supporting biodiversity in plantation forest management as do other countries such as Britain. There, private landowners have grants available to them that allow up to 20% of their land to be left unplanted and managed as open ground for biodiversity values (Hodge et al. unpublished report).

Despite the fact that New Zealand plantation forests are not managed specifically for biodiversity they can support a significant number of species in their understoreys. The importance of this biodiversity to crop production is not clear; however, it likely does contribute to an intangible level of environmental quality not present in densely-stocked plantation forests found in the other countries. Perhaps more importantly as far as the conservation of New Zealand’s indigenous biodiversity is concerned, most plantation managers are committed, through the Forest Accord and the Principles, to the conservation of indigenous biodiversity in natural reserve areas and to the conservation of threatened species known to occur in their plantations.

References
NIZFOA, 1997. NZ Forestry Facts and Figures 1997. New Zealand Forestry Commission to the conservation strategy, and operational exotic forestry, are emphasised our concern about the potential loss of some of our indigenous biodiversity, and an acceptance of our international obligations under the 1992 Biodiversity Convention. We should be particularly concerned about the loss of the flora, for its own sake and because plant communities largely constitute the ‘habitats’ in which other components of biodiversity live. In a very short span of evolutionary time the flora has been subject to enormous changes wrought by forest clearance, increased fire frequency, wetland drainage, urban and rural subdivision, introduced mammals, birds and insects, and an enormous influx of exotic plants. There is no clear end in sight for most of these effects, or for the new class of problems arising from global climate change. The latter is just one of several threats to biodiversity originating outside New Zealand, driven by a global economy on which we seem to have little influence.

Forests, both native and exotic, play a very significant role in maintaining indigenous biodiversity. Lowland forest ecosystems, now much depleted in extent, contain many of our more

Biogeography and Forest Biodiversity in New Zealand
John Ogden*

Abstract
The concept of biodiversity is scale dependent. Biogeography is concerned mainly with explaining the larger scale patterns - the origins of the New Zealand flora and the regional patterns of biodiversity within the country (gamma-diversity). Geological and climatic history are important at these scales. Altitudinal gradients illustrate variation at the landscape level (beta-diversity). The conservation of landscape processes is important for maintaining high biodiversity and ecosystem ‘services’. Ecologists cannot yet explain differences in diversity at the community scale (alpha-diversity), but arguably the disturbance regime is a key factor. A case study, comparing kauri (Agathis australis) and beech (Nothofagus solandri) forests, is outlined. If the indigenous conservation strategy, and operational exotic forestry, are to emphasise the sustainability of indigenous biodiversity, then the spatial and temporal scale effects which determine it must be explicitly addressed.

*Centre for Conservation Biology and School of Environmental and Marine Sciences, University of Auckland.

Introduction
The development of a New Zealand Biodiversity Strategy by the NZIF indicates our concern about the potential loss of some of our indigenous biodiversity, and an acceptance of our international obligations under the 1992 Biodiversity Convention. We should be particularly concerned about the loss of the flora, for its own sake and because plant communities largely constitute the ‘habitats’ in which other components of biodiversity live. In a very short span of evolutionary time the flora has been subject to enormous changes wrought by forest clearance, increased fire frequency, wetland drainage, urban and rural subdivision, introduced mammals, birds and insects, and an enormous influx of exotic plants. There is no clear end in sight for most of these effects, or for the new class of problems arising from global climate change. The latter is just one of several threats to biodiversity originating outside New Zealand, driven by a global economy on which we seem to have little influence.

Forests, both native and exotic, play a very significant role in maintaining indigenous biodiversity. Lowland forest ecosystems, now much depleted in extent, contain many of our more

Understorey biodiversity.

8 N.Z. FORESTRY NOVEMBER 1997
unusual plants and animals. Such systems have survived in varying extent and composition throughout the climatic oscillations of the Pleistocene. More recently, forest covered the majority of the landscape. The significance of this is that tree diversity general reflects total biotic diversity (e.g. Richardson and Lum 1980).

Hal Mooney, Secretary General of the International Council of Scientific Unions (ICSU), has recently emphasised that ecologists do not yet know how many species must be saved to keep an ecosystem functioning – providing the ‘services’ we take for granted. Such services include clean water from mountain catchments, regenerating fisheries and soul-sustaining landscapes (Mooney 1997; quoted by Basu 1997). Large-scale changes in such services can have global effects and unpredictable interactions. We do know that some species are more important than others (‘keystone’ and ‘umbrella’ species) and that different trophic levels are interdependent. Theoretically we can argue that high biodiversity ensures ecosystem stability, but the argument is bedevilled by definitions, semantics and scale effects. The challenge for ecology is to develop a new approach linking biodiversity and ecosystem functioning.

This paper starts by presenting the traditional concepts used in the discussion of plant biodiversity. Traditional views about the origins of our flora are currently being questioned. This debate is important because understanding the origins and wider relations of our plant communities underpins the scientific justification for their conservation. The well-known geographical patterns of biodiversity within New Zealand are outlined, and some comments made on the implications of this for our reserve network. Reserve priorities should be based not simply on species inventories, but on the quantitative understanding of change along major environmental gradients, and an appreciation of the dynamic nature of plant communities as mosaics on the landscape. To preserve biodiversity we must preserve the landscape processes which have selected for or against particular combinations of demographic characteristics in different species, thus determining their ability to grow together in natural communities.

Scales of plant species diversity - a conceptual framework

Scale is central to the perception of diversity. While the general trend of decreasing diversity from the equator to the poles is clear at one scale, at finer scales of resolution more detailed patterns may be superimposed. Whittaker (1972) presented a framework for the discussion of species diversity which, with minor modifications (e.g. Soulé 1986), has been widely adopted.

Alpha-diversity relates to the number of species coexisting within a uniform habitat. It is an inventory of the species present within a community, often referred to as ‘species richness’. In common with other authors (e.g. Huston 1979; Peet 1974; Wilson and Keddy 1988; Wilson and Sykes 1988), I recognise that equitability (the relative proportions of species) is a component of diversity, but species richness is a simpler statistic. Beta-diversity describes the between community level of diversity. It provides an index of the diversity across a landscape within one biogeographical region. For example, as the community composition changes along an altitudinal gradient new species are encountered and others drop out, and this species turnover rate is termed beta-diversity. Two stands with the same number of species present (same alpha-diversity) can differ in composition, and thus together they exhibit beta-diversity. Beta-diversity can be measured by a similarity coefficient (e.g. Sorensen’s coefficient) or using ‘between stand distance’ in ordination space (Druitt et al. 1990). Gamma-diversity can be regarded as the total species inventory for a large geographical region, e.g. regions of New Zealand or the country as a whole. Thus gamma-diversity is the broad-brush approach, dealing with the flora of large areas, and it can be further partitioned into gradients within regions (beta-diversity) and the composition of communities (alpha-diversity).

The origins of floristic biodiversity in New Zealand

The size of the New Zealand flora (about 2500 vascular plants), percentage of endemism (85%; Wardle 1991), amount of endemism and the prominence of small inconspicuous flowers (Dowson 1988) all reach intermediate levels between truly isolated oceanic islands (e.g. Hawaii) and much less isolated continental islands (e.g. Britain). The traditional biogeographic explanation for this, and for some striking similarities between the floras of all the southern land-masses, is that the flora comprises an ancient ‘Gondwanan’ element, and more recent elements derived by west-wind drift from Australia (Raven 1973) or by putative ‘island hopping’ from the more tropical islands to the north. This view, expressed first as an explanation for the observed faunal and floristic similarities between South America, Tasmania and New Zealand, appeared to be vindicated when the geological evidence of plate tectonics clearly indicated that these areas were formerly connected.

However, the idea of an ancient Gondwanan flora, rafted with the original continental fragments but remaining largely intact, has recently been questioned. Pole (1994) points out that some supposedly Gondwanan species are also present on true oceanic islands, which were not part of Gondwanaland, and to which the species in question must have migrated over long sea crossings. MacPhail (1997) further points out that, if the earliest fossil records of New Zealand plants are examined, the majority appear to have been present in Australia before they occurred in New Zealand, a feature which would be consistent with long-distance transport across the Tasman Sea. The dispersalist’s argument is that, if some supposedly Gondwanan elements can indeed cross huge oceanic distances, then given sufficient time, maybe all of them could have reached New Zealand by this means! So, once again the origins of the New Zealand flora are under debate. The new techniques of molecular biology may be the means whereby the debate will be settled.

Whatever the origin of our biota, it is clear that there are high levels of endemism at the specific level in most groups, for example, 90 - 100% for Angiosperm trees and Conifers. The supposed Gondwanan element of our flora contains several small genera in primitive families, suggesting that these species are ‘palaearctic endemics’ which have survived the extinction of most other members of the group in remote isolation. In contrast, suites of endemic species in some genera can be explained as ‘neoe- endemics’ resulting from speciation in new environments following immigration. Some of the larger genera of alpine plants, e.g. Celmisia (59 species), Ranunculus (45), and Epilobium (57), appear to have reached New Zealand and undergone adaptive radiation in the new mountainous habitats of the Pleistocene (Raven 1973). Such species, genetically close to each other and to related members of the genus elsewhere in the world, are generally thought of as having less ‘value’ than those taxonomically and geographically isolated organisms regarded as palaearctic endemics. Of course, there is always a problem with these arguments, illustrated by the tendency of taxonomists to split large genera into smaller ones. The attitudes of taxonomists (to the grouping of patterns of variation into species or genera) are central to any assessment of biodiversity.

The broad brush - patterns of biodiversity on the New Zealand landscape

Many species have localised distributions within New Zealand and are thus regional endemics. More widespread species often show marked disjunctions in their distributions. Although there is a general tendency for decreasing total diversity with increasing latitude, a much stronger pattern in which areas of high and
low endemism alternate down the country is superimposed (Fig. 1). These patterns have been discussed by Wardle (1963), Burrows (1965), McGlone (1985) and Atkinson (1994), with the two later authors drawing on additional field exploration and (unpublished) taxonomic work by A.P. Druce. Centres of endemicity are generally associated with topographically varied but geologically stable landform units (McGlone 1985).

Regions of high endemism are also regions in which the total floristic list (gamma diversity) is greatest. This is partly due to the inclusion of the endemics, and partly because some more widespread species are not present in the intervening areas of low endemicity. The two major disjunctions occur in the southern North Island and the central South Island. For example, nine northern tree species (e.g. Phyllocladus trichomanoides, Libocedrus plumosa) reappear in north-west Nelson and Marlborough (McGlone 1985), while several alpine species are absent from the waist of the South Island (Burrows 1965). Likewise, the genus Nothofagus is disjunct west of the Alps, though the environment there appears suitable and all species are found to both the north and the south. McGlone (1985) draws attention also to a third disjunction, between the high-altitude areas of the central North Island, and the South Island mountains (see also Rogers 1983; Park 1983). McGlone (1985) points to the geological and historical links between this area and the Nelson-Marlborough region, which have separated by movement along the Alpine fault during the Tertiary. Wardle (1963) and Burrows (1965) emphasise high rates of extinction in the intensely glaciated region between them. Both factors may have contributed to the pattern. The apparently low number of endemics on Stewart Island may simply reflect the much smaller area compared to the others under consideration.

The geographical patterns of plant diversity described above are paralleled by distribution patterns in geckos and skinks (Pickard and Towns 1988), large land snails (Powell 1979, Meads et al. 1984), large insects (Meads 1990) and earthworms (Lee 1959). Regions of relative tectonic stability, in some cases periodically isolated as islands or escaping the worst effects of the last glaciation, apparently constitute refuges for palaeo-endemics and local centres of diversity (Ross 1972). Park (1983) drew attention to the urgency for conservation efforts in the Northland and Nelson-Marlborough regions. This known biogeographic pattern, which is an overall feature of the New Zealand biota, should be specifically targeted in New Zealand’s biodiversity strategy.

In considering the contrasts between regions of high and low species diversity one can ask whether high diversity is a result of more species per community sample (high alpha-diversity), or a result of a finer mosaic of community types (higher beta-diversity). For example, are species more tightly packed along altitudinal gradients in Nelson than in the Tararuas? How do different forest types differ in alpha-diversity? These sorts of questions are conceptually straightforward, but they imply a considerable amount of field effort in geographically separated regions. Relevant data focused on such questions are not yet available, but some instructive comparisons can be made.

**The altitudinal gradient of plant species richness**

It is well known that the number of woody species generally declines with an increase in altitude (e.g. Whittaker 1967; Beals 1969; Peet 1981) and this has been described in several studies in New Zealand (e.g. Ogden 1971; Clarkson, 1986; Druitt et al. 1995). In a quantitative study of species distributions on nine South Island altitudinal sequences, Allen et al. (1991) indicated that altitude accounted for 44 ± 5% of the variation in species composition. Other components of variation included geologic substrate and disturbance history. Simple presence/absence data for the woody flora on altitudinal gradients throughout New Zealand are illustrated in Fig. 2. Forest clearance has resulted in few of the North Island sequences extending below 400 m altitude, but some ‘spot’ data from forests at lower altitudes have been included. The overall impression is of diversity decreasing in a fairly consistent way as altitude increases. The range of possible woody species diversity at any one altitude (i.e. the range of alpha-diversity) is about 35-40 species, truncated to a smaller range in the subalpine zone. Some of the spread in the data can be regarded as sampling error, but it is noteworthy that even geographically close sequences sampled with the same methods show marked differences in alpha-diversity at some altitudes (Ogden 1995).

The least squares linear regression line fitted to these data accounts for 65% of the variance, and indicates an average loss of 3.4 woody plant species for every 100 m of altitude gained. This should be regarded as an expected or null value of the rate of decline in ‘species packing’ with altitude against which individual sequences can be assessed.
If we take 60 species as the sea-level value, then reduce species number with altitude on the assumption that the number is proportional to the area available on a perfect cone, we get a 'regression' line which is indistinguishable from the least-squares line (lower line on Fig. 2). Of course, the mountains studied are not all conical, but the volcanic Mt Egmont comes close, and shows the closest fit to this line. This analysis, and other evidence (Ogden 1995), implies that the simplest explanation for woody species diversity at any altitude below tree-line is the area of land available at that altitude. This prediction agrees with the generalisations proposed in the general theory of island biogeography of MacArthur and Wilson (1967), and suggests that rare species will be present at rare altitudes (isolated mountain summits) and that maximum diversity will generally be reached in the lowlands. It also predicts that high altitude plateaux will be important centres of diversity.

![Fig. 2. Total numbers of woody species (trees, shrubs, sub-shrubs and lianes) in samples from different altitudes in New Zealand. The species number at any altitude includes any species recorded at both higher and lower altitudes. It is the number 'potentially present' (Drutt et al. 1990) at any altitude rather than the actual number in a sample plot. The upper regression line is defined by: \( y = 59.914 - 0.0432x \) (\( r^2 = 0.6655 \)). The lower line assumes 60 species at sea-level and a progressive reduction with altitude based on the declining surface area of a cone with its apex at 1500 m (see text). The solid data points are for Mt Egmont (unpublished data, but see Clarkson 1986). See Ogden (1995) for data sources.]

Case study: kauri and mountain beech

Kauri forest, characteristic of the high gamma-diversity region of the northern North Island, is generally regarded as the most diverse forest community in New Zealand. Canopy and sub-canopy composition varies across gradients of topography and altitude (Burns and Leathwick 1992). Kauri itself is distributed in patches, often on ridge crests or upper north-facing slopes. These patches have a cohort structure suggesting regeneration following local catastrophic destruction (Ogden et al. 1987). In contrast, mountain beech forest is almost a monoculture (low alpha diversity). The species can grow on a wide range of substrates and over a broad altitudinal range, thus 'preventing' the development of high beta-diversities also. In the Craigieburn Range (low gamma-diversity region of the central South Island), this type of forest clothes valleys, slopes and ridges, often with only one or two scattered representatives of other trees and shrubs per hectare. This aggressive occupancy of sites is maintained by a catastrophic or 'dieback' system of regeneration and relatively high juvenile growth rates (Ogden 1988; Ogden et al. 1993).

The differences in geographical extent, structure and diversity between kauri and mountain beech forest are outlined in Table 1. The comparison suggests that particular forest types, defined by particular canopy dominants, have characteristic levels of alpha-diversity. Though kauri forest is much more restricted in latitudinal and altitudinal range, it has about double the tree species diversity of mountain beech forest. Twice as many species in half the altitudinal range implies tighter species packing along that gradient in kauri compared to mountain beech forest. Burns and Leathwick (1992) demonstrated that, at Waipoua forest, the topographic gradient from ridges to gullies (controlling soil moisture and nutrient availability) is more important in determining species composition than is altitude alone. Thus, where more species are present we witness more niche differentiation along a variety of gradients. Communities with high alpha-diversity may be more 'stable' in the sense that their component populations show less fluctuation in abundance, but they are less 'resilient' than simpler systems: mountain beech forest springs back readily following destruction, while kauri forest undergoes a long succession. This suggests that high alpha-diversity can be sustained only in association with high beta-diversity - a landscape mosaic of successional stages.

Biodiversity and plantation forests

Pine plantations, mainly *Pinus radiata*, cover about 5% of the New Zealand landscape (Newsome 1987) - about the same as kauri. Such plantations are not the 'biological deserts' they are sometimes said to be. The exotic monocolonial appearance of these forests has obscured the fact that pine plantations in New Zealand sometimes have quite high plant species richness in their understoreys (Ogle 1976, 1989a; Allen et al. 1995a; Ogden et al. 1997), and have value in maintaining populations of native birds (Gibb 1961; Clout and Gaze 1984; Allen et al. 1995b). Plantation forestry involves the development of a crop and an associated assemblage of adventive and native species which increases in indigenous diversity through time. Because this is normally destroyed during harvest and replanting, the level of indigenous plant species richness in planted areas is largely a question of the rotation time and the proportion of the landscape left in indigenous cover (Ogden et al. 1997). As in native communities, the build-up of alpha-diversity on any one location is a function of the 'disturbance regime' (felling) in the surrounding landscape and the consequent level of beta-diversity. Plantation forestry can move to sustainable operational practices which retain indigenous biodiversity within the plantation forest matrix if consideration is given to rotation times (disturbance frequency), the proportions of the landscape in different aged pine cohorts (seral stages), and the amount and spatial pattern of areas left in native forest cover (the beta-diversity of the landscape).

Conservation of plant biodiversity

Since human effects are relatively recent, 'natural' and 'semi-natural' landscapes still cover about 50% of New Zealand (Norton 1989a,b), and the country possesses a range of ecosystems from highly-modified to near pristine (Towns and Ballantine 1993). Aspects of vegetation ('habitat') loss and modification by introduced plants and animals are reviewed by Norton (1989b) and Atkinson and Cameron (1993). Loss of indigenous species diversity is widely regarded as the most apparent and serious effect of browsing by mammals.

The introduction of mammalian pests has been halted, but alien plants continue to arrive in New Zealand at the rate of about 11 new species per annum. Huge efforts have been expended since the 1930s to control introduced mammals and techniques are researched and refined each year. Meanwhile, the import of exotic (garden) plants and their subsequent control as noxious weeds is accorded relatively low priority. Weeds are already changing many ecosystems, but continued importation implies an insidious long-term problem in sustaining native plant communities.
The Threatened Species Unit of the Department of Conservation has devised a system of priority ranking using criteria which can, supposedly, allow comparisons across major taxonomic boundaries (Molloy and Davies 1992). This system indicates that vascular plants comprise the largest group of threatened organisms in New Zealand, with 40 plants classed as highly endangered (category A) and a further 44 ranked as requiring some recovery work in the short term. Wilson & Given (1989) claim that ten per cent of New Zealand’s flora is under threat of extinction in the wild. In 1991 the threatened plant list numbered 368 species and varieties (about 16% of the vascular flora) (de Lange and Taylor 1991; Wassilieff 1992). Most conservation expenditure, however, is devoted to the protection of the endangered avifauna.

Detailed vegetation data covering all parts of New Zealand (Wardle 1991) emphasise the unrepresentative nature of existing reserves (Mark 1983, 1985). Mountains are high-altitude ‘islands’ in a ‘sea’ of lowland. However, this sea of diverse forest in the lowlands is now itself highly fragmented, imposing new problems for the conservation of its biodiversity.

Realisation that lowland forest below 300 m altitude, tussocklands, swamps, estuaries and dune systems were under-represented in reserves gave impetus to the Protected Natural Areas Programme (PNA) initiated in the early 1980s (Mark 1983). Using differences in landform, vegetation, biota and climate, the country was divided into 268 Ecological Districts, and these were to be surveyed for areas requiring protection (McEwen 1987).

Table 1. Comparison between kauri (Agathis australis) and mountain beech (Nothofagus solandri var. cliffortioides) forests.

<table>
<thead>
<tr>
<th>Structure</th>
<th>Kauri forest ((\text{m}^3/\text{ha}))</th>
<th>Mountain beech forest ((\text{m}^3/\text{ha}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basal area m(^3)/ha(^1)</td>
<td>51 - 117</td>
<td>42 - 56(^*)</td>
</tr>
<tr>
<td>Height m</td>
<td>30 - 40</td>
<td>16 - 19</td>
</tr>
<tr>
<td>Density, trees ha(^1)</td>
<td>337 - 821</td>
<td>1409 - 1685(^*)</td>
</tr>
<tr>
<td>Diversity</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vascular plant spp. per plot(^*)</td>
<td>20 - 44</td>
<td>4 - 29</td>
</tr>
<tr>
<td>Total vascular plant spp.</td>
<td>231</td>
<td>173</td>
</tr>
<tr>
<td>Tree species ha(^1)</td>
<td>8 - 27</td>
<td>2 - 23</td>
</tr>
<tr>
<td>Total tree species(^*)</td>
<td>69</td>
<td>44</td>
</tr>
<tr>
<td>Disturbance</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Frequency at landscape scale</td>
<td>500 - 1000 years</td>
<td>50 - 200 years</td>
</tr>
<tr>
<td>Recovery after windthrow</td>
<td>Succession back to forest</td>
<td>Rapid in-situ regeneration</td>
</tr>
<tr>
<td>Recovery after fire</td>
<td>Succession back to forest</td>
<td>Often loss of site to grassland</td>
</tr>
<tr>
<td>Geographical extent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Latitudinal range</td>
<td>4° (34 - 38°S)</td>
<td>10° (36 - 46°S)</td>
</tr>
<tr>
<td>Altitudinal range</td>
<td>700m</td>
<td>1500m</td>
</tr>
<tr>
<td>Area 10° ha(^*)</td>
<td>0.2</td>
<td>2.5</td>
</tr>
</tbody>
</table>

Notes: (1) Data from Ahmed (1984) and Ahmed and Ogden (1991) except where noted otherwise. Based on point-centred-quarter sampling of 25 1 ha stands in mature kauri forest throughout the latitudinal range of the species.
(2) Data from Wardle (1970 part 2), except where noted otherwise. Based on 2400 plot descriptions generally covering about 0.05ha. The data presented relate to the 19 associations with > 50% cover of trees.
(3) From Wardle (1984; Tables 11.3 and 11.4); average of all mountain beech forest types in proportion to the number of plots included.
(4) Based on 20 5m-diameter circular plots in each kauri stand, but plants < 1m tall not recorded, consequently the figures are an underestimate.
(5) From Cockayne (1928); mountain beech data from Cockayne’s sub-alpine (beech) forest community, and so the difference between the two communities is again underestimated.
(6) Trees > 10cm diameter (dbh) in the case of kauri, but > 5cm dbh in the case of beech.
(7) All tree species recorded in all plots.
There is a need in New Zealand to integrate the findings of plant ecology and biogeography into the broad conservation strategy. Patterns of plant species distribution and bio-diversity in New Zealand are known, but this information is not clearly reflected in the reserve network. As a generalisation we seek to conserve vegetation types which reflect the pre-European vegetation pattern. In aggregate these reserves will, we hope, preserve the character of the New Zealand landscape and its unique biota. High species diversity is regarded as desirable, and is achieved both by creating reserves which include a variety of different plant communities and by selecting examples of communities which retain high diversity. One approach would be to identify the largest remaining lowland forest reserves, and the most intact lowland to montane altitudinal sequences in each biogeographic area of different gamma-diversity in New Zealand, and accord them high priority for the control of introduced mammals and invasive plants.

Acknowledgements
Discussions with Rob Allen, Ian Atkinson, Bruce Burns, Ewen Cameron, Matt McGlone, Susan Timmins and Carol West helped to formulate the ideas on which my original paper (Ogden 1995) on this topic was based. The Robert C. Bruce Trust supported my research on diversity in pine plantations, and Carter Holt Harvey Forests gave permission for the work in Kineleith Forest.

References


changes in biodiversity. Implicit in that evolutionary terms) and irreversible human-directed influences can be managed, particularly the use of beetles as an indicator group (Hutcheson, 1994). The rationale for this work is to identify changes in the composition of insect communities that will reflect underlying changes in their habitat. Future research will include studies of indigenous and exotic understorey plant species as indicators of biodiversity.

Just as a large component of biodiversity is concerned with the richness expressed by species variability, so too is genetic diversity within a species of significant importance. The genetic diversity of a production crop needs to be protected, maintained, and if necessary enhanced as part of responsible management of our forest genetic resources.

The genetic resources of New Zealand's major exotic forest tree species have been managed since the 1950s, including low-intensity programmes of species and provenance introduction, test-