanswered questions on the role of habitat heterogeneity in maintaining high tree species richness in tropical rain forest ecosystems. To address some of these controversies, rigorous testing of habitat-related spatial aggregation in tropical forest communities is required, in order to provide experimental verification that tree species perform better in habitats in which they are aggregated. A previous study²⁰ also stressed the importance of gathering evidence to determine the performance of plants when they are exposed to native soils compared to non-native soils.

The 25-ha Forest Dynamic Plot (FDP) in Sinharaja is located within the lowland rainforest zone and has an elevation range from 424 to 575 m.a.s.l. The structure and floristic composition of the Sinharaja FDP show an interesting correlation across a small scale altitudinal gradient of 151 m²⁶. Based on the spatial heterogeneity of three physical parameters; viz. elevation, slope, and convexity, eight different topographic habitats were identified within the plot. Of the 125 tree species tested, 99 species showed at least one positive or negative association (specialists) to one or more of the habitats identified²⁶. Other tree species showed no special habitat

preferences (generalists) and were distributed more or less evenly throughout the landscape. The Sinharaja FDP provides an appropriate site to investigate soil-related habitat specialization exhibited by some of these tropical tree species along this small scale elevational gradient. The present study attempted to examine whether soil-related factors may have contributed to species-habitat association, by carrying out a pot experiment with two valley-dominant (*Agrostistachys hookerii* and *Semecarpus gardneri*) and two ridge-dominant (*Mesua nagassarium* and *Agrostistachys intramarginalis*) species. Seedlings of all four species were grown in their native soil as well as in non-native soils to test whether species have soil-related performance differences in accord with their spatial distributions in the natural forest. We hypothesize that seedlings of the four forest species will grow faster in their native soils than in non-native soils.

METHODS AND MATERIALS

Site description: The pot experiment was conducted in a specially constructed plant house at the Research Station in the Sinharaja World Heritage Site (6° 21' – 26°N, 80° 21-34'E; hereafter known as Sinharaja) located in the ever-wet south west lowlands of Sri Lanka. On a regional scale Sinharaja represents a mixed dipterocarp forest²⁷ and locally it is classified as a *Mesua-Doona* community. The mean annual rainfall at Sinharaja is 5016 mm and it varies from 4080 to 5907 mm²⁸. The annual minimum and maximum temperatures are 22 and 28 °C, respectively. The highest monthly temperatures are observed in April (25.3 °C), May (25.9 °C) and

Table 1: Variation in some soil chemical and physical characteristics (at 0-20 cm depth) of the ridge and valley at the Sinharaja FDP in Sri Lanka.

Soil parameter	Mean elevation	
	Ridge (>525 m)	Valley (<470 m)
Total C (%)	3.24 (0.22)a	2.48 (0.14)b
pH	4.42 (0.13)a	4.54 (0.11)a
Extractable P (µg/g soil)	1.94 (0.13)b	2.79 (0.22)a
Total N (%)	1.61 (0.1)b	2.13 (0.2)a
Clay (%)	21.0 (5.99)a	11.4 (0.79)b
C:N ratio	22.1 (2.24)a	12.8 (1.59)b
Water potential (kPa)*		
Wet season	-30 (11.7)	-96(81.2)
Dry season	-525 (253.4)	-118(80.8)

*Soil water potentials were measured using the Filter Paper method. Standard error of the mean (SEM) is given within parentheses. Different letters indicate significant differences ($p = 0.05$) between the two elevation levels.

January (25.1 °C) and it is least in December (23.8 °C)²⁸. Sinharaja also experiences a high annual temperature which is typical of the tropics with little seasonal variation and marked daily changes. The detailed information on floristic richness and diversity within the FDP has been given in Gunatilleke *et al.*²⁹.

Soil nutrient status: Some selected chemical and physical factors of soil on the FDP at Sinharaja are given in Table 1³⁰. The 'ridge' soils contain significantly higher clay (21%) and soil organic carbon (C) concentrations (3.24%) than the 'valley' soils. In contrast to this trend, total nitrogen (N) and extractable phosphorus (P) concentrations were significantly higher in the valley soils compared to ridge soils. Soil pH did not show any significant difference between ridge and valley soils. A separate soil study that has been conducted in Sinharaja has shown that valley soils have significantly higher pH and concentrations of total N and magnesium (Mg) than ridge soils³¹.

Test species: *Agrostistachys intramarginalis* (Euphorbiaceae) is a large shrub or treelet growing up to 7 m in height. It is an endemic to Sri Lanka and found in the understory tree layer in the Sinharaja forest. *A. hookeri* (Euphorbiaceae) is an understory shrub and also an endemic to Sri Lanka. *Mesua nagassarium* (Clusiaceae) is a native canopy tree. *Semecarpus gardneri* belongs to the family Anacardiaceae and is an endemic to Sri Lanka. It is commonly found in the sub-canopy layer of the Sinharaja forest.

Experimental design: In the cross-over pot experiment, the plants were subjected to two treatments, i. e. grown in their native as well as in their non-native soils. Seedlings in the two-leaved growth stage were collected from each species from their naturally growing habitats in the forest. For the pot experiment soil was collected from three randomly selected sites from the ridge and from the valley sites within the 25ha FDP. At each site, top soil was collected to a depth of about 15–20 cm and sub soil was collected from 20 – 35 cm depth separately. To prepare the potting medium, first, gravel particles were manually removed and then the soil from the different depths and from the ridge and the valley sites were separately mixed with sand in a ratio of 1:1. As the soils are very clayey in nature, sand was added in order to improve the texture. These soil-sand mixtures were then filled into black polythene bags (15.2 cm x 30.5 cm, 300 gauge); each poly bag contained approximately 3 kg of soil. Twenty five seedlings as replicates from each test species were then potted either in their native or non-native soil-containing poly bags. In total, 200 seedlings were planted (4 species x 2 soil types x 25 replicates). Transplanted seedlings were allowed to establish for

about a week, dead seedlings were replaced and poly bags were arranged in a completely randomized design inside the shade house (covered with coir mats) at the Sinharaja field station. The seedlings received sun flecks that penetrate through the coir mat which mimics somewhat comparable conditions on the forest floor. The experiment was set up in January 2004 and maintained for about 18 months. Poly bags were randomly rearranged within the shade house four times during the experimental period to avoid any growth variations due to edge effects.

Data collection and analysis: Ten seedlings were randomly selected to determine fresh and dry mass at the time of planting. Foliar nutrient concentrations and mycorrhizal colonization were determined using 10 randomly selected seedlings at the start (to determine the initial foliar nutrient concentrations and mycorrhizal colonization) and after 18 months of growth under experimental conditions. Tissue nutrient concentrations were determined on an oven-dried sub sample of seedlings from each treatment. Tissue samples were wet digested and analyzed for N and P concentrations using Kjeldhal and molybdate-blue methods, respectively³². Mycorrhizal colonization was determined using the Phillips and Hayman method³³.

After 18 months of growth (July 2005), harvested seedlings were separated into above- and below-ground components, dried at 70 °C to a constant weight, and weighed to a precision of 1 mg. Based on biomass measurements, root mass ratio (RMR) (root biomass as a fraction of the total biomass) were calculated³⁴. The relative growth rate (RGR) was determined using the following equation. Average initial seedling dry mass was used to compute the relative growth rates at the final harvest (after 18 months).

$$\text{RGR} = (\text{Log}_e m_2 - \text{Log}_e m_1) / (t_2 - t_1)$$

where m_1 and m_2 represent the dry masses (g) at planting date (t_1) and harvested date (t_2), respectively. The data were analyzed using an un-paired t-test for each species separately to determine how RGR differed between seedlings grown in native and non-native soils. The analyses were conducted using MINITAB Release 14.30 statistical software.

RESULTS

Survival of seedlings

The mycorrhizal colonization tests revealed that seedlings were not colonized by mycorrhizae either at

Table 2: Percentage seedling survival rates after 18 months of growth of four tree species in Sinharaja, based on 25 seedlings per species per treatment.

Species	Survival (%)		
	Native soil	Non-native soil	Difference
<i>A. hookerii</i> (Vd)	67 (valley)	61 (Ridge)	6
<i>S. gardneri</i> (Vd)	16 (valley)	11 (Ridge)	5
<i>A. intramarginalis</i> (Rd)	84 (ridge)	83 (Valley)	1
<i>M. nagassarium</i> (Rd)	72 (ridge)	89 (Valley)	17

Vd = valley dominant ; Rd = ridge dominant.

the start of the experiment or after 18 months of growth under experimental conditions. Varying survival rates between test plants do not provide any convincing differences to demonstrate that soil specificity does contribute to survival of seedlings (Table 2). Both ridge dominants species (*A. intramarginalis* and *M. nagassarium*) showed higher survival rates (> 72%) than the valley dominants, *A. hookeri* (61–67%) and *S. gardneri* (11–16%). *S. gardneri* had the lowest survival rates in both native (16%) and non-native (11%) soil types; hence it was excluded from further analyses. However, *M. nagassarium* showed fairly notable chance of survival in the non-native soil compared to native soil (17% difference).

Relative growth rate

A. hookeri showed significantly higher (19%) relative growth rate (RGR) when grown in native soils compared to non-native (Figure 2). *A. intramarginalis* also showed 25% higher growth in its native soil though the difference was not significant at $p = 0.05$. However, *M. nagassarium* has shown a significantly better growth (33%) in non-native soils than in native soil.

Total biomass

The total biomass of *A. hookeri* (valley dominant: Vd) at harvest was 19% higher in valley soils than that in ridge

soils, though the difference was not significant ($p = 0.122$). Similarly, *A. intramarginalis* (ridge dominate, Rd) showed a significantly higher total biomass in seedlings grown in ridge soil than that in valley soil ($p = 0.044$). Unlike the two *Agrostistachys* species, *Mesua*, a ridge species (Rd), gained a significantly higher biomass when it was grown in its non-native valley soil ($p = 0.036$) (Figure 3).

Biomass allocation pattern

In all three species, the root weight ratio (RWR) was higher in the seedlings grown in valley soil than those in the ridge soil and this difference was significantly greater in both *A. hookeri* and *M. nagassarium* (Figure 4; p values are 0.035 and 0.0003, respectively).

Foliar nutrient concentrations

The initial plant nitrogen (N) and phosphorus (P) concentrations of *A. hookeri*, *A. intramarginalis* and *M. nagassarium* varied significantly between species. The plant N and P concentrations were significantly higher in *A. hookeri* (Vd) than in the two ridge dominants, *A. intramarginalis* and *M. nagassarium* (Table 3). However, the shoot and root N and P concentrations in all three species measured at the time of the final harvest (after 18 months of growth) did not vary significantly irrespective of whether they had grown in their native or non-native soil types (Figure 5).

In both *A. hookeri* and *M. nagassarium*, shoot and root nutrient concentrations decreased significantly during the course of their growth, while *A. intramarginalis* showed a slight increase in foliar nutrient concentrations after 18 months of growth, irrespective of the soil medium.

DISCUSSION

The survival variations shown by the seedlings in the present study are possibly due to their varying abilities to adjust to new surroundings under shade house conditions. The poor growth and high mortality rates of

Table 3: Initial shoot and root nutrient contents of *A. hookeri*, *A. intramarginalis* and *M. nagassarium*. One-way ANOVA: different letters indicate significant differences between species at 5% probability level.

Species	Plant part	<i>A. hookeri</i> (Vd)	<i>A. intramarginalis</i> (Rd)	<i>M. nagassarium</i> (Rd)
N (%)	shoot	1.76 ^a	0.71 ^c	1.06 ^b
	root	1.38 ^a	0.67 ^b	0.53 ^b
P (%)	shoot	0.47 ^a	0.07 ^b	0.12 ^b
	root	0.36 ^a	0.05 ^b	0.16 ^c

Semecarpus may be attributed to its inability to adapt to new environmental conditions inside the shade house. Studies have shown that survival and growth of rainforest seedlings rely on plasticity (ability to adjust to varying environmental conditions) and mycorrhizal colonization during their early establishment phase under both natural and simulated conditions^{35, 36}. However, in the present experiment none of the species showed noteworthy home-soil advantage in their survival rates when grown in native and non-native soils. Observations also revealed that none of the test seedlings were infected with mycorrhizae during the experimental period. Therefore, the present results did not give any indication to suggest that the survival rates associate with their choice of habitat or else with mycorrhizal colonization. Previous studies^{20,37} also have shown no unequivocal relationships between soil specificity and survival (or mortality). However, in the present study, *M. nagassarium*, which is a ridge dominant, survived markedly better in valley soils than in the ridge soils indicating its ability to take the advantage of better quality soils (nutrient-rich valley soils) than other test species. The high plastic ability of *M. nagassarium* when exposed to shade and drought conditions has been demonstrated in an experiment conducted under shade house conditions³⁸.

The two *Agrostistachys* species with differing soil-dominance grew faster in their native soils than in the non-native soils. This trend was evident in terms of their RGR and total biomass indicating that for *Agrostistachys*

species edaphic factors do play a vital role in habitat preferences. A reciprocal experiment conducted in Sepilok Forest Reserve, Borneo⁷ showed that edaphic specialist species, when grown on their native soils, are capable of acclimatory shifts to achieve resource-use efficiency than species not specialized to that soil type. In the present study, *A. intramarginalis* performed better in its less fertile native ridge soils than fertile valley soils, indicating its high plastic ability (morphologically and/or physiologically) in less fertile soils to compensate for their overall growth. According to one study³⁹, plasticity is a morphological (and physiological) trait shown by some plants in response to edaphic variations that could be driven by scarcity of nutrients and/or water. The results suggest that being plastic, *A. intramarginalis* has higher chances of acclimatory shifts in order to accomplish resource-use efficiency under low-fertile soil conditions.

M. nagassarium on the other hand showed no growth preference in their native soils; in fact growing significantly well under non-native soils. Two studies^{37,40} have also observed that plants do not always grow faster and better on their native soils than in non-native soils. Another study⁴⁰ has suggested that the differing responses of plants to different soil types is mainly due to their inherent growth rates rather than responding

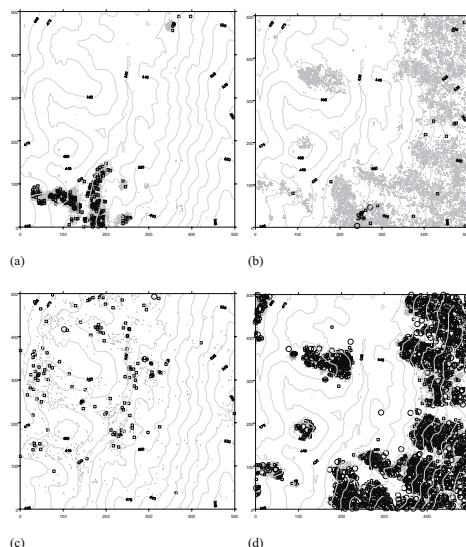


Figure 1: Spatial distribution maps of the four species selected for the study. (a) *A. hookeri* (valley dominant, Vd), (b) *A. intramarginalis* (ridge dominant, Rd), (c) *S. gardneri* (Vd) and (d) *M. nagassarium* (Rd).

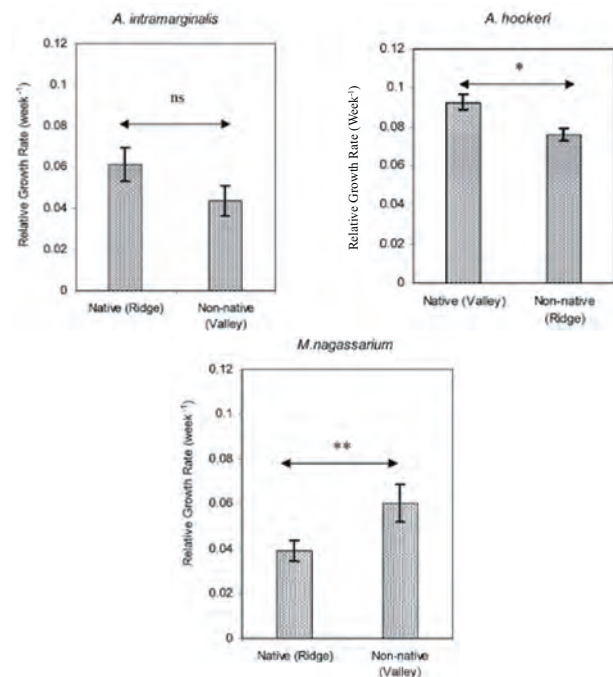


Figure 2: Relative Growth Rates (RGR) of *A. intramarginalis*, *A. hookeri* and *M. nagassarium* in their native and non-native soils. Untransformed data were used to plot the graph. Vertical bars indicate the standard errors of the mean (SEM).

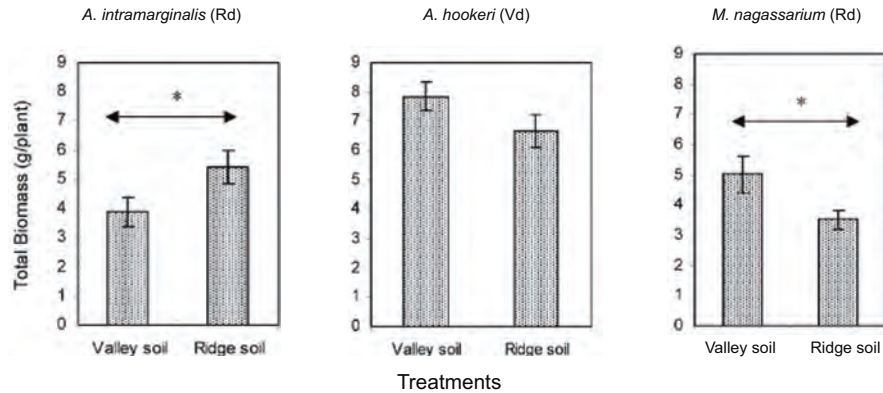


Figure 3: Total biomass of ridge-dominant and valley-dominant species on each soil type. Vertical bars indicate the standard error of the mean (SEM). Asterisks show significant differences between the two soil types at 5% probability level.

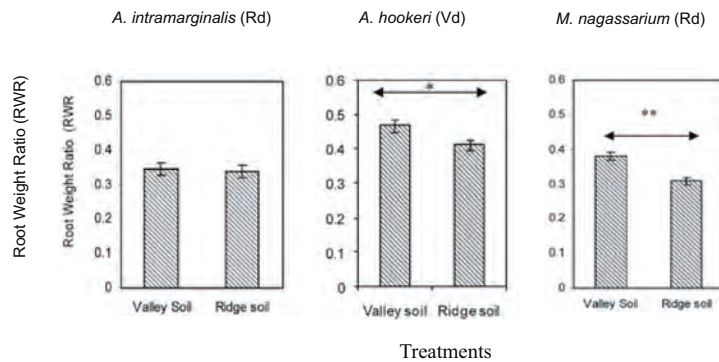


Figure 4: Root Weight Ratio (RWR) of ridge dominant (Rd) and valley dominant (Vd) species on each soil type. Vertical bars indicate the standard error of the mean (SEM).

to changes in soil quality. Therefore, our results also imply that not all plants always perform better in their home-soils than on non-home soils. These ambiguous results indicate that it is not only the edaphic factors that will determine plant distribution, but also other factors such as competition, herbivory, and climatic factors. A study³⁷ has also concluded that there may be potential mechanisms influencing the spatial distribution pattern of plant species other than soil nutrient availability. Some of these potential mechanisms are dispersal limitations and light, which are important unmeasured sources of variations and beyond the scope of this study. A previous study⁴¹ also stresses the importance of combination of factors when determining the spatial distribution of plants.

Seedlings of *A. hookeri* and *M. nagassarium* have shown greater biomass allocation to roots when grown in the nutrient-rich valley soils, irrespective of their

home-soil preferences. Their enhanced root allocations correlate with their better performances irrespective of their native habitats indicating increased nutrient assimilation when roots come across nutrient-rich soils. Evidence also suggests that plant roots tend to proliferate in contact with high-nutrient soils thereby increasing assimilation rates⁴²⁻⁴⁴. Nutrient availability perhaps is not the only reason for *A. hookeri* and *M. nagassarium* to produce more roots in valley soils than in ridge soils. The sandy nature of the valley soil allows easy penetration of the roots deep into the soil and easy spreading compared to clay-rich soil. The anoxic conditions created in the rooting zone due to the greater water holding capacity in the clay-rich ridge soil may be another reason for relatively lower allocation of biomass to the root system than to the shoot in ridge soil.

The higher initial concentrations of foliar N and P in the valley dominant *A. hookeri* are consistent with

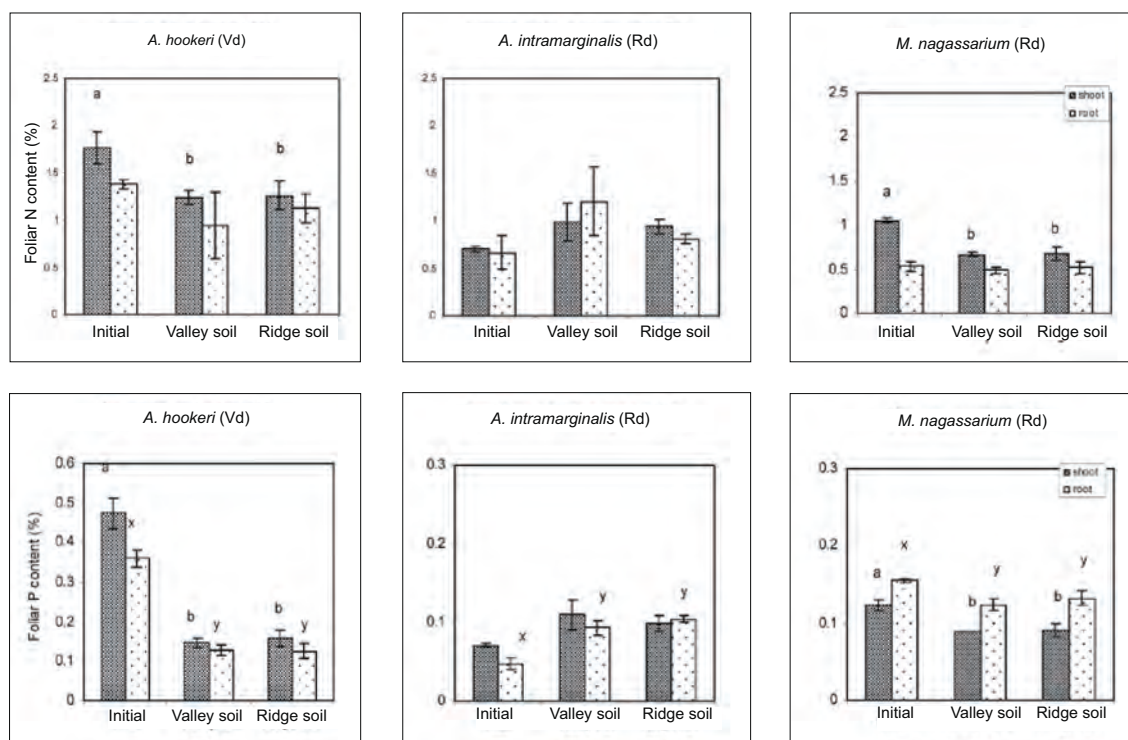


Figure 5: Foliar N and P concentrations in roots and shoots of *A. hookeri*, *A. intramarginalis* and *M. nagassarium* (initial and after 18 months of growth). Different letters indicate significant differences at 5% probability level. Vertical bars indicate standard error of the mean (SEM).

high N and P concentrations in the valley soils. However, the foliar nutrient concentrations did not directly reflect the soil nutrient status always, as factors other than soil nutrient availability would also determine the foliar nutrient levels in plants. Concentrations of N and P did not differ between plants grown in valley and ridge soils for any of the three species after 18 months of growth in their respective native and non-native soils. Therefore, present observations did not give us an opportunity to relate the performance differences in plants to their tissue nutrient concentrations directly. One study²⁰ suggested that differential growth responses of seedlings of some rainforest species were not related to soil nutrient availability and that the reduced growth of seedlings in their non-native soil may have resulted from occasional anoxic conditions in the rooting zone due to clay-rich soils, the absence of native mycorrhizal flora, and the toxic effects due to high Aluminium (Al) or Iron (Fe) concentrations, etc.

Out of the four species studied, two *Agrostistachys* species with different soil preferences, performed better when grown in their respective native soil types indicating that edaphic factors seem to have influenced the growth of plants and consequently the habitat

associations. *M. nagassarium* did not follow this trend and in fact performed even better when grown in non-native soils. From the present results, we could suggest that the performance variations of plants are not only determined by edaphic resources, but also by their inherent eco-physiological traits such as plasticity and other environmental factors.

A previous study⁴⁵ suggest that the spatial distribution of 'habitat specialists' are governed by environmental factors while the distribution of 'habitat generalists' are governed by dispersal processes, though these facts cannot be fully supported or disputed by the findings of the present study. However, it would be imperative to carry out long-term research in the natural environment taking into consideration other parameters such as competition, light, moisture content, understorey and forest gap environments in order to set more light on the underlying factors that eventually decide the spatial distribution of plant species with and without habitat preferences (habitat 'specialists' and 'generalists' respectively). The seedling growth should also be evaluated until at least the seedlings become saplings, as the trees give more information on their preferential traits at the mature stage than at the seedling stage.

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