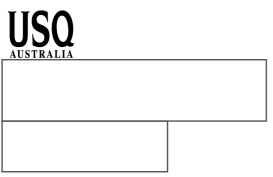
Pasture and habitat monitoring for grazing management 'USQ13' - LWW/AWI project

Progress Report: Vegetation changes following the exclusion of grazing in the Traprock region



February, 2008



|A collaborative research project involving:

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Pasture and habitat monitoring for grazing management

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Contents

Report Authors i Acknowledgements i Contents ii Summary iii 1. Introduction 1 1.1 General Introduction 1 1.2 Project Aims 3 2. Study Area & Methods 4 2.1 Study Area, Sample Sites and Experimental Design 4 2.2 Data Acquisition & Analysis 9 3. Interim Results 14 3.1 General Results 14 3.2 Floristic Composition 14 3.3 Species richness and plant biomass 19 4. Discussion 26

5. References 30



Summary

This study is the continuation of previous research that examined the effects of vegetation management on woodland communities in the Traprock region and will contribute to an understanding of the 'biodiversity potential' of managed land units on properties (including open paddocks, scattered treed areas, and remnant woodland) identified previously. In order to determine 'biodiversity potential', a number of grazing exclosures were established on selected properties with the aim to monitor biodiversity changes over-time following the removal of grazing.

Eighteen study sites across 10 properties in the Traprock region were selected for this study. The design consisted of: 2 vegetation types (grassy box (*Eucalyptus melliodora, E. microcarpa,* or *E. moluccana*) woodland (L) and ironbark (*E. crebra*)/gum (*E. dealbata*) (U) woodland); 3 mature (overstorey) tree densities (<6 trees/ha [low](L); 6-20 trees/ha [medium](M); >20 trees/ha [high](H), and; 3 exclosures (full exclosure (1) [2.5m complete fence], partial exclosure (2) [1.5m three-wire fence], open (3) [corner makers]). Exclosure plots were erected in January/February 2005 and sampled in April 2005 (two months after exclosure establishment), February 2006 (12 months after exclosure establishment), February 2007 (2 years after exclosure establishment), and February 2007 (3 years after exclosure establishment). Within each 6 x 6 m exclosure plot, a central 2 x 2 m quadrat was sampled for plant species cover. Above-ground vegetation ('biomass') was clipped in a separate 0.25m² sample and dry weight determined. Overstorey cover and recruitment were determined within each 6 x 6 m exclosure plot. Stand structural characteristics, including foliage projective cover of distinct strata, and cover of litter, logs and rocks, and general habitat condition were also determined at each site.

Preliminary patterns in floristic composition were determined using non-metric Multidimensional Scaling (nMDS). Two-way crossed Analysis of Similarity (ANOSIM) determined whether there were significant differences in floristic composition between exclosure types and mature tree density classes. nMDS was also used to assess patterns in cover data for growth forms. Two-way crossed Analysis of Variance (ANOVA) was used to determine if groups (density class, exclosure type and the interaction of density and exclosure) differed significantly for total, native, exotic, growth-form, perennial, and annual species richness and above-ground biomass. In addition, Spearman-rank correlations were performed to determine if biomass and estimates of ground cover were related.

A total of 151 plant species was recorded across all treatments with an average richness across treatments of 17 species per $4m^2$. No differences were apparent in overall plant composition (cover) between the exclosure treatments one year following grazing exclusion (ANOSIM, pr > 0.05). nMDS ordinations show no distinction between exclosure treatments, but patterns were observed in mature tree density treatments within vegetation types as found in previous research. Stand structure (cover of strata) showed much the same pattern as floristic composition.

There were generally no detectable differences (p > 0.05) in plant above-ground biomass between exclosures, although significant differences between tree density classes was indicated with a significantly higher plant biomass in low density treatments compared to high density for both vegetation types. Correlation results showed that estimates of grass cover provide a good indication of above-ground biomass (p<0.05). There were some differences between density classes for growth-form species richness, exotic species richness and annual species richness, however exclosure treatments did not differ. Overall, there were no differences in total or native species richness between groups.

While the distinction between vegetation type and mature tree density is observed in species composition, plant biomass and species richness, the exclusion of grazing (native and exotic) has not significantly altered composition after 3 years. The patterns in floristic composition are associated with different mature tree densities and vegetation type, which are consistent with earlier findings. There is some evidence to suggest that plant above-ground biomass

has responded to the removal of grazing in open paddock areas, although this is not consistent across mature tree density treatments. It is suggested that a longer period of exclusion will be necessary to detect changes (if any) in plant species composition.

1. Introduction

Woodland ecosystems can provide a number of important services for agricultural production such as soil formation and protection, nutrient storage and cycling, natural control of diseases and parasitic organisms, insect pollination for seed and fruit set, and the breakdown and absorption of pollutants (McIvor and McIntyre 2002). Woodlands also provide critical wildlife habitats (e.g. for possums, birds, and bats) (McIntyre 2002; Lumsden and Bennett 2005), and habitat diversity, which may facilitate the establishment of different native plant species (Chilcott *et al.* 1997). Unfortunately, much of Australia's woodland ecosystems have been cleared to make way for agricultural production. It is estimated that 500 000 km² of woodlands have been cleared since European settlement (AUSLIG 1990). As a result, some woodland communities are amongst the most poorly conserved ecosystems in Australia (Yates and Hobbs 1997).

It is recognised that where woodlands have little or no representation in nature reserves, biodiversity conservation may best be achieved if combined with current production systems (McIntyre 1994; Chilcott *et al.* 1997). Pastoral grazing lands that contain semi-intact woodlands (variegated landscapes) may be compatible with conservation outside reserves if managed appropriately (McIntyre and Barrett 1992; McIntyre 1994; Chilcott *et al.* 1997). This requires understanding how management practices within pastoral gazing lands affect vegetation characteristics in woodland communities.

Pastoral land management in Queensland often involves removing or reducing the tree layer to increase native grass production for livestock grazing (McIvor and McIntyre 2002). Two major anthropogenic disturbances influencing woodland communities in variegated landscapes are livestock grazing and the associated modification of natural tree densities (McIntyre and Barrett 1992; Clarke 2003). The combination of both livestock grazing & clearing in grassy woodlands can be detrimental to many native plant species (Prober and Thiele 1995; Clarke 2003).

Grazing by sheep and cattle represents the single greatest pressure on two-thirds of Australia's agricultural land (Hamblin 2001). Since the introduction of domestic livestock there has been a major increase in the rate of defoliation of native vegetation and consequently changes in the composition and diversity of plant communities (Wilson 1990). Grazing by domestic stock has altered the composition of understorey species (Prober and Thiele 1995; Clarke 2003), prevented seedling recruitment (Tothill 1971; McIntyre and Lavorel 1994), contributed to soil erosion and compaction (Wahren *et al.* 1994; Yates and Hobbs 1997) and enhanced the invasion of exotic species (Prober and Thiele 1995; Clarke 2003). Furthermore, significant changes in fire frequency have occurred as fire has often been excluded from pastoral lands (Wilson 1990). Grazing can also contribute to soil deterioration due to soil compaction and trampling of vegetation (Yates and Hobbs 1997). Soil compaction can impede root growth reducing the ability of roots to provide plants with water and nutrients (Willatt and Pullar 1983; Yates and Hobbs 1997). In heavily grazed areas, water infiltration may be decreased, which has important implications for plant growth, reproduction and seedling establishment (Willatt and Pullar 1983; Yates and Hobbs 1997). Grazing also creates greater areas of bare ground that are susceptible to soil erosion (Wahren *et al.* 1994).

In the last 200 years, clearing native vegetation on more productive soils has resulted in the loss of large areas of woodlands in eastern Australia (AUSLIG 1990; Hobbs and Hopkins 1990; McIvor and McIntyre 2002). Remnants with near natural understoreys are now rare, with biodiversity often restricted to patches that vary considerably in size, quality and isolation (Prober and Thiele 1995; Yates and Hobbs 1997; Prober *et al.* 2002). Small woodland remnants are particularly vulnerable to loss of native species, changed soil conditions and additional disturbances, while the isolation of remnant patches can result in changes to the normal dispersal and reproductive success of both plants and animals (Hobbs 1987; Prober

and Thiele 1995; Gilfedder and Kirkpatrick 1998; Ross et al. 2002; Godefroid and Koedam 2003).

In grazing landscapes, clearing trees often has a significant effect on grass production (McIvor and McIntyre 2002). Numerous studies have shown that tree density is inversely related to pasture yield in many Australian woodland communities, with often a significant increase in pasture yield when all trees are removed or killed (Walker *et al.* 1986; Harrington and Johns 1990; Scanlan and Burrows 1990; McIvor and Gardener 1995; McIvor 2001). While increased grass production is ideal for livestock grazing, the removal of trees from grazing landscapes can have negative impacts on original woodland understories. For example, Gibbs *et al.* (1999) report that clearing trees from grazing lands may result in a change in dominant grass species, from shade-tolerant grasses (e.g. *Microlaena, Danthonia, Poa*) to species that dominate long-cleared pastures (e.g. *Aristida ramosa*). The change in floristic composition has been attributed to altered microclimatic and competitive regimes, and lower soil fertility (Gibbs *et al.* 1999).

1.1 Project Aims

This project aims to ascertain the biodiversity *potential* of managed land units on properties (including open paddocks, scattered treed areas, and remnant woodland). To gauge this, a number of grazing exclosures were established on selected properties with the aim to monitor biodiversity changes over time following the removal of grazing.

Studies that have examined the response of native vegetation to grazing exclusion elsewhere in Australia have reported diverse results (e.g. Tremont 1994; Pettit and Froend 2001; Spooner *et al.* 2002). These studies demonstrate that it is often hard to predict the response of vegetation to grazing exclusion and results may largely depend on differences in environment, grazing history and length of grazing exclusion.

The specific objectives of this study are to:

- 1. determine the response of vegetation to the short-term exclusion of grazing in the Traprock region by examining the floristic composition, species richness and above-ground biomass of exclosure plots (complete, partial and open/control).
- 2. determine if tree density and/or vegetation type are important in explaining patterns in plant communities.
- 3. determine if tree density, vegetation type or exclosure type (complete, partial and open/control) interact to influence floristic patterns.

2. Study Area & Methods

2.1 Study Area, Sample Sites & Design

The study is being undertaken in the Traprock wool-growing region, west of the Stanthorpe-Wallangarra granite belt in southern Queensland. The region is approximately bounded by the major towns of Warwick and Stanthorpe to the east and Inglewood and Texas to the west. The climate within the region is influenced by both tropical and temperate weather patterns. The average minimum and maximum temperatures for the region range between 2.6 - 30.2°C for Warwick (28°22′S, 152°03′E) and 0.9 - 27.4°C for Stanthorpe (28°66′S, 151°93′E) (Bureau of Meteorology 2005). Rainfall is generally higher in summer months, although the winter proportion can be significant (Wills 1976). The average rainfall ranges from 701 mm at Warwick to 770 mm at Stanthorpe, but typically declines to the west of the study area (Wills 1976; Queensland Murray Darling Committee 2004).

Currently, the Traprock region supports approximately 300 000 hectares of sheep grazing country at a stocking rate of about 1-2 dse (dry sheep equivalent) per hectare (Queensland Murray Darling Committee 2004). Wool production is the dominant land-use, with limited winter and summer cropping and horticulture (Wills 1976; Queensland Murray Darling Committee 2004).

The vegetation of the Traprock region is predominately grassy eucalypt woodland mainly comprised of narrow-leaved ironbark (*Eucalyptus crebra*), tumbledown gum (*Eucalyptus dealbata*), and white box (*E. albens*) and, on the lower slopes, yellow box (*E. melliodora*), grey box (*Eucalyptus microcarpa*) or gum topped box (*Eucalyptus moluccana*). Remnant patches have been mapped by the Queensland Herbarium and classified as Regional Ecosystems 13.11.3/13.11.8, respectively (Environment Protection Agency 2003). Both regional ecosystems are listed as of concern as a result of both grazing and clearing within the region (Environment Protection Agency 2003).

Native pastures are the main source of forage for grazing livestock within the study area (Wills 1976). Past land management practices have largely consisted of ring-barking woody plants to encourage the growth of grasses and burning regularly to control woody regrowth and to remove unpalatable dry herbage (Wills 1976). Natural grasslands are not considered to have been a common component before settlement (Wills 1976). Regional ecosystem mapping by the Queensland Herbarium shows approximately 22 % of remnant vegetation remains in the Traprock region (Queensland Murray Darling Committee, 2004).

To ensure the major community types were sampled, site selection was stratified across the study area according to vegetation type and density of mature trees. Satellite imagery was used to select potential sample sites and actual sites were chosen in the field if all criteria were met. Two dominant woodland communities were recognised using the regional ecosystem mapping by the Queensland Herbarium. The grassy box woodlands (RE 13.11.8) typically occur on lower slopes and are distinguished from ironbark/gum woodlands (RE 13.11.3) that typically occupy upper slopes and ridge lines. The box woodlands are dominated by *E. melliodora* (yellow box) and *E. microcarpa* (grey box), while *E. tereticornis, E. albens* and *Angophora floribunda* are occasional components of the community (Wills 1976). The box woodland community generally lacks a well developed shrub layer, but occasionally *Acacia* spp. and *Cassinia* spp. may form a dense shrub layer to two metres tall (Wills 1976). The ground

layer component is moderately dense and dominated by *Cymbopogon* spp., *Bothriochloa* spp., *Austrodanthonia* spp., *Dichelachne* spp., *Stipa* spp. and *Aristida* spp. (Wills 1976). For the purposes of this study, the box woodlands are labelled as lower (L) slope vegetation. The ironbark/gum woodlands are dominated by *E. crebra* and *E. dealbata* and occasionally *Angophora costata* (Wills 1976). *E. crebra* may be replaced by *E. sideroxylon* in some areas. The ground layer is typically sparse, but has a well developed shrub layer consisting of *Acacia* spp., *Jacksonia* spp., *Leucopogon* spp., *Daviesia* spp. and *Olearia* spp. (Wills 1976). For the purposes of this study the ironbark/gum woodlands are labelled as upper (U) slope vegetation.

Within each of these vegetation types, sites were assigned to one of three mature tree density classes: low (<6 trees/ha), medium (6-20 trees/ha) or high (>20 trees/ha) based on the number and cover of mature trees observed on the satellite imagery. Only sites with an absence of woody regrowth in the understorey were included. Potential sites were excluded if the vegetation patch was less than 5 ha in size, if fence lines and water points were less than 250 m from the patch, and if areas were recently cleared (<5 years ago) or burnt (< 10 years ago). Ease of access to sites, spatial spread of sites across the study area, and landholder's permission to access properties were also considered in site selection.

The final experimental design consisted of:

- (two vegetation types (ironbark/gum woodlands; box woodlands); and
- (three mature tree densities (<6 trees/ha [low]; 6-20 trees/ha [medium]; >20 trees/ha [high]).

Six treatment combinations were recognised (Table 1) and a total of 18 sites across ten properties were selected (Figure 1). Three 6 x 6 metre exclosure plots were erected in January/February 2005 at each of the eighteen sites (totalling 54 exclosure plots). The exclosure plots consist of:

- **1. complete exclosure**: 2.5 metre fence to exclude both sheep and large herbivores (e.g. kangaroos);
- 2. partial exclosure: 1.5 metre three-wire fence to exclude sheep only;
- 3. open (control): corner markers, no fence (to allow grazing).

Exclosure plots were erected at locations representative of the vegetation at each site. Plots were a minimum distance of 20 m apart and a minimum of 20 m from the boundary of the patch to minimise edge effects.

Table 1. Description of treatment combinations.

Abbreviated description (label), the number (n) of replicates for each treatment combination and site numbers are indicated.

Site description	Label	n	Site numbers	
Low density; ironbark/gum woodland	LU	3	1, 2, 3,	
Low density; box woodland	LL	3	4, 5, 6	
Medium density; ironbark/gum woodland	MU	3	7, 8, 9	

Medium density; box woodland	ML	3	10, 11, 12	
High density; ironbark/gum woodland	HU	3	13, 14, 15	
High density; box woodland	HL	3	16, 17, 18	

Figure 1. Map of Traprock study area showing location of exclosure sites.

2.2 Data Acquisition & Analysis

Sites were sampled in April 2005 (two months after exclosure establishment), February 2006 (12 months after exclosure establishment), February 2007 (2 years after exclosure establishment), and in February 2008 (3 years after exclosure establishment). Within each 6 x 6 metre exclosure plot, a central 2 x 2 metre quadrat was sampled for ground cover and vascular plant species composition. Ground cover was determined by subjectively estimating the percent cover of forbs/herbs/other (non-woody), graminoids (grasses/sedges), logs and branches >10cm in circumference, course litter (twigs and branches, 4-10 cm circumference), fine litter (leaf and twigs, < 4 cm circumference), rock cover, bare ground and cryptogams within each 2 x 2 metre quadrat. The average heights (centimetres) of grasses were also recorded within each 2 x 2 metre quadrat by measuring the height of four different grasses with a ruler and taking the mean.

The composition of plant species was determined by estimating the percent cover of vascular plants within each 2 x 2 metre quadrat. Plant species that could not be identified in the field were collected and later identified using the nomenclature of Harden (1991), Stanley & Ross (1983, 1986, 1989) and Henry *et al.* (1995). Some plant species that were not flowering or seeding at the time of sampling were identified to genus level. Exotic species were distinguished as any plant species that has been introduced into Australia and identified using the nomenclature of Stanley & Ross (1983, 1986, 1989) and Auld & Medd (1987).

Overstorey cover and recruitment were also recorded for each exclosure plot. Overstorey cover was determined by subjectively estimating the percentage foliage cover of trees >10 m, trees <10 m, shrubs >2 m and shrubs <2 m within each 6 x 6 m exclosure plot. Recruitment was determined by recording the number of individual juvenile trees >1 m, juvenile trees <1 m, and shrubs, and by scoring epicormic regrowth and suckering from 0 (none) to 3 (high) according to their percent cover within each 6 x 6 m quadrat (where 0=0%; 1=<10%; 2=10-20%; 3=>20%).

Above-ground vegetation was clipped in a separate 0.25 m^2 quadrat within each 6 x 6 m exclosure plot. Plant biomass (gm/0.25m²) was determined by drying the samples in paper bags in a Thermoclime at 50°C for 4 days in 2005, and at 60°C for minimum of 3 days in 2006 and 2007. Samples were left to cool for five minutes to room temperature before weighing on scales to determine biomass in grams.

At each site, stand structure and condition were recorded. Site stand structure was determined by using the modified Specht (1981) structural classification. Based on the vegetation of the study area, seven strata were pre-defined: trees >20m, trees 10-20 m, trees <10 m, shrubs >2 m, shrubs <2 m, Forbs/Herbs/Other (non-woody species), and graminoids (including grasses, sedges and others). Trees are defined as single or multi-stemmed woody plants greater than 3 m in height and shrubs are defined as multi-stemmed plants less than 3 m in height. The percentage foliage cover of each stratum was subjectively estimated at each site. In addition, the cover of logs >100 cm in circumference, rock cover, and the cover of course litter (twigs and branches, 4-10 cm circumference) and fine litter (leaf and twigs, < 4 cm circumference) were estimated and recorded for each site. The condition of the site was determined by recording evidence of disturbance. Disturbances included grazing, clearing, logging, erosion, weeds, feral animals, soil compaction, bare ground and canopy death and were subjectively scored from 0 (no evidence) to 3 (high) depending on the level of impact at each site.

All analyses described below have been performed on 2005-2007 only. 2008 monitoring data is currently being collated and processed.

Non-metric multidimensional scaling (nMDS) was performed on preliminary plant cover data and stand structure data using the Primer v.5.2.9 for Windows computer program (Primer-E Ltd 2001) to determine the dissimilarity relationship between sites (Clarke and Warwick 2001). nMDS is an ordination method that constructs a map or configuration of the sites in a specified number of dimensions, with sites closer together more similar (eg. in species composition) than those further apart (Clarke and Warwick 2001; Quinn and Keough 2002). The stress value provided with the ordination indicates how well the ordination shows the relationship between sites (McGune and Grace 2002). Higher dimensions (e.g. 3-D and 4-D) tend to decrease stress; however, a 2-dimensional ordination allows a reasonable visual summary of site relationships if the stress value is 0.2 or less (Clarke and Warwick 2001). nMDS was performed on all exclosure and site plant cover data, as well as on a subset to determine the relationship between sites in ordination.

Prior to performing nMDS, the Primer v.5.2.9 for Windows computer program (Primer-E Ltd 2001) was used to compute Bray-Curtis similarity matrix on species cover data to allow sites to be represented

graphically and to discriminate sites from each other (Clarke and Warwick 2001). Bray-Curtis similarity coefficient is widely accepted as a satisfactory coefficient for biological data on community structure (Clarke and Warwick 2001).

Two-way crossed analysis of similarity (ANOSIM) was performed on plant cover data for ironbark/gum woodlands and box woodlands using the Primer v.5.2.9 for Windows computer program (Primer-E Ltd 2001). A Bray-Curtis similarity matrix was computed and the two-way crossed ANOSIM test performed on the cover data to determine if there were differences between exclosure plots, allowing for the fact that there may be density differences or (vice versa) to determine if there were differences between tree densities (Clarke and Gorley 2001). ANOSIM is a hypothesis testing procedure comparing between-group and within-group variation using rank similarities (Quinn and Keough 2002). The test statistic (R) is scaled to be within the range +1 to -1 and is not overly affected by the number of replicates in the two groups being compared, whereas the statistical significance is dominated by group sizes (Clarke and Warwick 2001). ANOSIM provides a global R value for overall differences between groups, with large values (close to 1) indicating complete separation of groups are. An R value >0.75 indicates groups are well separated, R >0.5 indicates groups are overlapping but clearly different, and R<0.25 indicates groups are barely separable (Clarke and Gorley 2001).

SPSS® for Windows version 11.5 (SPSS Inc. 2002) was used to perform two-way crossed ANOVA (Analysis of Variance) to determine if there were differences in total species richness, shrub species richness, graminoid species richness, herb/forb species richness, native species richness, exotic species richness, perennial species richness, annual species richness and plant biomass for each vegetation type due to density class (low, medium, high) or exclosure type (complete, partial, open) or the interaction of density and exclosure (all species richness data were log transformed to improve normality and reduce the influence of any outliers).

Prior to performing two-way crossed ANOVA, the Levene's statistic and residual plots were used to test homogeneity of variances in SPSS® for Windows version 11.5 (SPSS Inc. 2002). In addition, Spearman-rank correlations were performed using SPSS® for Windows version 11.5 (SPSS Inc. 2002) to determine whether biomass and estimates of ground cover were related and the degree to which they vary together (Ashcroft and Pereira 2003).

3. Interim Results

3.1 General Results

A total of 151 plant taxa (135 native, 16 exotic) were recorded across the study area. Total species richness ranged from 2 to 29 species, with an average richness over all samples of 17 species per 4 m².

3.2 Floristic Composition

Non-metric multi-dimensional scaling (nMDS) ordination of floristic composition data (Figure 4) (stress=0.21) shows a clustering of sites to the left of the diagram, with one high density ironbark/gum woodland site (site 15) separated from this major cluster. The ordination shows a clustering of samples according to site rather than exclosure type. The large cluster shows that sites with similar mature tree densities tend to cluster together (with some exceptions). Low tree density sites (LU and LL) are largely grouped to the left, with medium density ironbark/gum woodlands (MU) immediately to the right. Medium and high density box woodlands (ML and HL) are slightly separated from this cluster with high density ironbark/gum woodland sites (HU) separated towards the end of the major cluster. Based on previous evidence of differences in vegetation types (see Goodhew 2005), floristic data were separated by vegetation type and a nMDS ordination performed on each data set.

nMDS ordination of floristic composition data for ironbark/gum woodland sites (Figure 5) (stress=0.17) shows a gradient of increasing mature tree density from left to right across the diagram. Low density sites (LU), regardless of exclosure type were shown to cluster to the left of the diagram, with medium density sites (MU) in a loose cluster immediately to the right. High density sites (HU) were reasonably separated from low and medium density sites, representing the other end of the cluster.

nMDS ordination of floristic composition data for box woodland sites (Figure 6) (stress=0.23) shows a closer clustering of samples. With some exceptions, the ordination shows low density sites (LL) clustering towards the left of centre with medium (ML) and high density sites (HL) in a separate cluster towards the right of centre.

Two-way crossed ANOSIM of floristic composition data for each vegetation type (Table 2) shows that the test for differences between mature tree densities was significant for both ironbark/gum woodlands (p=0.007) and box woodlands (p=0.037). However, the exclosure differences overall were not significant (see Table 2). For ironbark/gum woodlands, high mature tree density sites (HU) were significantly different in floristic composition to low (LU) and medium (MU) tree density sites. For box woodlands,

high mature tree density sites (HL) were significantly different in floristic composition to low tree density sites (LL). Medium density box woodland sites (ML) were not significantly different in floristic composition to either high or low tree density sites.

Figure 4. Non-metric multi-dimensional scaling ordination of floristic data. See Table 1 for descriptions of treatment codes.

Figure 5. Non-metric multi-dimensional ordination of floristic data for ironbark/gum woodland sites. See Table 1 for descriptions of treatment codes.

Figure 6. Non-metric multi-dimensional scaling ordination of floristic data for box woodland sites. See Table 1 for descriptions of treatment codes.

 Table 2. Two-way crossed ANOSIM of floristic data for ironbark/gum woodland sites and box woodland sites.

Global R value and significance level shown (groups sharing the same superscript letter are not significantly different).

	R value	Significance	Pairwise tests		
Ironbark woodland					
Density	0.262	0.007*	Lowa	Mediuma	Highb
Exclosure	-0.192	0.953	Completea	Partiala	Opena
Box woodland					
Density	0.193	0.037*	Lowa	Mediumab	Highb
Exclosure	-0.147	0.894	Completea	Partiala	Opena

*results are significant at the 0.05 level

3.3 Species richness and plant biomass

Two-factor ANOVA results for total plant species richness and growth-form species richness are shown in Table 3. The results indicate some significant differences in shrub species richness, graminoid species richness and herb/forb species richness between mature tree density classes (although variances are unequal). Exclosure type and the interaction of exclosure type and tree density did not contribute to differences between groups.

Two-factor ANOVA results for native and exotic species richness, perennial and annual species richness and plant biomass are shown in Table 4. The results indicate some significant differences in exotic species richness, annual species richness and plant biomass between mature tree density classes. In particular, for ironbark/gum woodlands annual species richness was significantly lower in high density sites compared to low and medium density sites. Exclosure type and the interaction of exclosure type and tree density did not contribute to differences between groups.

 Table 3. Two-factor ANOVA of total species richness and growth-form richness for each vegetation type.

 Homogeneity of variances indicated by N (no) or Y (yes). [df: 2=density; 2=exclosure; 4=density*exclosure.] Post-hoc results where applicable include means and standard errors in parentheses. Means sharing the same superscript letter are not significantly different (L=low, M=medium, H=high tree density).

*significant at the 0.05 level

Table 4. Two-factor ANOVA of native, exotic, perennial and annual species richness and aboveground plant biomass for each vegetation type.

Homogeneity of variances indicated by N (no) or Y (yes). [Degrees of freedom: 2=density; 2=exclosure; 4=density*exclosure.] Post-hoc results where applicable include means and standard errors in parentheses. Means sharing the same superscript letter are not significantly different (L=low, M=medium, H=high tree density).

*significant at the 0.05 level

Mean above-ground plant biomass $(gm/0.25m^2)$ per exclosure type for each mature tree density class and vegetation type across years are shown in Figures 7 to 12. An increase in mean above-ground biomass from April 2005 to February 2006 was indicated for low tree density treatments for both vegetation types, and medium and high tree density box woodlands sites. In the low density ironbark/gum woodland sites, mean above-ground biomass increased from an average of 28 gm/0.25m² (SE ±13.7) in 2005 to an average of 73.2 gm/0.25m² (SE ±11.5) in the complete exclosure (Figure 7). In the low density box woodland sites, the most notable increase in mean above-ground biomass was in the partial exclosure with an average of 14.8 gm/0.25m² (SE ±5.7) recorded in 2005 and an average of 62.3 gm/0.25m² (SE ±32.5) recorded in 2006 (Figure 8). There was no notable increase in average above-ground biomass between years for medium and high density ironbark/gum woodland sites.



Figure 7. Mean above-ground plant biomass (gm/0.25m²) for treatment LU (low density ironbark/gum woodland) for successive years per exclosure type

(error bars represent standard error)

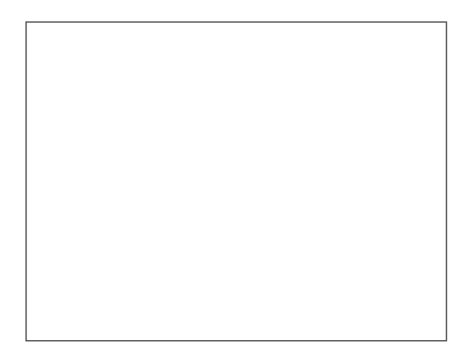


Figure 8. Mean above-ground plant biomass (gm/0.25m²) for treatment LL (low density box woodland) for successive years per exclosure type (error bars represent standard error)

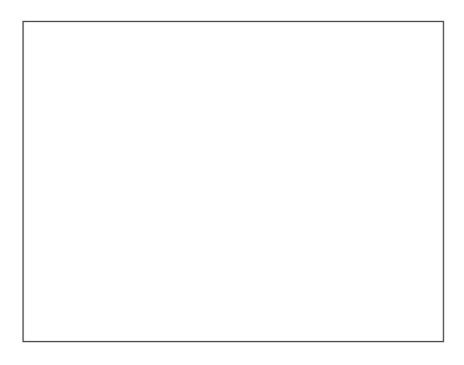


Figure 9. Mean above-ground plant biomass (gm/0.25m²) for treatment MU (medium density ironbark/gum woodland) for successive years per exclosure type (error bars represent standard error)



Figure 10. Mean above-ground plant biomass (gm/0.25m²) for treatment ML (medium density box woodland) for successive years per exclosure type (error bars represent standard error)



Figure 11. Mean above-ground plant biomass (gm/0.25m²) for treatment HU (high density ironbark/gum woodland) for successive years per exclosure type (error bars represent standard error)



Figure 12. Mean above-ground plant biomass (gm/0.25m²) for treatment HL (high density box woodland) for successive years per exclosure type (error bars represent standard error)

Estimates of ground cover (per $4m^2$) and total ground cover (per $4m^2$) were positively correlated with above-ground plant biomass (gm/0.25m²) for both sampling years (Table 5). Estimates of herb/forb cover (per $4m^2$) were not correlated with either above-ground biomass (gm/0.25m²) or estimates of grass cover and total ground cover (per $4m^2$).

nMDS ordination of stand structure data revealed that the structure of treatments were reasonably consistent across sites (Figure 13; stress 0.07). With a few exceptions, high tree density ironbark/gum woodlands were well separated from high and medium density box woodlands. Medium density ironbark/gum woodlands were separated from this cluster, while low tree density sites clustered together regardless of vegetation type, and were well separated from all other clusters.

4. Discussion

The interim results suggest that while the distinction between vegetation type and tree density is observed in overall plant species composition, the exclusion of grazing has not significantly altered composition after 12 months. This suggests that the vegetation has had little time to recover after 12 months of grazing exclusion, which agrees with results reported from other studies in Australia (Pettit and Froend 2001; Spooner *et al.* 2002). For instance, Pettit and Froend (2001) concluded that initial patterns in floristic composition after 3 years of grazing exclusion were only temporary and highly influenced by fluctuations in annual climatic patterns when exclosure plots were sampled again after 7 years (Pettit and Froend 2001). Pettit and Froend (2001) recommended that long-term monitoring is needed to detect long-term trends in floristic composition after grazing is excluded.

There was some evidence to suggest that above-ground plant biomass has responded to the removal of grazing in open paddock areas, in particular there was an increase in biomass from 2005 to 2006 in the complete exclosure for low density ironbark/gum woodland sites. However, no difference in above-ground biomass between exclosure plots was found for ironbark/gum woodland sites or box woodland sites, and there was no interaction between exclosure plot and tree density. Significant differences in above-ground biomass between tree density classes was indicated with a significantly higher plant biomass in low density box woodlands compared to medium and high density treatments, and a significantly higher plant biomass in low density ironbark/gum woodlands compared to high density treatments (although there were unequal variances).

This result does confirm conclusions made in earlier research (Goodhew 2005) and reported elsewhere in Australia, that grass cover (or biomass) declines significantly with increasing total tree cover (e.g. Walker *et al.* 1986; Harrington and Johns 1990; Scanlan and Burrows 1990; McIvor and Gardener 1995; McIvor 2001). Interestingly, the results from the correlation of grass cover (per 4 m²) and above-ground biomass (gm/0.25 m²) indicate that estimating the cover of grasses within exclosure plots will provide a good indication of plant above-ground biomass. This result is important in that more confidence can be given to subjective estimates of grass cover which are often less time consuming and costly. If landholders monitor exclosure plots in the future this result is particularly relevant.

The number of plant species recorded during this study is comparable to previous studies on woodland communities in Australia. Mean species richness has been reported as 20 species per 25 m² for woodlands in Western Australia (Yates and Hobbs 1997), 28 per 30 m² for grazed temperate grassy woodlands in New South Wales (McIntyre and Martin 2001) and 25 per 20 m² for grazed hill woodlands in south-eastern Queensland (Fensham 1998). In this study, mean species richness was 17 per 4 m² indicating the Traprock region has a significant biodiversity component (in terms of plant species). However, little evidence of overstorey species recruitment suggests that the wooded sites sampled in this study may not be viable over-time.

The recruitment of overstorey species depends on a number of factors such as suitable environmental conditions, adequate seed supply, and absence of herbivory (Clarke 2002). Possible explanations for the lack of recruitment are that there may not have been suitable environmental conditions for the germination of overstorey species since exclosure establishment (Clarke 2002), and the fact that exclosure plots were

only established 12 months ago. For example, Spooner *et al.* (2002) found a significant positive correlation between tree recruitment densities and time since fencing in woodland remnants of northern NSW. However, another possibility is that disturbance (e.g. fire) may be necessary to enhance germination. Clarke (2002) found that no natural recruitment of native shrub species had occurred after 5 years in the grassy woodlands of NSW and concluded that recruitment is episodic and disturbance driven. Experiments that examine mechanisms of shrub and tree recruitment could be important for implementing management practices specific for tree and shrub establishment in the Traprock region.

The patterns in floristic composition found in this study are similar to those reported in earlier research (Goodhew 2005) where sites were separated according to vegetation type (either ironbark/gum or box woodland) and tree density (low, medium or high). As reported previously (Goodhew 2005), high density ironbark/gum woodlands are floristically and structurally different to both low and medium density ironbark/gum woodlands, yet low and medium density ironbark/gum woodlands do not differ floristically (yet are different structurally). This result suggests that both grazing and tree clearing have resulted in a vastly different understorey in low and medium density patches, with a higher annual species richness (in low and medium density treatments), compared to that of more 'natural' ironbark/gum woodlands. However, it should be noted that there were no differences in total or native species richness between groups.

The higher annual species richness in low and medium density treatments is possibly related to the more intense grazing and tree removal practices that have historically occurred in these areas. Annuals are reported to be more tolerant of disturbance than perennials due to their fast growth rates and early and prolific seed set (Grimes 1974 cited in Pettit *et al.* 1995) and many studies within Australia have reported an increase in annual species in areas that were frequently grazed by livestock (Pettit *et al.* 1995; Prober and Thiele 1995; Clarke 2003). Similarly, the higher herb/forb richness in medium density treatments compared to high density treatments may be a response to reduced tree density/cover. Walker *et al.* (1986) reported that forb density, while not influenced by cattle grazing did increase with increasing tree thinning approximately 3 years after the experimental manipulation of original tree densities in the *E. crebra* woodlands of south-east Queensland. However, due to unequal variances caution must be applied when making assumptions relating to differences in growth-form richness between density treatments.

For the box woodlands, the results show that medium density patches are floristically similar to both low and high density patches. This result is different to that found previously where medium density patches were floristically similar to high density but dissimilar to low density (Goodhew 2005). This result may be attributable to site 11 which was shown to be more floristically similar to low density sites in ordination space than the other two medium density sites. Site 11 differs from the other medium density sites in that the native perennial grass *Bothriochloa decipiens* largely dominates the ground layer, similar to low density sites. *Bothriochloa decipiens* is recognised as a widespread grass species and reported to be very resistant to heavy grazing (Henry *et al.* 1995). However, a high proportion of *Bothriochloa decipiens* can indicate a drop in condition on more fertile soils. Therefore, the similarity between these treatments may be the result of a historically similar intensity of grazing.

After 2 years, the exclusion of large herbivores and livestock from grazing has resulted in little change in the understorey of woodland communities in the Traprock region. The patterns in floristic composition are associated with different mature tree densities and vegetation type, which are consistent with earlier findings (Goodhew 2005). It is suggested that a longer period of exclusion may result in changes in

floristic composition. Long-term monitoring of exclosure plots is essential to determining the biodiversity 'potential' of management units.

A long history of livestock grazing and tree removal in the region has contributed to significant differences in floristic composition between management units, yet this is not reflected in differences in total or native species richness between groups. As concluded in earlier research (Goodhew 2005), the heterogenous nature of vegetation management practices in the Traprock region maintains different communities of plant species and therefore, each management unit (or treatment combination) has value in contributing to regional plant diversity in the Traprock region.

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