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Vincent Allard, 2003

Effects of Elevated Atmospheric CO₂ Concentrations on Carbon and Nitrogen Fluxes in a Grazed Pasture

Effets de l’Elévation de la Concentration en CO₂ Atmosphérique sur les Flux de Carbone et d’Azote en Prairie Pâturée

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“Je suis le luxe des civilisations occidentales”

Guillaume Clémentine (1999), *Le Petit Malheureux*  
éditions “Le serpent à plumes”
Abstract

Predicting the response of grazed grasslands to elevated CO₂ is of central importance in global change research as grasslands represent 20% of the world's land area and grassland soils are a major sink for carbon (C). Grasslands responses to elevated CO₂ are strongly controlled by the availability of other nutrients and nitrogen (N) in particular. There have been many previous studies of N cycling in grasslands exposed to elevated CO₂ but none of these experiments were grazed. In this thesis I present data on CO₂ effects on N cycling from an experimental system (FACE: Free Air Carbon dioxide Enrichment) that enabled grazing to be included. The thesis focuses on the effects of elevated CO₂ on the different processes involved in organic matter (OM) returns from the plant to the soil and the consequences for N availability. In Chapter 1, it was shown that elevated CO₂ modified N returns by grazing animals by altering the partitioning of N between faeces and urine creating a potential for enhanced N losses at elevated CO₂. Plant litter decomposition rates were, at the ecosystem scale, not affected by elevated CO₂ (Chapter 3), but a marked increase in the organic matter fluxes, from roots, led to an accumulation of coarse OM in the soil (Chapter 4). In Chapter 5, using ¹⁴C and ¹⁵N labelling, I compared short-term (plant mediated) and long-term (soil mediated) effects of elevated CO₂ on soil OM dynamics and concluded that soil OM accumulation under elevated CO₂ was not caused by C or N limitation but probably by the availability of other nutrients. The thesis demonstrates that the inclusion of grazing animals can strongly modify N cycling under elevated CO₂. As most grasslands are grazed, the prediction of grassland responses to elevated CO₂ must be derived from systems in which animals are an integral part.
Résumé

Prédire la réponse des prairies pâturées à l’élévation de la concentration en CO₂ revêt une importance majeure dans la mesure où cet écosystème représente environ 20% de la surface terrestre non immergée mais aussi, parce que les sols prairiaux représentent un important puit de carbone (C). La réponse des prairies au CO₂ est fortement contrôlée par la disponibilité des autres nutriments et en particulier l’azote (N). De nombreuses expériences ont par le passé étudié le cycle de l’azote en prairie sous CO₂ enrichi mais aucunes de ces études n’a inclus le pâturage. Dans le cadre de cette thèse, je présente des données concernant les effets du CO₂ sur le cycle de l’N provenant d’un système expérimental (FACE: enrichissement en dioxyde de carbone à l’air libre) permettant d’inclure des ruminants. Cette thèse est dédiée à l’étude des effets de l’élévation en CO₂ sur les différents processus impliqués dans les retours de matière organique (MO) de la plante vers le sol et leurs conséquences pour la disponibilité en N. Dans le Chapitre 1, il a été montré que le CO₂ pouvait modifier les retours d’N par les ruminants en affectant la partition d’N entre l’urine et les fæces, ce qui induisait des pertes d’N potentiellement accrues. La décomposition de la litière végétale, considérée à l’échelle de l’écosystème, n’a pas été affectée par le CO₂ (Chapitre 3) mais une forte augmentation du volume de MO retournant au sol depuis les racines a induit une accumulation de MO grossière dans le sol (Chapitre 4). Au cours du Chapitre 5, à l’aide d’un double marquage isotopique ¹⁴C et ¹⁵N, nous avons comparé les effets court terme (transmis par la plante) et long terme (transmis par le sol) du CO₂ sur la dynamique de la MO du sol et il a été conclu que l’accumulation de MO n’était pas causée par une limitation en C ou en N mais probablement par la disponibilité des autres nutriments. Cette thèse démontre que les ruminants peuvent fortement modifier la réponse des prairies au CO₂. Dans la mesure où ce mode d’utilisation des pâtures est largement majoritaire, prédire les réponses des pâtures à un enrichissement en CO₂ doit provenir de systèmes où les ruminants sont partie intégrante.
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First, I would like to thank my four supervisors, Paul Newton, Cory Matthew, Jean-François Sousana and Philippe Grieu. All of them provided me help from the start of this work, made excellent facilities available to me both in New Zealand and France and provided ideas, comments and criticisms all through these three years.

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Structure of the thesis

This thesis is based on a series of papers. Chapter 2 has been accepted for publication in Global Change Biology. Chapters 3 and 4 have been prepared for submission in Global Change Biology and Plant and Soil respectively. A decision on submission of Chapter 5 for publication is pending. The references relevant to individual chapters are at the end of each chapter.
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1.1. Introduction

Since the beginning of the industrial revolution, the intensive use of fossil energy sources (coal, oil and natural gas) and the continuing conversion of forests into agricultural land, has profoundly affected the chemical composition of our atmosphere by simultaneously increasing the sources and decreasing the sinks of biospheric carbon (C). This has led to an exponential increase of the atmospheric CO₂ concentration from a pre-industrial value of \(280 \mu l \, l^{-1}\) to about \(373 \mu l \, l^{-1}\) today (Fig. 1.1; Keeling and Whorf, 2003). Should this trend continue, CO₂ concentrations by the end of the 21st century are expected to be between 540 and 970 \(\mu l \, l^{-1}\) (IPCC, 2001).

The increase in atmospheric CO₂ concentration has two major implications for the terrestrial ecosystems that should be considered separately. As a greenhouse gas, the rise in atmospheric CO₂ concentration could lead to an increase of global average air temperature and consequently strongly modify the functioning of the climate. It is estimated that surface temperatures have increased by \(0.6^\circ C\) since 1860 and that temperatures will be between \(0.6\) and \(6^\circ C\) warmer in 2100; this will have major implications for global and local climatic patterns, as well as for terrestrial ecosystem processes (IPCC, 2001). This thesis will not address the indirect effects of an increase in atmospheric CO₂ on air temperature and its numerous implications for plants but will focus on its direct effects on ecosystem functioning that occurs both at the single plant and community scales because CO₂ is the primary C substrate of photosynthesis.

In this context, the question of the response of grasslands to an increase in atmospheric CO₂ is of particular importance. Grasslands (temperate and savannas) account for more than 20% of the terrestrial surface and therefore predicting their response to elevated CO₂ has both major economic and ecological relevance. In addition, grasslands have a particular place in the total terrestrial C budget. Compared
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to tropical forests, the C contained in grassland biomass is relatively limited, but when soils are taken into account their importance is dramatically increased (Fig. 1.2). Grasslands soils have been shown to sequester large amount of atmospheric C (Kim et al., 1992; Franck et al., 2000), a phenomenon that could contribute to buffer the increase in atmospheric CO₂.

Both grassland plant response to elevated CO₂ and sequestration of atmospheric CO₂ may be controlled by the availability of other nutrients and nitrogen (N) in particular. This thesis focuses on the C-N interactions in a grazed temperate grassland with an emphasis on how organic matter returns may affect soil N availability and organic matter content. In the following literature review I present results from previous studies about the effects of elevated CO₂ on single plant growth and show the difficulty of extrapolating these results to the ecosystem scale. I then give a brief description of the experimental site used in this work and show the advantages of the techniques used for understanding ecosystem responses to elevated CO₂.

1.2. Plant response to elevated CO₂

Because CO₂ concentration at its current level limits the rate of photosynthesis and hence C inputs into plants, elevated CO₂ has numerous effects on plant physiology. In this first part, I discuss the direct effects of elevated CO₂ on plant photosynthesis and growth. Attention is placed on the numerous controls affecting plant growth responses to elevated CO₂.

1.2.1. Photosynthesis stimulation

Increased photosynthetic activity at the leaf level under elevated CO₂ has been observed in a wide range of plants; including trees such as Pinus sylvestris (Jach and Ceulemans, 2000) and Populus tremuloides (Curtis et al., 2000); crop species such as Triticum aestivum (Brooks et al., 2000) and Glycine max (Ziska et al., 2001); and
Figure 1.3: CO₂ dependency of photosynthesis of ryegrass (*Lolium perenne*) plants grown under ambient (full circles) or elevated (open circles) CO₂ for 3 months. The negative photosynthetic acclimation of plants grown under elevated CO₂ causes their lower assimilation rate.
pasture species like *Lolium perenne* (Rogers *et al.*, 1998) and *Trifolium repens* (Greer *et al.*, 2000).

As described by Farquhar (1980), C₃ photosynthesis, over the range of CO₂ concentrations occurring over a geological time scale (150-600 μl l⁻¹), is mainly limited by two phenomena; (i) the C fixation rate, which depends on the activity of ribulose 1,5-bisphosphate carboxylase-oxygenase (Rubisco) and (ii) on the regeneration of ribulose 1,5-bisphosphate (RuBP), which ultimately is limited by the availability of energy rich-compounds (ATP and NADPH). Under the current ambient CO₂ concentration Rubisco activity is unsaturated and therefore its activity increases with higher atmospheric levels of CO₂. In addition, because Rubisco has a double function (catalysing simultaneously RuBP carboxylation and oxygenation) the increase in the CO₂ concentration not only stimulates Rubisco activity but also limits photorespiration by displacing the competitive equilibrium between CO₂ and O₂ more towards carboxylation, hence reducing the amount of assimilated C immediately lost through this process.

The primary stimulation of photosynthetic rates under elevated CO₂ is partially buffered by a reduction of the photosynthetic capacity upon long-term exposure to high levels of CO₂ (Fig. 1.3), which is associated with reduced levels of Rubisco and organic N per unit leaf area (Long *et al.*, 1993). This down regulation or acclimation of photosynthesis usually increases with the duration of elevated CO₂ exposure and is most pronounced in plants grown under low nutrient supply (Sage, 1994). The causes of photosynthetic acclimation are both direct CO₂ control over photosynthesis and/or indirect CO₂ effects on plant N nutritional status (see review by Stitt and Krapp, 1999). Regardless of the main driver of acclimation and its physiological meaning, it is clear that the accumulation of non-structural carbohydrates usually observed under elevated CO₂ (see later) acts as a negative feedback on the synthesis of photosynthetic enzymes (Stitt, 1991).

Despite the acclimation phenomenon, plants growing under elevated CO₂ concentration usually maintain a higher photosynthetic rate per unit leaf area than plants growing under ambient CO₂ conditions, when they are studied in their respective growth environments. For example, recent data has shown that *Lolium*
perennial swards from a grass/legume mixture grown under elevated CO₂ for ten years sustained photosynthetic rates per unit leaf area about 40% higher than ambient plants, and with very little variation over the course of the experiment (Ainsworth et al., 2003). In order to determine whether this increase in C assimilation per unit leaf area under elevated CO₂ is totally or only partially translated into more biomass, it is necessary to discuss the possible physiological controls over photosynthesis under elevated CO₂.

1.2.2. Plant growth responses to elevated CO₂

Although photosynthesis and growth are intricately linked, it does not mean that the CO₂ dependence of plant growth can be simply derived from the CO₂ dependence of photosynthesis (Lloyd and Farquhar, 1996). In a compilation of 14 long-term ecosystem-scale CO₂ studies, Körner (1996) clearly pointed out there was a missing link between the CO₂-induced stimulation of leaf photosynthesis and the amount of C in the biomass. In this compilation, 13 of the 14 experiments exhibited a disproportional increase in photosynthesis compared to biomass production when exposed to elevated CO₂. In a calcareous grassland, Leadley and Körner (1996) observed an increase in CO₂ assimilation under elevated CO₂ but this resulted in no above- or belowground biomass stimulation. Using a modelling approach to study this phenomenon, Luo et al. (1997) predicted that the observed increase of 70% in leaf photosynthesis in a Californian grassland community grown under elevated CO₂ should translate into an increase of 97% in plant biomass if no physiological adjustments in the plants occurred. This certainly did not match the observed 5, 13 and 40% biomass increases during the 3 years of the experiment.

The most likely reason for the discrepancy between growth and photosynthesis enhancement is that plants exhibit increased photosynthesis to the extent that they cannot fully incorporate the newly fixed C into additional growth. Firstly any supplementary photosynthetic C acquisition must be accompanied by increased maintenance respiratory costs (Lloyd and Farquhar, 2000). Secondly, plants grown under elevated CO₂ tend to accumulate non-structural carbohydrates in the leaves (Stitt, 1991; Casella and Soussana, 1997; Poorter et al., 1997), usually taken as a sign
Figure 1.4: Plant response to elevated CO$_2$; distribution of the weight ratio (plant biomass$_{elevated}$/plant biomass$_{ambient}$). Data from Poorter (1993) after compilation of the literature (total number of species is 156). The vertical dotted line indicates a weight ratio of 1, i.e. no biomass response to elevated CO$_2$. 
that the plant is unable to convert all the newly fixed C into structural biomass. Thirdly, due to this accumulation of non-structural carbohydrates, a decrease in plant specific leaf area (SLA) is one of the major adjustments found upon prolonged exposure to elevated CO2 (Lambers et al., 1998). By having more biomass in a given leaf area, the photosynthetic increase is partially buffered by having less light capture per unit biomass. Consequently growth stimulation is less than the photosynthetic stimulation per unit area (Evans and Poorter, 2001).

Despite these adjustments, single plant growth is usually greater under elevated CO2. Poorter (1993), in a compilation of literature sources based on 156 different species found an average 37% stimulation of plant biomass (Fig. 1.4). Poorter also identified some functional traits, strongly linked with plant strategy, which were correlated with plant response to elevated CO2. For example, species with a high relative growth rate (i.e. fast-growing species) were found to exhibit a higher response to elevated CO2. This was supported by other studies also based on plant strategies (Hunt et al., 1991). More generally, the plant response to elevated CO2 might depend on the capacity to functionally use the newly fixed C (i.e. plants with strong C sinks) through: (i) greater accumulation of C in storage organs (Diaz, 1995), (ii) increased C allocation towards symbiotic microorganisms, for example mycorrhizal infection (Rillig et al., 1998), or (iii) enhanced allocation belowground through higher root growth, turnover or exudation (Paterson et al., 1996). Based on this information, it is intuitive that the fate of the extra C fixed under elevated CO2 not only depends on a plants inherent capacity to use it, but also on the availability of other nutrients.

1.2.3. Nutrient dependency of plant response to elevated CO2

CO2 is just one of the many inorganic substrates that are required by plants to grow. Their response to elevated CO2 will thus depend on the availability of other nutrients and the way they are utilised by the plant. Bazzaz (1990) stated that nutrient-limited plants respond less to increases in atmospheric CO2 than plants grown at high nutrient availability. Under elevated CO2, the higher rates of growth create a higher demand for other nutrients relative to C; these become relatively more limiting for
plant growth. In an experiment assessing the CO₂ response of a grass (*L. perenne*) under different soil nitrogen regimes, the CO₂-induced photosynthetic response was 3-fold greater with high N availability (Casella and Soussana, 1997). More generally, meta-analyses compiling numerous studies to assess the interactions between CO₂ and nutrient availability tend to show that on average, whole plant biomass stimulation under elevated CO₂ is less at low nutrient levels (Poorter, 1998; Poorter and Pérez-Soba, 2001). One reason for this positive relationship between CO₂ and N availability relies on the fact that, regardless of the elevated CO₂ issue, low-nutrient plants consistently show an accumulation of non-structural carbohydrates in the leaves (Chapin, 1980). This might therefore accentuate the photosynthesis acclimation phenomenon, which is partially induced by non-structural carbohydrates. Photosynthetic acclimation has indeed often been seen to be stronger under low N supply (Sage, 1994; Bowler and Press, 1996).

Nevertheless, plant C and N metabolic processes are in close interrelation and therefore there are numerous feedbacks between them (Stitt and Krapp, 1999). In particular assuming that an increased C availability simply increases the relative need for N or other nutrients would be over-simplistic, because elevated CO₂ might also affect the use of the existing nutrients. For example, a classic outcome of CO₂ studies is increased N use efficiency (NUE) i.e. the rate of growth per unit N taken up increases under elevated CO₂ (Davey *et al.*, 1999; Midgley *et al.*, 1999; Osborne *et al.*, 1998). Because under elevated CO₂ a given rate of C assimilation can be achieved with lower activities of Rubisco or other photosynthetic enzymes (Stitt, 1991; Rogers *et al.*, 1996), this allows the reinvestment of photosynthetic N into more limiting processes (Stitt and Krapp, 1999). When these complex physiological controls between C and N metabolisms are taken into account, the relationship between plant CO₂ response and N supply is not clear-cut and this may explain the results of certain studies where there were higher responses to CO₂ under low N supply (see for example Wong *et al.* 1992; Hocking and Meyer, 1991). Lloyd and Farquhar (1996), using an approach based on photosynthetic models, predicted that there was no simple relationship between N and CO₂ because of the variability of plant nutritional status and/or strategy of nutrient economy.
The issue of N control over the plant response to CO2 assumes another dimension when the possible effects of CO2 on soil N availability are taken into account: observed effects have been both positive (Zak et al., 1993) and negative (Diaz et al., 1993). This highlights two important needs in elevated CO2 studies: the necessity of whole ecosystem studies to take into account possible feedbacks through soil processes, and the need for long-term experiments in order to allow these feedbacks to occur.

1.3. The difficulty in predicting ecosystem responses to elevated CO2

When studying the effects of elevated CO2 on complex ecosystems such as grasslands, interspecific competition and resource limitation become important (Körner, 1996). Prediction at the ecosystem scale based upon extrapolation of data obtained from a simpler system (typically at the single plant scale) has potential pitfalls. In particular it is important to consider spatial variability and possible time issues as these can profoundly alter system responses to elevated CO2.

1.3.1. CO2 effects in multispecific systems

A good example of the difficulty in extrapolating community responses to elevated CO2 from single plant studies is the question of the relative response to elevated CO2 of C4 plants compared to C3 plants. Early studies of this issue appeared to confirm the hypothesis that C4 plants should not show significant growth responses to elevated CO2 (Curtis et al., 1989), due to the CO2-concentrating mechanism in their bundle sheath cells. This mechanism increases the effective concentration of CO2 at the carboxylation site, thereby masking photorespiration and apparently assuring the saturation of Rubisco at ambient CO2 concentration. Therefore C4 plants should not benefit from increased CO2 concentration and may suffer reduced competitive abilities over C3 species. However, in a meta-analysis of the response of C4 and C3 grass species to elevated CO2, Wand et al. (1999) showed that this issue was not as clear-cut as previously thought. Both C4 and C3 species increased total biomass under elevated CO2 by 33% and 44% respectively. When the two types of plants are in
competition, the problem of the relative response of C\textsubscript{3} and C\textsubscript{4} plants can even be more complex. An experiment comparing the response to elevated CO\textsubscript{2} of cotton (*Gossypium hirsutum*, C\textsubscript{3}) and sorghum (*Sorghum bicolor*, C\textsubscript{4}) showed that even though the C\textsubscript{3} plant had a greater response to CO\textsubscript{2} in monoculture, the C\textsubscript{4} was still a better competitor in the mixture (Derner *et al.*, 2003). Similar outcomes can be found in non-arable ecosystems: after 8 years of enrichment in a tallgrass prairie ecosystem, Owensby *et al.* (1999) observed little modification of the C\textsubscript{4}-C\textsubscript{3} balance. This is particularly true under some sort of environmental stress situation (e.g. water limitation) in which case C\textsubscript{4} plants tend to maintain their CO\textsubscript{2}-induced stimulation to a greater extent than C\textsubscript{3} plants. Predicting the response of multispecific systems to elevated CO\textsubscript{2} requires more than simple compilation of single-species studies and therefore, species diversity must be included within any experiments designed to obtain valid results on community scale responses to elevated CO\textsubscript{2}.

In temperate grasslands, one of the strongest trends emerging from multispecific experiments concerns legume abundance. Numerous studies in a range of conditions have measured an increased legume abundance under elevated CO\textsubscript{2}: e.g. permanent grasslands (Clark *et al.*, 1997; Teyssonneyre, 2002; Edwards *et al.*, 2001a); or *Lolium perenne* – *Trifolium repens* mixture (Lüscher *et al.*, 1996; Hebeisen *et al.*, 1997). The increased competitive ability of the legumes is generally attributed to their capacity to fix atmospheric N that allows them to meet the enhanced requirements for N induced by the CO\textsubscript{2} stimulation effect (Lusher *et al.*, 2000). The CO\textsubscript{2}-driven increase in legume abundance under elevated CO\textsubscript{2} is nevertheless not ubiquitous (Navas *et al.*, 1995; Leadley and Körner, 1996); this may be explained to some extent by phosphorus limitation, a nutrient that is often underestimated as a key driver of the legume CO\textsubscript{2} response (Stöcklin and Körner, 1999; Körner, 2000).

From the literature it appears that even two strongly deterministic physiological traits that are intuitively of primary importance in the plant CO\textsubscript{2}-response process (photosynthetic type and N\textsubscript{2}-fixing ability), cannot be used as universal predictors of the CO\textsubscript{2} effects at the community level. The difficulty in predicting community response to elevated CO\textsubscript{2} is even greater when temporal aspects are taken into account.
1.3.2. *Duration of the CO₂ enrichment*

Some of the processes discussed above (e.g. the acclimation phenomenon or the discrepancy between photosynthesis and biomass production responses to elevated \( \text{CO}_2 \)) intuitively lead to the problem of *time* in elevated \( \text{CO}_2 \) experiments. \( \text{CO}_2 \) affects photosynthesis within seconds after \( \text{CO}_2 \) enrichment (the time for stomatal \( \text{CO}_2 \) concentration to adjust with the atmospheric concentration), while acclimation only occurs within days since it is (partly) controlled by an accumulation of non-structural carbohydrates in the leaves. The time problem is even more acute at higher levels of spatial complexity, increasing the number of possible feedbacks. Therefore the duration of the \( \text{CO}_2 \) enrichment applied to a given community can have a central importance when predicting its future functioning.

A valuable resource for assessing the differences between long-term and short-term plant responses to elevated \( \text{CO}_2 \) are natural \( \text{CO}_2 \) springs (Raschi *et al.*, 1997). They provide situations where both plants and soil have been exposed for long periods of time to high levels of \( \text{CO}_2 \) and therefore the system is likely to have adapted to this environment and to have reached a steady state. If such naturally \( \text{CO}_2 \)-enriched sites have nearby a standard site with similar characteristics and ambient \( \text{CO}_2 \) concentration it is possible to compare short- and long-term responses to elevated \( \text{CO}_2 \). Using such a natural \( \text{CO}_2 \) source and a cross-over experiment (i.e. transplanting soil cores from enriched locations to ambient locations and *vice versa*), Newton *et al.* (2001a) clearly showed the discrepancy existing between transient and equilibrium responses to elevated \( \text{CO}_2 \), particularly because of the long-term alterations of soil substrate and/or decomposer communities.

Similarly, changes in botanical composition under elevated \( \text{CO}_2 \) may require the integration of numerous phenomena that can be affected by \( \text{CO}_2 \) at different time scales. As discussed previously, legume proportion usually increases in temperate grasslands under elevated \( \text{CO}_2 \). But long-term changes in community botanical composition not only occur through vegetative process but also through difference in the response of reproductive mechanisms to elevated \( \text{CO}_2 \). Edwards *et al.* (2001a, 2001b) clearly showed that seed production and seedling recruitment and performance were all affected by elevated \( \text{CO}_2 \) in a temperate grassland and that this effect was
In particular, under elevated CO2, *Trifolium repens*, a species exhibiting a strong response to elevated CO2 in this particular system exhibited the following effects: (i) *T. repens* produced seeds with higher germination %, (ii) the resulting seedlings had a greater mass and (iii) seedlings grown from elevated CO2 seeds had a higher survival rate when sown into the pasture. Consequently, changes in species composition not only reflect the effects of elevated CO2 on the abundance of existing plants through vegetative competition, but also CO2 effects on the recruitment of new plants from seeds. Because these processes do not affect the community at the same time scale and because they are inter-dependent, excluding one of them in a study by extrapolating short-term results to long-term community response may be seriously misleading.

1.3.3. The step increase issue

A second aspect of elevated CO2 studies dealing with time relies on the methods used to simulate an increase in elevated CO2, in particular the potential initial artefact caused by a step increase in CO2. In the majority of CO2 experiments, a single concentration step CO2 increase is used to simulate future atmospheric CO2 concentration. However, plants in the “real world” are not, and will not, be exposed to such an abrupt increase in CO2 but rather to a gradually rising concentration at a rate of about 1.5 \( \mu \text{mol mol}^{-1} \text{ year}^{-1} \) (IPCC, 2001). In response to a step increase in CO2, photosynthetic rates usually dramatically increase (see above). The resulting large increment in photosynthetic C influx may exert different effects on the physiological processes compared to small increments in C influx observed with a gradual CO2 increase (Luo, 2001). In an 80 day experiment comparing the effects of a step increase of elevated CO2 (700 \( \mu \text{mol mol}^{-1} \)) with a gradual increase of 5 \( \mu \text{mol mol}^{-1} \text{ day}^{-1} \) (reaching 700 \( \mu \text{mol mol}^{-1} \) by the end of the experiment) on *Plantago lanceolata* photosynthesis and growth, Hui et al. (2002) showed that the step CO2 increase resulted in an immediately higher leaf photosynthesis rate and induced a large N demand and stress that led to considerable down-regulation of photosynthesis. In comparison, the gradual increases stimulated photosynthesis gradually and created limited N stress. Nevertheless growth response under the two CO2 treatments seemed
to converge at the end of the experiment showing a possible attenuation of the step-increase artefact with time.

Another aspect of the artefact created by a step-increase in elevated CO₂ relies on the fact that ecosystem responses to elevated CO₂ may depend on the concentration of the “elevated CO₂ treatment”. Indeed the response of some measured variables can be non-linearly related (i.e. not proportional) with the simulated increase of the CO₂ concentration. A good illustration of this phenomenon is given by studies using more than two CO₂ treatments (ambient and enriched), (e.g. Hunt et al., 1991; Sims et al., 1998, Clark et al., 1997; Granados and Körner, 2002); or a gradient of concentrations from sub-ambient to super-ambient CO₂ concentrations (Gill et al., 2002). For example Gill et al. (2002) showed that organic matter accumulation in a grassland soil was more sensitive to an increase from sub-ambient to ambient CO₂ concentration than to an increase from ambient to super-ambient CO₂ concentration. This implies the existence of specific thresholds that influence the magnitude of the change in some ecosystem processes. Granados and Körner (2002) also showed that the relative stimulation of growth by elevated CO₂ was larger at low as compared to higher ranges of CO₂ concentration. But in their experiment this phenomenon was species-specific with one species in particular expressing a negative effect of CO₂ at the highest CO₂ concentration used.

Predictive studies intrinsically require anticipation of future CO₂ conditions and therefore some sort of step-increase artefact is inevitable. But the examples given above clearly show that short-term plant responses to a step increase in CO₂ are far from being equilibrium responses. Therefore long-term studies should be preferred because they might allow systems to reach equilibrium, or at least show a reliable trajectory of response.

1.3.4. The need for large spatio-temporal studies to assess C-N interactions in grasslands under elevated CO₂

An important aspect of ecosystems response to elevated CO₂ dealing with both space and time are the numerous interactions existing between the C and N cycles.
We have seen so far that N is an important driver of both single plant response and community response to elevated CO$_2$ since the relative availability of C and N could induce feedbacks on photosynthesis at the plant scale but also shifts in botanical composition. The question of feedbacks in elevated CO$_2$ studies has already been described previously but lets discuss more exhaustively how the CO$_2$-induced changes occurring at these different scales may induce feedback through soil N availability.

It has long been recognised that soil N availability could be a key driver of long-term plant responses to elevated CO$_2$ through increased immobilisation (Diaz et al., 1993) or mineralisation (Zak et al., 1993) of soil N under elevated CO$_2$. Various organic matter fluxes in the soil may alter organic matter dynamics under elevated CO$_2$: decreased quality of leaf litter (Strain and Bazzaz, 1983; Cotrufo and Ineson, 1996) and increased biomass allocation to roots (Cotrufo and Gorissen, 1997), both leading to lower decomposition rates and subsequently limit N availability for plant growth. In addition, increased root exudation of readily decomposable C (Paterson et al., 1996) might stimulate the growth of soil microorganisms leading to higher N immobilisation. These processes will be reviewed more comprehensively later in this thesis but I will now set out why their existence requires a high level of integration.

Firstly, as implied by many of the concepts discussed above, N cycling can be affected by elevated CO$_2$ both at the plant scale through a decrease in leaf N concentration (Cotrufo et al., 1998), or at the community scale through an alteration of botanical composition. Secondly, soil N mineralisation not only depends on organic matter deposition through plant senescence (i.e. processes that can be affected in the relative short-term by elevated CO$_2$) but also by soil micro- and meso-fauna activity. Because soil organisms are further down in the chain of reactions through which elevated CO$_2$ affects ecosystem functioning, enough time is needed for them to reach a new equilibrium and for possible feedbacks to occur.
Table 1.1: Major elevated CO₂ experiments in grasslands.

<table>
<thead>
<tr>
<th>Location</th>
<th>Vegetation type</th>
<th>Reference</th>
<th>Facility</th>
<th>[CO₂]</th>
<th>Experimental duration</th>
<th>Past management</th>
<th>Defoliation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colorado, USA</td>
<td>Shorthgrass steppe (C3-C4)</td>
<td>Morgan <em>et al.</em>, 2001</td>
<td>OTC</td>
<td>Amb-720</td>
<td>About 3 years</td>
<td>Low to moderate grazing by cattle</td>
<td>2 cuts per year</td>
</tr>
<tr>
<td>Kansas, USA</td>
<td>Tallgrass prairie (C3-C4)</td>
<td>Owensby <em>et al.</em>, 1999</td>
<td>OTC</td>
<td>Amb-twice amb</td>
<td>8 years</td>
<td>Winter grazing by cattle</td>
<td>1 cut per year</td>
</tr>
<tr>
<td>Texas, USA</td>
<td>Grassland (C3-C4)</td>
<td>Polley <em>et al.</em>, 2002</td>
<td>Tunnels</td>
<td>200-550 gradient</td>
<td>About 3 years</td>
<td>Grazed grassland</td>
<td>1 cut per year</td>
</tr>
<tr>
<td>California, USA</td>
<td>Annual grassland</td>
<td>Hungate <em>et al.</em>, 1997</td>
<td>OTC</td>
<td>Amb-Amb+350</td>
<td>4 years</td>
<td>n.a.</td>
<td>Single regrowth</td>
</tr>
<tr>
<td>California, USA</td>
<td>Sown annual grassland</td>
<td>Cardon <em>et al.</em>, 2001</td>
<td>OTC</td>
<td>Amb-Amb+350</td>
<td>Various</td>
<td>New</td>
<td>Single regrowth</td>
</tr>
<tr>
<td>California, USA</td>
<td>Annual grassland</td>
<td>Shaw <em>et al.</em>, 2002</td>
<td>Mini-Face</td>
<td>Amb-680</td>
<td>Since 1998</td>
<td>n.a.</td>
<td>Single regrowth</td>
</tr>
<tr>
<td>Minnesota, USA</td>
<td>Sown perennial grassland</td>
<td>Reich <em>et al.</em>, 2001</td>
<td>FACE</td>
<td>Amb-550</td>
<td>Various</td>
<td>New</td>
<td></td>
</tr>
<tr>
<td>Clermont-Fd, France</td>
<td>Sown mixtures</td>
<td>Teyssonneyre <em>et al.</em>, 2002</td>
<td>Glasshouse</td>
<td>Amb-700</td>
<td>1 year</td>
<td>New</td>
<td>2 treatments : 3 or 6 cuts per year</td>
</tr>
<tr>
<td>Clermont-Fd, France</td>
<td>Sown ryegrass</td>
<td>Soussana <em>et al.</em>, 1996</td>
<td>Glasshouse</td>
<td>Amb-700</td>
<td>2 years</td>
<td>New</td>
<td>6 cuts per year</td>
</tr>
<tr>
<td>Clermont-Fd, France</td>
<td>Perennial grassland monoliths</td>
<td>Teyssonneyre, 2002</td>
<td>Mini-Face</td>
<td>Amb-600</td>
<td>About 4 years</td>
<td>Grazing by sheep</td>
<td>2 treatments : 4 or 8 cuts per year</td>
</tr>
<tr>
<td>Basel, Switzerland</td>
<td>Calcareous grassland</td>
<td>Niklaus <em>et al.</em>, 2001</td>
<td>SACC (OTC)</td>
<td>Amb-600</td>
<td>5 years</td>
<td>Extensively grazing by cattle</td>
<td>2 cuts per year</td>
</tr>
<tr>
<td>Basel, Switzerland</td>
<td>Calcareous grassland monoliths</td>
<td>Stöcklin <em>et al.</em>, 1998</td>
<td>Glasshouse</td>
<td>Amb-600</td>
<td>2 years</td>
<td>Extensively grazing by cattle</td>
<td>2 cuts per year</td>
</tr>
<tr>
<td>Basel, Switzerland</td>
<td>Sown simulated grasslands</td>
<td>Stöcklin <em>et al.</em>, 1999</td>
<td>Glasshouse</td>
<td>Amb-600</td>
<td>2 years</td>
<td>New</td>
<td>2 cuts per year</td>
</tr>
<tr>
<td>Zurich, Switzerland</td>
<td>Lolium-Trifolium mixture</td>
<td>Glöser <em>et al.</em>, 2000</td>
<td>FACE</td>
<td>Amb-600</td>
<td>8 years</td>
<td>New</td>
<td>2 cuts per year</td>
</tr>
<tr>
<td>Bulls, New Zealand</td>
<td>Perennial pasture turves</td>
<td>Newton <em>et al.</em>, 1994</td>
<td>Growth chamber</td>
<td>350-700</td>
<td>217 days</td>
<td>Grazed grassland</td>
<td>Cut every 3 weeks</td>
</tr>
<tr>
<td>Bulls, New Zealand</td>
<td>Perennial pasture turves</td>
<td>Clark <em>et al.</em>, 1997</td>
<td>Growth chamber</td>
<td>350-525-700</td>
<td>324 days</td>
<td>Grazed grassland</td>
<td>2 treatments: every 3 or 6 weeks</td>
</tr>
<tr>
<td>Bulls, New Zealand</td>
<td>Perennial grassland</td>
<td>Edwards <em>et al.</em>, 2001a</td>
<td>FACE</td>
<td>Amb-475</td>
<td>Since 1997</td>
<td>Grazed grassland</td>
<td>Grazed</td>
</tr>
</tbody>
</table>
1.4. The New Zealand grazed pasture FACE experiment

1.4.1. Free air CO₂ enrichment (FACE) experiments

In order to study the effects of elevated CO₂ on communities it is therefore essential to take into account the issue of time and scale discussed above. Free Air CO₂ Enrichment (FACE) experiments were developed to meet this need. This is a technology that delivers consistent CO₂ concentration, to large plots of an intact ecosystem without walls, and therefore without altering the microclimate or disturbing the soil (Allen, 1992; Hendrey et al., 1993). In particular FACE does not modify environmental factors such as incident solar radiation, temperature, humidity and wind compared with methods that grow plants under elevated CO₂ in some sort of enclosure (i.e. glasshouses, tunnels and open-top chambers (OTC)). FACE systems release CO₂ to area of vegetation ranging from 1 to 27 m in diameter. Various technologies are used but all the systems control the CO₂ concentration in the target area using feedbacks that take into account of wind speed and direction and the CO₂ concentration (Plate 1.1). Enrichment may be to a set point or as a percentage addition to the ambient level (McLeod and Long, 1999). The FACE technique has been successfully used to grow a variety of vegetation types under elevated CO₂ (Plate 1.2): including rice (Okada et al., 2001), pastures (Hebeisen et al., 1997) and trees (Matamala and Schlesinger, 2000).

1.4.2. Specifications of the New Zealand FACE

Worldwide there have been numerous FACE experiments on grassland, which have together with smaller scale studies (Table 1.1) produced useful information at scales ranging from plant physiology to ecosystem ecology under elevated CO₂. Nevertheless none of these studies, often for practical limitations, have used grazing animals to achieve defoliation, whereas the majority of grassland ecosystems worldwide are grazed, at least to some extent, by wild or domestic animals. Table 1.1 highlights the fact that all existing experiment studying the effect of elevated CO₂ on grassland excluded grazing even when it was part of the normal management of the ecosystem.
Plate 1.1: General view of an elevated CO\textsubscript{2} ring at the New Zealand pasture FACE. Depending on the wind direction measured in the centre of the ring, CO\textsubscript{2} enriched air is emitted by uprights encircling the ring that are in the wind. The CO\textsubscript{2} concentration is monitored by a CO\textsubscript{2} sensor at the centre of the ring.

Plate 1.2: FACE systems have been designed to simulate CO\textsubscript{2} enrichment on various ecosystems including (a) an Aspen forest (Wisconsin, USA), (b) a loblolly pine forest (North Carolina, USA), (c) a Mojave desert ecosystem (Nevada, USA) and (d) crops such as rice (Shizukuishi, Japan).
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The New Zealand FACE experiment starts from the premise that the inclusion of grazers and ruminants in grassland studies may be of central importance for a number reasons. First, concerning CO₂ effects on pasture botanical composition, the homogeneous removal of the vegetation by cutting does not take into account possible selective grazing by ruminants (Parsons et al., 1994). Such a supplementary pressure on the selected species might be of importance in determining community response. More complex interactions between CO₂ and management (cutting or grazing) may also occur (Newton et al., 2001b). Second, regarding the important question of N cycling under elevated CO₂, grazed grassland will differ significantly from cut grasslands both in terms of the amount of nutrient and the heterogeneity of these returns (Haynes and Williams, 1993), leading to the likelihood that these systems will differ in their response to elevated CO₂. Because, at a global scale, most of the grasslands are at least partially managed with domestic livestock, the prediction of grassland response to elevated CO₂ must at some stage include ruminants or other grazers in the scope of study. Compared to other grassland FACE experiments, the inclusion of grazing by sheep within the normal experimental management is the main distinguishing feature of the New Zealand pasture FACE.

1.4.3. Description of the facility

The New Zealand pasture FACE was set up in a pasture at Bulls, Manawatu, New Zealand (40°14'S, 175°16'E). The pasture had been under permanent grazing since the 1940s by cattle, sheep and goats with occasional hay cuts taken. An exhaustive botanical composition determination of the pasture in spring 1996 found 25 vascular plant species including the C₃ grasses Agrostis capillaris L., Anthoxanthum odoratum L., Lolium perenne L., the C₄ grasses Paspalum dilatatum Poir. and Cynodon dactylon (L.) Pers., the legumes Trifolium repens L., Trifolium subterraneum L. and Trifolium dubium L. and the herbs Hypochaeris radicata L. and Leontodon saxatilis Lam syn. L. taraxacoides. All the species found were non native. The soil at the site is a Pukepuke black sand (Mollic Psamment) with a 0.25-m black loamy, fine sand top horizon underlain by greyish-brown, fine sand textured
Figure 1.5: Twenty years temperature and rainfall average at the experimental site recorded from a nearby weather station (Flock House, Bulls). Average monthly temperature is indicated by the full line and maximum and minimum monthly average are also represented. Average monthly rainfall is indicated by the full bars.
horizons. Mean values of some physico-chemical characteristics of this soil in 1997 were: pH = 5.8, K = 0.15 cmol c kg soil⁻¹, P (phosphate) = 20 μg ml soil⁻¹ and S (sulphate) = 7 μg ml soil⁻¹ (data previously compiled by Edwards et al., 2001b). Mean rainfall and air temperature at the experimental site are shown in Fig. 1.5.

The FACE facility consists of six rings, each 12 m in diameter. Three blocks of two rings were grouped based on initial botanical composition and soil properties. One ring in each block was selected to be enriched with CO₂, the other being left at ambient CO₂ concentration. Each ring was fenced off in order to control sheep access and the duration of the grazing events. The FACE system was installed during the first nine months of 1997 and enrichment began on 1 October 1997. The target CO₂ concentration for elevated CO₂ plots is 475 μl l⁻¹ CO₂ during the photoperiod. Enrichment takes place through 24 standpipes, each 0.05 m in diameter, located on the perimeter of each ring. Enriched air is blown across the ring from the up wind direction. The rings are intermittently grazed by sheep. Grazing begins when aboveground herbage biomass reach 1.8-2 t DM ha⁻¹ and continues until the residual biomass is about 0.5-0.7 t DM ha⁻¹. Standing dry matter and botanical composition of the rings is measured before and after each grazing event.

1.4.4. Objectives of the work

As discussed above, N cycling in grasslands submitted to elevated CO₂ is central to predict the response of this ecosystem to our changing climate. Elucidation of this forms the core objective of this thesis which is based on four separate experiments.

- As discussed previously, elevated CO₂ affects pasture both at the plant and the community scale. With regard to the question of N cycling, the decrease in leaf N concentration usually observed at the single plant scale under elevated CO₂ is accompanied, at the community scale, by an increase in legume proportion, possibly counterbalancing the leaf N concentration decrease. Because ruminants, through grazing, integrate both these effects of elevated CO₂ on pasture N status I
hypothesised that N returns by the ruminants grazing in a pasture under elevated CO₂ might be altered.

- A second pathway of organic matter and N return to the soil is litter decomposition. Though it is usually accepted that litter decomposition per se is not affected by elevated CO₂, there is still debate over the possibility that indirect effects of elevated CO₂ on botanical composition and shoot/root biomass partitioning might alter decomposition rates at the ecosystem scale. Therefore I compared in situ decomposition rates of leaf litter from different species, root material and faeces in order to integrate the different levels at which elevated CO₂ may act.

- The effects of elevated CO₂ on litter decomposition rates are not the only drivers of possible alterations of N availability and soil organic matter accumulation. The size of the litter fluxes returning to the soil is also of importance. Therefore I assessed the question of the quantity of plant material returning to the soil under elevated CO₂ and in particular, attention was paid to root growth and turnover, processes that are expected to be strongly enhanced under elevated CO₂. I hypothesised that if organic matter flux size was greater under elevated CO₂, it may lead to a detectable accumulation of young organic matter in the soil under elevated CO₂ and to a subsequent immobilisation of mineral N and accumulation of C in the soil.

- In order to test whether long-term feedbacks through the possible CO₂-driven changes in soil organic matter could affect plant response to elevated CO₂ a glasshouse experiment was set up. Soil from ambient and enriched rings were used in a cross-over design under ambient or enriched atmospheric CO₂ concentration and the growth and N yield of a model species, *Lolium perenne* was measured.

Finally, in a general discussion, some questions raised by the previous result chapters are developed and some perspectives about this work are discussed.
1.5. References


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pasture turves exposed to two levels of elevated CO$_2$. *Journal of Applied Ecology*, **34**, 304-316.


Chapter 1


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Plate 2.1: General view of an elevated CO$_2$ FACE ring during grazing by adult sheep wearing faeces collection bags.

Chapter 2: Nitrogen Cycling in Grazed Pastures at Elevated CO$_2$: N Returns by Ruminants
Chapter 2: Nitrogen Cycling in Grazed Pastures at Elevated CO₂: N Returns by Ruminants

Allard V, Newton PCD, Lieffering M, Clark H, Matthew C, Soussana JF, Gray YS.

2.1. Abstract

In pastures grazed by large herbivores, nutrients cycle both through litter and animal excreta. We compared nitrogen (N) returns from sheep grazing a temperate pasture exposed to ambient or elevated CO₂ (475 µl l⁻¹) in a FACE (Free Air CO₂ Enrichment) experiment established in the spring of 1997. In the spring of 2000 and 2001 we measured the chemical composition of the diet, sheep faeces and of individual plant species before grazing to characterise feed intake and to compare the intake of N to the N produced in faeces. In both years under elevated CO₂, leaves of the individual species exhibited lower N concentrations and higher water soluble carbohydrate (WSC) concentrations. There was a significantly greater proportion of legume in the diet at elevated CO₂ but, together with the changes in chemical composition of individual species, this resulted in diets that had similar N but higher WSC and digestibility for both ambient and elevated CO₂. We calculated that a greater proportion of dietary N was partitioned to urine at elevated CO₂, probably because of the higher proportion of legume N in the diet, with possible differences in protein quality. A potentially significant consequence of this change in partitioning is greater N loss through volatilisation at higher CO₂ levels.
2.2. Introduction

Grasslands cover a fifth of the terrestrial surface of the world (Hadley 1993) and most of this area is grazed by domestic livestock. The response of these grasslands to the predicted continuing increase in atmospheric CO₂ (Keeling et al., 1995) is potentially important, first because of their economic significance and, second, because grasslands soils are a major sink for carbon (C) (Thornley et al., 1995). The response of grasslands to elevated CO₂ depends, in the long-term, partly on how elevated CO₂ modifies nutrient cycling. It has been proposed that the fertilisation effect of elevated CO₂ on plant communities (Newton 1991) could be constrained by a relative decrease in the availability of other nutrients, in particular nitrogen (N) (Diaz et al., 1993). One of the major mechanisms implicated in this negative feedback is a reduction in litter quality (i.e. increased C/N ratio) resulting from reduced N concentrations in plant tissues at elevated CO₂ (Poorter et al., 1997; Cotrufo et al., 1998) and, therefore, reduced N mineralisation rates in soils (Strain and Bazzaz 1983; Norby et al., 1986; van Ginkel and Gorissen 1998). However, it has become apparent that the observed decrease in N concentration in green leaves at elevated CO₂ (Cotrufo et al., 1998) is not, or is only partially, reflected in the senescing tissues; consequently only small changes in litter quality (Hirshel et al., 1997; Hartwig et al., 2000; Norby et al., 2001) and decomposition rates (Dilustro et al., 2001; Van Vuuren et al., 2000) appear as a result of elevated CO₂. Thus, in ecosystems in which nutrient cycling depends heavily on shoot litter decomposition it seems unlikely that any feedback on N availability will occur (Norby and Cotrufo 1998).

However, in grasslands grazed by large herbivores only part of the nutrient cycling occurs through litter decomposition and this fraction decreases with higher herbage utilisation rates. In a well managed temperate grassland the optimum herbage utilization is about 50% of aboveground biomass production (Parsons and Chapman 2000). In this situation, due to the lower N concentration of senescing leaf material compared to green herbage and the poor N utilization by ruminants (Jarvis 2000), about 75% of the N returned to the soil occurs through the dung and urine of the grazers. In terms of potential CO₂ effects on these N cycling pathways it is important to note that grazers eat primarily live plant material and, therefore, any CO₂-induced
changes in green tissue composition may influence nutrient returns. In contrast, the litter pathway is sensitive to the chemical composition of dead/senescent material.

The literature indicates that changes in the chemical composition of ruminant diet at elevated CO₂ may arise from two sources: 1) a change in the chemical composition of individual species, especially the well documented reduction in N concentration of leaves (Poorter et al., 1997; Cotrufo et al., 1998), and 2) a change in plant species composition in the pasture, in particular, a shift to a higher content of N-rich legumes (Hebeisen et al., 1997; Jongen and Jones 1998; Teyssonneyre et al., 2002). Through the grazing process, grazers integrate the CO₂ effects that occur at these different scales (plant and community levels). Although most grasslands are grazed, our current understanding of nutrient cycling at elevated CO₂ is based exclusively on cut grasslands i.e. in which animals are excluded. There is good reason to expect that cycling under these conditions will differ significantly from grazed grasslands both in terms of the amount of nutrients returned and the heterogeneity of these returns (Haynes and Williams 1993) leading to the likelihood of cut and grazed systems responding very differently to elevated CO₂ (Newton et al., 2001). In this paper we present data on N returns by sheep grazing a pasture exposed to elevated CO₂.

### 2.3. Materials and methods

#### 2.3.1. Experimental site

The study was carried out in November 2000 and November 2001 (spring in the Southern Hemisphere) in a temperate pasture on the west coast of the North Island of New Zealand (40°14'S, 175°16'E). The pasture had been under permanent grazing by sheep, cattle and goats since at least 1940. The mean annual rainfall at the site is 875 mm and average air temperature at a nearby weather recording station ranges from 8.0°C in July to 17.4°C in February. More details about the site botanical and physical characteristics can be found in Edwards et al., (2001). The experimental facility consisted of six FACE (Free Air CO₂ Enrichment) rings (McLeod and Long 1999), each 12 m in diameter; the rings were paired into three blocks based on initial
soil and the botanical characteristics. In each block, one ring was enriched with CO₂ at a target value of 475 μl l⁻¹ CO₂ (elevated CO₂) during the photoperiod, the second being left at ambient atmospheric CO₂ concentration (ambient CO₂). Enrichment began on 1 October 1997. Enriched rings were labelled R1, R2 and R3 (in blocks 1, 2 and 3 respectively) and ambient rings R4, R5 and R6 (blocks 1, 2 and 3). Each ring was fenced off individually in order to control sheep access and the duration and intensity of each grazing event.

2.3.2. Sheep grazing procedure

The usual grazing procedure throughout the experiment was intermittent grazing by adult sheep. Grazing started when the average aboveground biomass reached 1.8-2.0 t dry matter (DM) ha⁻¹ and continued until the residual aboveground biomass was reduced to approximately 0.5-0.7 t DM ha⁻¹. Details of the grazing methodology used for the two spring grazing reported here are given below.

November 2000

During the first grazing event, a single set of five mature wethers grazed all the rings sequentially; in this case individual sheep were used as replicates for the CO₂ treatment. The sheep were held indoors and starved for 24 h prior to grazing in order to remove the effects of their previous uncontrolled diet on dung composition, the average transit time of feed in the animals being 24 to 48 h (Haynes and Williams 1993). The sheep then grazed sequentially the three ambient rings (R4, R5 and R6) followed by the three enriched rings (R1, R2 and R3). Sheep were allowed to stay in each ring for 24 h and moved each morning. This sequential grazing began in R4 on 20 November and finished in R3 on 25 November 2000.

November 2001

In the second grazing event, adult wethers were contained within the same ring for the duration of grazing. Rings were taken as the replicate unit for analysis of CO₂ effects. The sheep were starved for 48 h prior to grazing in a devegetated pen. Since the initial aboveground biomass varied greatly between blocks, different numbers of animals were allocated to each block (2, 3 and 5 sheep per ring, in block
1, 2 and 3 respectively). Sheep were placed in the rings on 10 November and the experiment itself started on 11 November to minimize possible artefacts due to the 48 h starving period. Two sheep per ring were randomly selected for faecal collection (see below).

2.3.3. Collection and analysis of faeces

The five sheep in November 2000 and the two selected sheep per ring in November 2001 were equipped with harnesses and dung bags to allow total faecal collection. Sheep were trained with this equipment for at least 48 h during the week prior to the beginning of each grazing sequence to avoid excessive stress and any induced behaviour modification. Dung bags were fitted before sheep were allowed to enter in the rings and were emptied every 24 h. In 2000, sheep wore dung bags for six days, in 2001 for two days. All faecal samples were weighed and a sub-sample was taken and oven dried (60°C, to constant mass) for dry matter determination. This sub-sample was then ground to a fine powder and the C and N contents were determined at Lincoln University (New Zealand) with a mass spectrometer (PDZ Europa, UK).

2.3.4. Herbage analysis

Two randomly placed 1 m × 0.078 m quadrats were cut to 2 cm above ground level with powered hand shears from each compass quarter of each ring (total of eight per ring) on the day prior to and on the day after each grazing. The samples were bulked to give one sample per ring, and a sub-sample was immediately taken, placed briefly in a microwave oven (2 minutes at 600W) to arrest metabolism (Popp et al., 1996) in order to later determine the chemical composition of the mixed herbage. The dry mass of this sub-sample was later added to the dry mass of the bulked sample for biomass calculations. A sub-sample from the bulked herbage was sorted into species; these were oven dried separately (60°C, 48h). The remainder of the bulked sample was also dried, thus allowing calculation of species composition of the total biomass. To determine single species chemical composition, green leaves of the most abundant species of different functional groups - *Lolium perenne* L., *Agrostis capillaris* Sibth.,
*Anthoxanthum odoratum* L. (all C3 grasses), *Trifolium subterraneum* L. and *T. repens* L. (both legumes), *Paspalum dilatatum* Poir. (C4 grass) and *Hypochoeris radicata* L. (forb) - were sampled randomly in each ring on the day prior to grazing, dried on site in a microwave oven as above and transferred to the lab to be oven dried (60°C, 48h).

Herbage chemical composition relevant to ruminant diets was measured by NIRS (Near Infrared Reflectance Spectroscopy) with the feedTECH system (Corson *et al.*, 1999). The components measured were: nitrogen (N), acid detergent fibre (ADF), neutral detergent fibre (NDF), water soluble carbohydrates (WSC) and *in vitro* organic matter digestibility (OMD). NIRS results had been previously calibrated against wet chemistry analysis with herbage samples from the same site. The same calibration curves were used for both CO₂ treatments since no effect of elevated CO₂ could be found in previous experiments on correlations between wet chemistry and NIRS values.

2.3.5. Determination of the N budget

In both years, herbage data were measured in each ring so rings were used as the replicate to estimate CO₂ effects. In 2000 a single group of sheep grazed the six rings sequentially and, for this grazing, we considered that the best estimates of faecal production and composition were those measured once the sheep had grazed for 48 h in the same CO₂ treatment and were still in a ring of the same treatment. Thus we used faeces collected on the last day of sheep presence in a given CO₂ treatment (respectively in R6 and R3 for ambient and enriched treatments) and sheep were used as replicates to test the CO₂ effects on these variables. For this grazing we calculated intake quality as the average of the diet on offer in all three rings in each treatment. In 2001 we used faecal samples collected on the second day of grazing in the rings to measure faeces production and N excretion and rings were used as replicates after averaging data by ring for the two selected sheep. The diet quality values were in this case extracted from each ring separately. The use of the quality of herbage on offer as a measure of the sheep diet may not represent the actual diet if there is strong species selection, in particular positive selection for legumes (Parsons *et al.*, 1994). In 2000, despite the relatively high clover content in the pasture, no positive selection for
Table 2.1: Aboveground biomass prior to grazing events in two years and the contribution of different plant functional groups to this biomass (%) taken from pre-grazing cuts from pastures exposed to ambient or elevated (475 ppm) CO$_2$. Values are mean of three replicates± standard deviation. Data analysed as ANOVA using a split-plot model with CO$_2$ as the main plot and year as a subplot. P values shown when p<0.05 and n.s. when p>0.05.

<table>
<thead>
<tr>
<th></th>
<th>2000</th>
<th></th>
<th>2001</th>
<th></th>
<th>Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ambient</td>
<td>Elevated</td>
<td>Ambient</td>
<td>Elevated</td>
<td>CO$_2$</td>
</tr>
<tr>
<td>Biomass (g.m$^{-2}$)</td>
<td>184.6±63.2</td>
<td>171.7±58.7</td>
<td>199.1±54.2</td>
<td>201.3±37.6</td>
<td>n.s.</td>
</tr>
<tr>
<td>Proportion (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C3-grass</td>
<td>71.7±9.2</td>
<td>61.5±1.8</td>
<td>87.1±3.9</td>
<td>78±8.1</td>
<td>0.018</td>
</tr>
<tr>
<td>Legume</td>
<td>21.7±7.4</td>
<td>34.1±3.8</td>
<td>5±2.7</td>
<td>10.7±6.3</td>
<td>0.019</td>
</tr>
<tr>
<td>Forbs</td>
<td>3.5±4.3</td>
<td>2.7±1.2</td>
<td>3.3±2.5</td>
<td>6.1±2.9</td>
<td>n.s.</td>
</tr>
<tr>
<td>C4-grass</td>
<td>3.1±3.2</td>
<td>1.7±2.2</td>
<td>4.6±1.2</td>
<td>5.3±1.4</td>
<td>n.s.</td>
</tr>
</tbody>
</table>
legumes was measured. Indeed, a comparison of pre- and post-grazing botanical composition showed no difference in clover content ($p=0.138$) in the ambient or elevated CO$_2$ treatments. In addition the N concentration of the diet on offer before grazing was not different to the herbage remaining after grazing ($p=0.524$) showing that the animals did not preferentially remove N-rich species, legumes in particular. It was therefore considered that the apparent lack of diet selection enabled us to use diet on offer as an estimate of the ingested diet for 2000. In 2001 the very low legume content of the herbage on offer did not allow for intense sheep selection for clover, thus the quality of the pasture was again considered as a good estimate of the actual diet.

Sheep intake over 24 h was calculated from faecal production and the OMD values of the herbage:

$$\text{Intake (g DM sheep}^{-1} \text{ day}^{-1}) = \text{Faeces (g DM sheep}^{-1} \text{ day}^{-1}) / (1-\text{OMD})$$

Nitrogen intake was calculated from daily herbage intake and diet N concentration. N excreted in faeces was determined from individual daily faeces production and faeces N concentration. We calculated N excreted in urine as the difference between ingested N and N excreted in faeces assuming a fixed 5 % N retention; this value was used because N retention in animals is low on a yearly basis (Lambert et al., 1982), particularly in adult sheep (Ball, 1982). The potential for differences in N retention between treatments is discussed later.

2.3.6. Statistical analysis

Analysis of variance was used to analyse CO$_2$ effects on diet chemical composition in this randomised block design ($n=3$). For botanical composition, year was used as a sub-plot within CO$_2$ across both grazing events ($n=3$). A split-plot in time was appropriate since no covariance symmetry problems could arise as there were only two dates in the experiment. A split plot design was used to analyse single species chemical composition with CO$_2$ as a main plot ($n=3$) and species as a subplot and linear contrasts were used to compare functional groups. N partitioning data were analysed by ANOVA using sheep as replicates for the CO$_2$ treatment in 2000 ($n=5$) and in 2001 using rings as replicate for CO$_2$ and sheep as a blocking factor ($n=3$).
Table 2.2: Some leaf chemical characteristics (nitrogen (N), acid detergent fibre (ADF), neutral detergent fibre (NDF), water soluble carbohydrates (WSC) and organic matter *in vitro* digestibility (OMD)) of plant species growing under ambient or elevated (475 ppm) CO$_2$ (values are mean of three replicates in % of total dry matter) in two years, 2000 and 2001. Species were combined into functional groups for linear contrasts: grasses = *Paspalum dilatatum*, *Agrostis capillaris*, *Anthoxanthum odoratum*, *Lolium perenne*; legumes = *Trifolium repens*, *Trifolium subterraneum*; forb = *Hypochaeris radicata*. Data analysed as ANOVA using a split-plot model with CO$_2$ as the main plot and species as a subplot. Linear contrast comparing functional groups are also shown for 2000 data.

<table>
<thead>
<tr>
<th></th>
<th>N (%)</th>
<th>ADF (%)</th>
<th>NDF (%)</th>
<th>WSC (%)</th>
<th>OMD (%)</th>
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</thead>
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<tr>
<td></td>
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<td>Elevated</td>
<td>Ambient</td>
<td>Elevated</td>
<td>Ambient</td>
</tr>
<tr>
<td>2000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Agrostis capillaris</em></td>
<td>4.2</td>
<td>3.9</td>
<td>18.2</td>
<td>18.4</td>
<td>41.3</td>
</tr>
<tr>
<td><em>Anthoxanthum odoratum</em></td>
<td>3.8</td>
<td>3.5</td>
<td>14.4</td>
<td>14.7</td>
<td>34.5</td>
</tr>
<tr>
<td><em>Lolium perenne</em></td>
<td>3.1</td>
<td>2.5</td>
<td>15.4</td>
<td>16.3</td>
<td>35.9</td>
</tr>
<tr>
<td><em>Hypochaeris radicata</em></td>
<td>3.5</td>
<td>3.3</td>
<td>13.3</td>
<td>12.6</td>
<td>26.8</td>
</tr>
<tr>
<td><em>Paspalum dilatatum</em></td>
<td>3.3</td>
<td>3.6</td>
<td>27.4</td>
<td>26.0</td>
<td>52.8</td>
</tr>
<tr>
<td><em>Trifolium repens</em></td>
<td>5.4</td>
<td>4.6</td>
<td>14.0</td>
<td>13.3</td>
<td>23.8</td>
</tr>
<tr>
<td><em>Trifolium subterraneum</em></td>
<td>5.0</td>
<td>4.3</td>
<td>15.6</td>
<td>15.2</td>
<td>26.9</td>
</tr>
</tbody>
</table>

Origin of variance (p values)

- **CO$_2$**: 0.089, 0.588, 0.761, 0.055, 0.753
- **Species*: <0.001, <0.001, <0.001, <0.001, <0.001
- **Contrast grass/legumes**: <0.001, <0.001, <0.001, <0.001, <0.001
- **Contrast grass/forb**: 0.483, <0.001, <0.001, 0.154, 0.109
- **Contrast legume/forb**: <0.001, 0.019, 0.908, 0.858, <0.001

<table>
<thead>
<tr>
<th></th>
<th>N (%)</th>
<th>ADF (%)</th>
<th>NDF (%)</th>
<th>WSC (%)</th>
<th>OMD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ambient</td>
<td>Elevated</td>
<td>Ambient</td>
<td>Elevated</td>
<td>Ambient</td>
</tr>
<tr>
<td>2001</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Anthoxanthum odoratum</em></td>
<td>3.4</td>
<td>3.6</td>
<td>17.5</td>
<td>15.9</td>
<td>33.5</td>
</tr>
<tr>
<td><em>Hypochaeris radicata</em></td>
<td>3.5</td>
<td>3.3</td>
<td>15.8</td>
<td>15.9</td>
<td>29.8</td>
</tr>
<tr>
<td><em>Paspalum dilatatum</em></td>
<td>3.4</td>
<td>3.4</td>
<td>26.2</td>
<td>26.5</td>
<td>48.6</td>
</tr>
<tr>
<td><em>Trifolium subterraneum</em></td>
<td>4.1</td>
<td>3.0</td>
<td>19.7</td>
<td>22.0</td>
<td>33.2</td>
</tr>
</tbody>
</table>

Origin of variance (p values)

- **CO$_2$**: 0.469, 0.699, 0.674, 0.150, 0.729
- **Species**: 0.912, <0.001, <0.001, <0.001, <0.001
Linear regression was used to describe the relationship between diet WSC and digestibility and between legume content and proportion of N excreted in faeces. Genstat v 6.1 (Genstat, 2002) was used for all analysis.

2.4. Results

2.4.1 Botanical composition

The total pre-grazing aboveground biomass did not differ between treatments or years but the composition of this biomass was different (Table 2.1). Across both years legume content was significantly greater at elevated CO₂ (57% in 2000 and 114% in 2001); although much lower in absolute terms in 2001. The very low legume content in 2001 was primarily due to a very low presence of *T. repens* during the period in which it usually makes the major contribution to the legume pool (data not shown).

2.4.2 Single species chemical composition

In 2000 we examined the effect of atmospheric CO₂ concentration on the leaf chemical composition of seven species; comparisons were also made between functional groups by combining single species data (Table 2.2). N concentration was dependent on species (p<0.001) (Table 2.2) and tended to be lower under elevated CO₂ (9.2% less on average) but this was only a trend (p=0.089). If considered in terms of functional groups, the legume N concentration was about 40% higher than that of the grasses (p<0.001) and 42% higher than that of the forbs (p<0.001).

Across all species WSC concentration tended to increase under elevated CO₂ (p=0.055) by 6.8% (Table 2.2). WSC concentration differed between species (p<0.001) but the changes were similar between functional groups (Table 2.2).

ADF and NDF concentrations of the different species were not significantly affected by elevated CO₂ but differed markedly between species. The linear contrasts
Figure 2.1: Effects of elevated CO$_2$ on the concentration (% dry matter) of a) nitrogen, b) acid detergent fibres, c) neutral detergent fibres, d) water soluble carbohydrates and e) *in vitro* organic matter digestibility. Values are mean of three replicates ± standard deviation. Black bars: ambient CO$_2$, open bars: elevated CO$_2$. 
showed lower fibre concentrations of forbs and particularly of legumes compared to the grasses.

OMD was unaffected by elevated CO₂, and ranged from 70% for *P. dilatatum* to about 90% for the highly digestible species *L. perenne* and *T. repens*. The analysis by functional groups highlighted the significantly higher digestibility of legumes compared to the other two groups (Table 2.2).

In 2001 only four species (*A. odoratum*, *H. radicata*, *P. dilatatum* and *T. subterraneum*) could be sampled individually due to the very low abundance of *T. repens* in all rings and of *A. capillaris* and *L. perenne* in one ring. For this reason data were not analysed on the basis of functional groups as in 2000. The trend showing a decrease in N concentration at elevated CO₂ observed in 2000 was not evident in 2001 (Table 2.2). The average reduction in N concentration for all four species was 7.2% but this decrease was almost entirely attributable to the reduction in *T. subterraneum* N concentration. The N concentration in *T. subterraneum* was much lower in 2001 than in 2000 in both treatments but of a similar magnitude in the other species. WSC concentration under elevated CO₂ showed a trend consistent with results from 2000 (4.8% greater under elevated CO₂) but the difference was not statistically significant. Other chemical composition parameters were not affected by elevated CO₂. Interspecific variations were consistent with those observed in 2000, in particular the high fibre concentration of *P. dilatatum* and its low digestibility.

### 2.4.3. Chemical composition of the offered diet

When examined as a mixture i.e. integrating both changes in individual leaf chemical composition and species presence and representing the average diet offered to the sheep, the average N concentration of the herbage was unaffected by elevated CO₂ (Fig. 2.1) in either year but was lower in 2001 (2.4%) than in 2000 (2.8%).

In 2001 both NDF and ADF concentrations of the herbage were significantly reduced by elevated CO₂ (16% (p<0.01) and 16.2% (p<0.01) respectively (Fig. 2.1). In 2000 no statistically significant difference was detected between treatments but a similar trend occurred for both ADF and NDF. In both years WSC and OMD were
Figure 2.2: Correlation between herbage water soluble carbohydrates (WSC) content (in % dry matter) and herbage in vitro organic matter digestibility (OMD) (in % dry matter) for samples taken from pasture growing at ambient or elevated (475 ppm) CO₂. Fitted linear regression line, its equation and regression coefficient are shown.

$$\text{OMD} = 42.76 + 1.45 \times \text{WSC}$$

$$R^2 = 0.8134$$

Figure 2.3: Correlation between the proportion of legumes in herbage (% of total dry matter) and the proportion of dietary N excreted as urine by animals grazing pastures exposed to ambient or elevated (475 ppm) CO₂ during spring of 2000 and 2001. Fitted linear regression line and its regression coefficient are shown.

$$R^2 = 0.76$$
higher at elevated CO$_2$ but the difference was only significant in 2001. Digestibility data from both years were well correlated with WSC concentration ($r^2=81.3$, $p<0.001$) (Fig. 2.2).

2.4.4. Nitrogen partitioning in the excreta

Herbage intake was calculated from the in vitro digestibility of the diet and faecal production. In both years the mass of faeces produced was lower under elevated CO$_2$ (by 21 and 5% in 2000 and 2001 respectively) but this difference was significant only in 2000 ($p=0.008$) (Table 2.3). Since OMD was higher under elevated CO$_2$, as described above, the calculated intake was unaffected by the CO$_2$ treatment but was about 40% lower in 2001 than in 2000. Because of the similar N concentration of the feed on offer regardless of the CO$_2$ level, the assumed dietary N was also similar for CO$_2$ treatments but again was lower in 2001. Faecal N output was calculated from daily faecal production and faecal N concentration. No statistically significant difference was observed between the CO$_2$ treatments in term of N concentration in the faeces despite a clear trend in 2000 ($p=0.051$) showing a N decrease under elevated CO$_2$. As a result, faecal N output was significantly lower under elevated CO$_2$ in 2000 (33%, $p=0.006$) but was unaffected in 2001 (Table 2.3). Faecal N output expressed as a proportion of ingested N was clearly lower under elevated CO$_2$. In 2000 the proportion of ingested N excreted in faeces dropped from 33.6% under ambient CO$_2$ to 24.4% under elevated CO$_2$. In 2001, the shift was of a similar magnitude (41.2% and 34.5% under ambient and elevated CO$_2$ respectively) but was only a trend ($p=0.07$). As shown in Fig. 2.3, the proportion of N excreted in faeces was strongly related to the proportion of legumes in the diet. These two variables were negatively correlated ($r^2=0.76$, $p<0.001$). Since the proportion of N excreted in urine was calculated by difference (see Materials and Methods section) it exhibited the opposite response to faeces and was higher under elevated CO$_2$. 

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Table 2.3: Nitrogen budget for sheep grazing pastures exposed to ambient or elevated (475 ppm) CO₂. Values are means with n=3 for digestibility and N content of the herbage, n=5 for other data in 2000 and n=6 for other data in 2001. Data analysed as ANOVA (see Materials and Methods section for information on the models).

<table>
<thead>
<tr>
<th></th>
<th>Ambient CO₂</th>
<th>Elevated CO₂</th>
<th>CO₂ effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>2000</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of sheep</td>
<td>5</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Intake</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>In-vitro digestibility of herbage (%)</td>
<td>73</td>
<td>76.1</td>
<td>p=0.07</td>
</tr>
<tr>
<td>Calculated intake (g DM.sheep⁻¹.day⁻¹)</td>
<td>1528</td>
<td>1408</td>
<td>p=0.13</td>
</tr>
<tr>
<td>N% in herbage</td>
<td>2.8</td>
<td>2.8</td>
<td>p=0.8</td>
</tr>
<tr>
<td>Dietary N (gN.sheep⁻¹.day⁻¹)</td>
<td>42.8</td>
<td>39.4</td>
<td>p=0.126</td>
</tr>
<tr>
<td><strong>Faeces output</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mass excreted (g.sheep⁻¹.day⁻¹)</td>
<td>412.4</td>
<td>325.4</td>
<td>p=0.008</td>
</tr>
<tr>
<td>N% in faeces</td>
<td>3.5</td>
<td>3.1</td>
<td>p=0.051</td>
</tr>
<tr>
<td>Faecal N (gN.sheep⁻¹.day⁻¹)</td>
<td>14.4</td>
<td>9.6</td>
<td>p=0.006</td>
</tr>
<tr>
<td>Faecal N (% of ingested N)</td>
<td>33.6</td>
<td>24.4</td>
<td>p=0.001</td>
</tr>
<tr>
<td><strong>Urine output</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N retention (% of ingested N)</td>
<td>5</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Calculated urine-N (% of ingested N)</td>
<td>61.4</td>
<td>70.6</td>
<td>p=0.001</td>
</tr>
<tr>
<td><strong>2001</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of sheep</td>
<td>6</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Intake</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>In-vitro digestibility of herbage (%)</td>
<td>66.8</td>
<td>73.3</td>
<td>p=0.003</td>
</tr>
<tr>
<td>Calculated intake (g DM.sheep⁻¹.day⁻¹)</td>
<td>836</td>
<td>985</td>
<td>p=0.42</td>
</tr>
<tr>
<td>N% in herbage</td>
<td>2.4</td>
<td>2.4</td>
<td>p=0.92</td>
</tr>
<tr>
<td>Dietary N (gN.sheep⁻¹.day⁻¹)</td>
<td>20.3</td>
<td>24</td>
<td>p=0.39</td>
</tr>
<tr>
<td><strong>Faeces output</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mass excreted (g.sheep⁻¹.day⁻¹)</td>
<td>277</td>
<td>263.2</td>
<td>p=0.83</td>
</tr>
<tr>
<td>N% in faeces</td>
<td>3</td>
<td>3.1</td>
<td>p=0.55</td>
</tr>
<tr>
<td>Faecal N (gN.sheep⁻¹.day⁻¹)</td>
<td>8.3</td>
<td>8.3</td>
<td>p=0.99</td>
</tr>
<tr>
<td>Faecal N (% of ingested N)</td>
<td>41.2</td>
<td>34.5</td>
<td>p=0.07</td>
</tr>
<tr>
<td><strong>Urine output</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N retention (% of ingested N)</td>
<td>5</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Calculated urine-N (% of ingested N)</td>
<td>53.8</td>
<td>60.5</td>
<td>p=0.07</td>
</tr>
</tbody>
</table>

*Urine-N was calculated by difference from faecal N therefore p values for CO₂ effect are identical for the two variables.
2.5. Discussion

The main objective of this study was to assess if elevated CO₂ effects on pasture chemical and botanical composition would affect nitrogen cycling through ruminants. We showed that N partitioning between faeces and urine returns was affected by elevated CO₂ and this has possible implications for increased N losses. We used data from two non consecutive grazing events, the first in 2000 and the second in 2001. In order to discuss overall patterns observed during these two grazing, a brief summary of the inter-annual differences is required. Despite the fact that both grazing events took place in spring under similar climatic conditions, the pasture in 2001 was still expressing a carry over of an extreme spring drought that occurred 2 months before our experiment. The total rainfall in September 2001 was 3 mm while the 20 years average for this month is 60 mm. *Trifolium repens* is particularly vulnerable to spring drought (Brock 1988) and this was reflected in the extremely low legume abundance in 2001 compared to 2000. A second difference between years was the lower apparent feed intake observed in 2001; we are unable to identify the cause of this difference but intake rates can vary for many reasons (e.g. environmental, feed or animal related) (McDonald et al., 1981). If the cause was a difference in the physiological state of the animals this could potentially influence N retention; however, as mature wethers were used in both years we do not consider this to be relevant to our results.

2.5.1. CO₂ effects on chemical composition of single species

In a survey of 27 C₃ species, Poorter et al., (1997) found that the main changes in leaf chemical composition in plants grown under elevated CO₂ were an increase in non-structural carbohydrates and a decrease in leaf N concentration. While the increase in non-structural carbohydrates under elevated CO₂ was large (49% on average), there was high interspecific variability with responses ranging from almost zero to over 100%. The decrease in N concentration of these plants was 15% at elevated CO₂, again with high variability between species. In a survey of 67 species Cotrufo et al., (1998) calculated a 14% decrease in N concentration at elevated CO₂ averaged over different experimental conditions, plant type and tissues. In our
experiment, averaged across species leaf N concentration dropped by 9% and 8% and the soluble sugars increased by 6% and 5% in 2000 and 2001 respectively (Table 2.2). These differences were not significant but reflect similar significant differences measured on this pasture (von Caemmerer et al., 2001) and in other temperate grassland studies e.g. for L. perenne (Soussana et al., 1996) and T. repens (Hartwig et al., 2000). These data therefore support the view that leaf N and soluble sugar concentrations are the leaf chemistry parameters that exhibit the strongest response to elevated CO₂.

Nitrogen decrease in tissues, especially leaves, exposed to elevated CO₂ may occur through a combination of effects (Cotrufo et al., 1998); two of the most likely mechanisms being: 1) a metabolic down-regulation of enzymes involved in photosynthesis as a regulatory response to the CO₂-stimulated photosynthesis rate, and 2) a dilution effect with N concentration reduced by an accumulation of non-structural carbohydrates (Gifford et al., 2000). However, if the N concentration of single species in our experiment is recalculated on a soluble sugars-free basis then N was still lower at elevated CO₂ (8.3% in 2000 and 7.8% in 2001). This suggests that the elevated-CO₂ plants had lower N concentration due to a reduction in the amount of photosynthetic enzymes. In support of this view, data from the same experimental site has shown a strong photosynthetic acclimation to elevated CO₂ by both legumes and grasses (von Caemmerer et al., 2001).

Other leaf chemistry responses to elevated CO₂ that might alter forage quality are increases in the synthesis of secondary products, in particular lignin (Gifford et al., 2000). Although we did not directly assess the question of lignin concentration, the fact that both the ADF and NDF concentration were not affected under elevated CO₂ conditions allows us to hypothesise that highly recalcitrant compounds like lignin were not significantly increased under elevated CO₂ in our experiment (Fig. 2.1). Moreover, OMD of individual species was not affected by elevated CO₂, showing that at the species scale, no major modification of tissue quality (from the ruminant perspective) occurred. The elevated CO₂-induced changes in WSC and N concentration reported here would have opposite effects on digestibility if taken alone. A decrease in the herbage N concentration usually leads to a limitation of microbial development in the rumen and thus a reduced capacity to digest compounds.
like cellulose and hemicellulose. On the other hand, increases in highly digestible soluble sugars in the diet increase digestibility. Here, the CO$_2$ driven changes observed in both N and WSC seem to counterbalance each other, leaving the digestibility of individual species unaffected (Table 2.2).

2.5.2. Effects of elevated CO$_2$ on botanical composition

We observed a significant increase in the proportion of legumes in the pasture at elevated CO$_2$ (Table 2.1) as shown in many other studies and systems e.g. sown bispecific mixture (Hebeisen et al., 1997; Soussana and Hartwig 1996) and multispecific complex pasture ecosystems (Newton et al., 1994; Teyssonneyre et al., 2002). Although a positive response of legumes to elevated CO$_2$ is frequently observed, it is not an invariable response (Leadley et al., 1999), since the legume response can be modified by other factors such as phosphorus availability (Stöcklin and Kömer 1999), temperature (Newton et al., 1994), cutting frequency and N fertilisation (Hebeisen et al., 1997). A potentially important modifying factor in our experiment was the presence of large herbivores that are known to express a positive preference for legumes (Parsons et al., 1994). No active selection could be measured in this short term experiment (see Materials and Methods section) but is clearly an issue on a yearly basis and raises the possibility of differences arising in composition between grazed and cut swards, and, indeed, data from this site confirms the expectation of lower legume content under grazing (Newton et al., 2001). Moreover, it appears that the relative clover suppression by grazing allows for greater expression of the CO$_2$ response as the difference between treatments in legume proportion was much greater under grazing than cutting (Newton et al., 2001).

2.5.3. Integration of direct and indirect effects of elevated CO$_2$ on herbage quality

One important characteristic of large herbivores feeding in a pasture exposed to elevated CO$_2$ is their ability to integrate, through grazing, CO$_2$ effects that occur both at the plant and community levels. In this instance, the shift in the botanical composition towards a higher proportion of legumes counterbalanced the N decrease
observed at the single species scale, resulting in an N concentration of the overall diet that was unaffected by elevated CO₂ (Fig. 2.1). A similar phenomenon was observed in a cut grassland FACE experiment where a grass (*L. perenne*) and a legume (*T. repens*) were grown alone or in mixture (Hartwig *et al.*, 2000).

In contrast to N, changes at the species level and at the sward level appeared to combine additively in relation to WSC. As there was a significant correlation between WSC and digestibility (as previously observed by Dent and Aldrich, 1963; Humphreys, 1989), there was also an increase in digestibility of the high CO₂ forage. This result matches that found in a Mini-FACE experiment under cutting (Teyssonneyre 2002; Picon-Cochard *et al.*, 2003); here digestibility also increased in response to CO₂ despite reduced crude protein concentration. These data, and the strong relationship between soluble sugars (rather than N) and digestibility (Fig. 2.2) suggest that the widespread response to CO₂ of increased soluble sugars might lead to an increase in forage digestibility.

2.5.4. Implications of a modified forage quality for nutrient cycling through ruminants

Our data show that despite having diets containing similar N concentrations, sheep grazing the elevated CO₂ pasture excreted proportionally less N in their dung and more in their urine. Because these sheep were mature they could be considered at maintenance i.e. had no significant N sink, we calculated urinary N as the difference between ingested and faecal N assuming a low N retention (5%). For this shift in N partitioning to be buffered by an increased N retention under elevated CO₂, it would require retention rates of about 14% and 12% in 2000 and 2001 respectively under elevated CO₂ conditions; i.e. a 2-3 fold increase in N retention under elevated CO₂. Given that N retention in non-pregnant, non-lactating animals is mainly driven by animal physiological needs, such an increase does not seem reasonable and our calculations probably reflect a real shift in N partitioning under elevated CO₂.

In general, the proportion of N excreted in the urine is positively correlated to N concentration of the diet (Jarvis 2000); however, in our case no difference in
dietary N concentration was apparent and we need to look elsewhere for an explanation for the change in partitioning. One strong candidate is the higher proportion of legumes ingested by the sheep on the elevated CO$_2$ pasture, particularly in 2000. A number of studies on a range of animals have shown increased allocation of N to urine as the proportion of clover in the diet increases even though the total N in the diet remains unchanged. For example MacRae and Ulyatt (1974), in a study of fresh forage digestion by sheep, measured a higher proportion of ingested N excreted in urine when sheep were fed with white clover (*T. repens*) (77.7%) compared to perennial (*L. perenne*) (68.2%) or annual ryegrass (*Lolium multiflorum* L.) (59%), despite a similar amount of ingested N in the three diets. The same trend was observed on ewes fed with pure ryegrass or ryegrass/white clover mixed swards by Orr *et al.* (1995). Again, data in Button *et al.*, (1967) show that variation in the N concentration of dairy cows’ urine throughout a milking season was strongly correlated with the proportion of clover in the diet but not with the N concentration of the diet.

The reason for the enhancement of urinary N output for a given amount of ingested N is still unclear. It has been hypothesised that a legume increase would create an imbalance between protein intake and rapidly fermentable energy, thereby altering the efficiency of N use in the rumen (Beever 1993) and enhancing the production of ammonia that will eventually be excreted in the rumen (Evans *et al.*, 1996). This does not fit with our data since the diet WSC concentration slightly increased under elevated CO$_2$ (Fig. 2.1). We may thus hypothesise that N partitioning within the ruminant is affected by the type of N offered in the diet, a variable that is possibly affected by elevated CO$_2$ beyond the shift in botanical composition. This would explain the results obtained in 2001 with a low legume content. It is known that N use efficiency in the rumen can be affected by the degree of protein protection, by tannins for example (Waghorn 1985) or the amount of nitrate-N in the diet (Marais 1980). In a recent paper, Kebreab *et al.*, (2001) discussed the complex interactions between energy source and protein digestibility, both of which being potentially affected by elevated CO$_2$, on N partitioning in dairy cows. This highlights the need for more comprehensive experiments to elucidate the basis of the CO$_2$ effect reported here and whether this effect is purely driven by the increase in legumes in the herbage.
2.6. Conclusion

In this study we have shown that elevated CO₂ affects forage quality, both through changes in single species chemical composition and through altering the balance of plant species in the pasture in favour of legumes. Taken together, this leads to bulk forage with a slightly higher digestibility and similar N intake. Importantly, in relation to nutrient cycling, we estimated an increased proportion of excreted N in the urine relative to the faeces at elevated CO₂. N losses from urine through leaching and volatilisation can reach nearly 30% and 20% respectively of deposited N, making grazed pastures far from closed systems with respect to N (Ball et al., 1979). In contrast, N losses from faeces are small (Haynes and Williams 1993) due to a low proportion of water-soluble N. Therefore we anticipate less efficient N cycling under grazing at elevated CO₂. The pathway of N loss is strongly soil dependent and in our pasture losses occur most readily through ammonia volatilisation, reaching approximately 20 kg N ha⁻¹ year⁻¹ under ambient CO₂ i.e. 15-20% of the N cycled through the animals (RA Carran, personal communication). Given the 15% increase in N partitioning towards urine shown here at elevated CO₂ and leaving aside any potential concentration dependent increase in volatilisation, an extra 3 kg N ha⁻¹ year⁻¹ of ammonia-N would be volatilised at high CO₂. As about 20% of the terrestrial biosphere is grazed by livestock and as between 40 and 75% of N cycles through animals rather than leaf litter, we suggest that this is a potentially important global response to elevated CO₂.

2.7. Acknowledgements

Financial support from the New Zealand Foundation for Research, Science and Technology (Contract C10X205) is gratefully acknowledged. V. Allard was funded by a Massey University Doctoral Scholarship. His stay in New Zealand was also supported by a grant from the Scientific and Cultural service of the French Embassy in New Zealand. We want to thank S. Brock, S. Dunn, E. Lawrence, T. Rayner for their technical assistance, F. Potter for statistical advice, G. Waghorn for advice on animal physiology, D. Corson for NIRs analysis and R. Cresswell for C-N
analysis. We also want to thank S. Ledgard, J. Crush, H. Jones and three anonymous referees for useful comments on the manuscript.

2.8. References


Chapter 2


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Plate 3.1: Litter incubation bags in the field. Leaf material is placed at the soil surface and root material is buried.

Chapter 3: Elevated CO₂ Effects on Decomposition Processes in a Grazed Grassland
Chapter 3: Elevated CO₂ Effects on Decomposition Processes in a Grazed Grassland

Allard V, Newton PCD, Lieffering M, Matthew C

3.1. Abstract

In situ decomposition of plant litter and animal faecal material was studied, over two years, in the New Zealand Free Air CO₂ Enrichment (FACE) facility. As expected, elevated CO₂ did not affect decomposition rates of plant litter because quality of the litter at the species scale was not altered. Nevertheless, indirect effects of elevated CO₂ on the pasture botanical composition and biomass allocation were found and these effects tended to increase decomposition rates at the ecosystem scale. Decomposition of faecal material was slower under elevated CO₂ during a summer experiment but not during winter. This highlights the complex relationships between animal returns and other ecosystem components in the context of elevated CO₂ and also the importance of taking grazers into account when predicting the effect of elevated CO₂ in grasslands. When direct and indirect effects of elevated CO₂ were taken into account, the two counterbalanced each other, and average organic matter decomposition rate in this ecosystem was not appreciatively altered by elevated CO₂.
3.2. Introduction

Understanding the response of organic matter (OM) decomposition to the rise in atmospheric CO₂ (Keeling et al., 1995) is central to the prediction of ecosystem functioning under our ongoing changing climate. In grasslands, carbon (C) sequestration occurs mostly belowground and because grasslands cover about 20% of the land surface, grassland soils are major C sinks (Thomley et al., 1991). Thus, increased C stocks in grassland soils could potentially buffer the atmospheric CO₂ increase, but this process is largely dependent on the capacity of OM to retain C during decomposition. In addition, the nitrogen (N) immobilization-mineralisation balance during OM decomposition is a key process determining ecosystem functioning and may act as a feedback to the ecosystem response to elevated CO₂ (Diaz et al., 1993; Zak et al., 1993).

Decreased quality of litter at the individual species level under elevated CO₂ was long considered the major mechanism altering litter decomposition (Strain and Bazzaz, 1983; Coutéau et al., 1996; Cotrufo et al., 1995). However, this hypothesis has now been discarded because the reduction in plant quality observed in green material (Cotrufo et al., 1998) is only partially reflected in senescing tissue (Norby and Cotrufo, 1998). Therefore no changes, or at most only small changes in individual species litter decomposition rates have been observed under elevated CO₂ (Hirschel et al., 1997; Van Vuuren et al., 2000). Litter decomposition at the ecosystem scale might nevertheless be strongly altered if possible indirect effects of elevated CO₂ on botanical composition and biomass partitioning between shoots and roots are taken into account. Indeed, it was concluded that CO₂ driven shifts in species composition could change average decomposition rates in a C₃-C₄ grassland due to the relative difference in degradability of these functional groups (Kemp et al., 1994). Similar findings were obtained from a natural CO₂ spring (Ross et al., 2002). In addition, belowground biomass allocation is usually increased under elevated CO₂ as shown by lower shoot: root ratios (Cotrufo and Gorissen, 1997; van Ginkel et al, 1997) and increased root turnover (Canadell et al., 1996; Fitter et al., 1996). This might alter decomposition rate at the system scale if root decomposition is slower than leaf litter decomposition (Gorissen and Cotrufo, 2000).
Chapter 3

The current understanding of organic matter decomposition processes in grasslands under elevated CO₂ is exclusively based on data for simulated cut grasslands (Newton et al., 2001) in spite of the infrequent use of this management at a global scale. In the presence of herbivores a major part of organic matter returns to the soil occurs through faeces and urine (Parsons et al., 1991; Orr et al., 1995), therefore faeces decomposition may also be of importance in the context of elevated CO₂. Indeed herbivores feed mainly on green tissue, material that exhibits stronger effects of elevated CO₂ than senescing material and in addition grazing animals also integrate CO₂ effects at the community level in particular possible changes in botanical composition (Chapter 2).

This study compares elevated CO₂ effects on the decomposition rates of the different materials that contribute to organic matter inputs to a grassland soil and their capacity to release N during decomposition. To this end, the results of three different decomposition experiments that took place in the New Zealand pasture FACE experiment between 2000 and 2002 are presented. In particular, attention is focussed on the possible effects of CO₂-induced shifts in botanical composition and shoot:root biomass partitioning that might significantly alter decomposition rate at the ecosystem scale. In addition, to assess an important process in grazed grassland, the question of potential effects of elevated CO₂ on animal returns through changes in plant chemical composition and botanical composition will be addressed. We then bring these fluxes together to compare their relative importance and the net outcome in term of decomposition rate and N release.

3.3. Materials and Methods

3.3.1. Experimental site

The study was carried out in a temperate pasture on the west coast of the North Island of New Zealand (40°14′S, 175°16′E). The pasture had been under permanent grazing by sheep, cattle and goats since at least 1940. The mean annual rainfall at the site was 875 mm and average air temperature at a nearby recording station ranges from 8°C in July (minimum) to 17.4°C in February (maximum).
Species diversity in this pasture is high; more than 20 vascular species were found during an inspection of the vegetation in 1996. The pasture is dominated by the C\textsubscript{3} grasses *Lolium perenne* L., *Agrostis capillaris* L. and *Anthoxanthum odoratum* L., the C\textsubscript{4} grass *Paspalum dilatatum*, the legumes *Trifolium repens* L. and *Trifolium subterraneum* L., and the forbs *Hypochaeris radicata* L. and *Leontodon saxatilis* L. The experimental facility consisted of six FACE (Free Air CO\textsubscript{2} Enrichment) rings (McLeod and Long 1999), each 12 m in diameter; which had been paired based on initial soil and botanical characteristics. In each pair (or block), one ring was fumigated with CO\textsubscript{2} at a target value of 475 μl l\textsuperscript{-1} CO\textsubscript{2} (enriched treatment) during the photoperiod, and the second was left at atmospheric CO\textsubscript{2} concentration (ambient treatment) with enrichment beginning on 1 October 1997.

### 3.3.2. Sheep faeces collection and decomposition

Two separate faeces decomposition experiments were set up in 2000 and 2001. Because the first experiment took place under a relatively extreme drought, a second was set up under more common climatic conditions. Methodology details for these two experiments are given below.

In November 2000 a single group of five sheep grazed all the rings sequentially. In this case, individual animals were used as replicates for the CO\textsubscript{2} treatments. The sheep were held indoors and starved for 24 h prior to the grazing in order to remove the effect of their previous diet on faeces composition. The sheep first sequentially grazed the three enriched rings followed by the three ambient rings. They were allowed to stay in each ring for 24 h and moved each morning to the next ring. Throughout this grazing, sheep wore harnesses and faecal collection bags. The bags were emptied every 24 h. For this experiment, we collected faeces after the sheep had grazed for 48 h in the same CO\textsubscript{2} treatment (i.e. were still in a ring of the same treatment). We thus collected five replicate samples of faecal material from both CO\textsubscript{2} treatments. Hence, individual animals were used as replicates for the CO\textsubscript{2} treatments.
A sub-sample of the faeces was taken and oven-dried (60°C, to constant mass) to determine dry matter content, and then finely ground before analysis of initial C and N concentration at Lincoln University with a mass spectrometer (PDZ Europa, UK). The remaining material was kept in a cold room (4°C) until preparation of the incubation bags. The incubation bags (5 cm × 5 cm) were made of polyethylene mesh (mesh size = 1mm) and filled with a known weight of fresh faecal material (approximately 5 g fresh weight). A cross-over design was used to compare the effects of the CO₂ concentration where the faeces originated from (CO₂ origin) and the CO₂ concentration during incubation (CO₂ incubation). In addition, enough bags were set up to allow for four retrieval dates (2, 4, 6 and 16 weeks of incubation). Therefore, 40 incubation bags (2 CO₂ origin × 4 dates × 5 sheep) were placed in each ring. A nail was used to keep each incubation bag in close contact with the soil surface. The vegetation was cut to about 1 cm prior to the bag placement in December 2000. At each of the four sampling times, the remaining faecal material was separated from adhering soil and newly grown vegetal material, oven dried (60°C, until constant mass), weighed and ground prior to C and N concentration determination.

In June 2001 sheep were kept in the same ring for the duration of grazing. Thus, in this instance, rings were the replicate units for detection of the CO₂ effect. Two sheep were allocated to the rings of the first block and three sheep per ring in the second and third blocks. This methodology was used because pre-grazing herbage biomass varied between blocks. In this second collection, animals did not wear faeces collection bags. Instead, after three days of grazing in the rings, fresh faecal samples were collected directly from the ground. These samples were then processed and placed in incubation bags, 40 per ring (2 CO₂ origin × 4 dates × 5 samples), as described above using the same experimental design. There were four retrieval dates with sampling taking place after 4, 6, 8 and 12 weeks of incubation.

3.3.3 Plant leaf and litter collection and decomposition

A third experiment was set up to study vegetal material decomposition. This experiment dealt mainly with senesced leaf litter and root decomposition but green
leaf material was also included in the study in order to highlight how the resorption processes during leaf senescence modified decomposition characteristics. Furthermore, green material decomposition was considered to be a good standard for comparison with faeces decomposition. In November 2001, green leaves of four species belonging to different functional groups—Anthoxanthum odoratum (C3 grass), Trifolium subterraneum (legume), Hypochaeris radicata (forbs) and Paspalum dilatatum (C4 grass)—were sampled in each ring. These species were chosen from diverse functional groups in order to assess possible species effects on decomposition. In addition, leaf litter of the same species was collected. The methodology used to collect litter varied according to species. From patches where P. dilatatum and A. odoratum were highly dominant, newly fallen dead leaves were collected. However, attached dead leaves of T. subterraneum and H. radicata were collected as the litter sample, since no loose dead leaves originating from those plants were found at the soil surface. All plant samples were oven dried (60°C, to constant mass) and were kept in a cold room (4°C) until preparation of the incubation bags. To facilitate sampling, sets of 16 incubation bags (2 CO2 origin × 4 species × green or litter) were made from a single piece of polyethylene mesh (30 cm × 30 cm, mesh size = 1 mm). Individual bags within each group were sealed with a soldering iron and filled with a known weight of approximately 0.35 g dry matter (DM) of material. A sufficient number of bags were set up to enable four retrieval dates. Litter bags were nailed in contact with the soil at the end of January 2001, and retrieval was every two months thereafter. After removal, samples were processed as described above for faecal samples and C and N content was analysed in INRA (France) with an elemental analyser (Carlo Erba, Italy).

3.3.4. Plant root collection and decomposition

Root material was collected from another experiment (Chapter 4) investigating the effect of elevated CO2 on root growth using the “ingrowth core” method between January and October 2001. The root material used was thus derived from newly grown roots (maximum 2 months old) and did not include dead roots. The potential implications of this are discussed later. Due to the high botanical diversity in the
Table 3.1: Effects of elevated CO₂ on N concentration (%) and C/N ratio prior to incubation of green leaf material and leaf litter from *Trifolium subterraneum* (Tri), *Hypochaeris radicata* (Hyra) and *Paspalum dilatatum* (Pasp) and *Anthoxantum odoratum* (Anth), root material and faeces from the summer experiment and the winter experiment. Values are mean of 3 replicates ± s.d. Effects of the treatments are presented in table 3.2.

<table>
<thead>
<tr>
<th>[CO₂] Origin</th>
<th>Material</th>
<th>N</th>
<th>C/N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Green Anth</td>
<td>2.8±0.1</td>
<td>3±0.5</td>
<td>15±0.7</td>
</tr>
<tr>
<td>Green Hyra</td>
<td>2.6±0.3</td>
<td>2.5±0.4</td>
<td>15.5±1.8</td>
</tr>
<tr>
<td>Green Pasp</td>
<td>3.1±0.2</td>
<td>3.1±0.4</td>
<td>14.2±1.2</td>
</tr>
<tr>
<td>Green Tri</td>
<td>3.5±0.3</td>
<td>3±0.4</td>
<td>12.5±0.9</td>
</tr>
<tr>
<td>Litter Anth</td>
<td>1.2±0.2</td>
<td>1.3±0.2</td>
<td>34.7±5.8</td>
</tr>
<tr>
<td>Litter Hyra</td>
<td>1.2±0.1</td>
<td>1.2±0.2</td>
<td>29±3</td>
</tr>
<tr>
<td>Litter Pasp</td>
<td>1.2±0.1</td>
<td>1.1±0.2</td>
<td>34.8±4.3</td>
</tr>
<tr>
<td>Litter Tri</td>
<td>2.5±0.1</td>
<td>2.1±0.2</td>
<td>17.9±0.6</td>
</tr>
<tr>
<td>Roots</td>
<td>1.1±0.2</td>
<td>1.1±0.04</td>
<td>31.6±5.4</td>
</tr>
<tr>
<td>Faeces Summer</td>
<td>3.48±0.37</td>
<td>3.1±0.22</td>
<td>13.27±1.49</td>
</tr>
<tr>
<td>Faeces Winter</td>
<td>2.48±0.47</td>
<td>2.39±0.29</td>
<td>16.66±2.83</td>
</tr>
</tbody>
</table>
rings, root material from particular plant species could not be collected separately. Six pseudo-replicates per ring were nevertheless sampled to take into account the botanical variability. The incubation bags were placed in groups of 12 (2 CO₂ origin x 6 samples), using a technique similar to that described above for dung samples, but with bags buried vertically with the upper edge 2 cm below the soil surface. As with the leaf material, incubation began at the end of January 2001 and a set of bags was retrieved after 2, 4, 6 and 8 months of incubation, except that bags for the third sampling date (6 months of incubation) were removed from the study because of problems during sampling.

3.3.5. Relative N loss by decomposing material

In order to assess the rate of N release by the different types of decomposing material the relative N loss was calculated as follow:

Relative N loss = (N₁ - Nₜ) / (DM₁ - DMₜ)

Where N₁ and Nₜ are the amount of N in the material before incubation and after sampling respectively (g) and DM₁ and DMₜ the mass of the material at the same dates (g). The relative N loss was only calculated for one retrieval date for each type of material: after four months of incubation for plant material and 16 and 6 weeks for faecal material in the summer and winter experiment respectively.

3.3.6. Summary scenarios

To discuss the overall effect of elevated CO₂ on litter decomposition at the ecosystem scale, different scenarios based on data from this experimental site were used. The effect of a doubled proportion of dicots in the pasture under elevated CO₂ (scenario 1) was inspired by a result obtained in October 2000 (Chapter 2). The second scenario (scenario 2) assesses the effects of an enhanced biomass allocation to roots under elevated CO₂ on decomposition rate based on data obtained in chapter 4. Scenario 3 is a combination of scenarios 1 and 2 and simulates what might be the overall decomposition rate in a pasture under elevated CO₂ without considering any
Table 3.2: P-values from the analysis of variance of the effects of CO₂, species (Trifolium subterraneum, Hypochaeris radicata, Paspalum dilatatum and Anthoxantum odorantum) and their interaction, on the N concentration and C/N ratio of green leaf material and leaf litter prior to the decomposition. P-values from the analysis of variance of the effects of CO₂ on the N concentration and C/N ratio of root material and faeces material from the summer and winter experiment prior to the decomposition, are mentioned.

<table>
<thead>
<tr>
<th>Material</th>
<th>CO₂ N</th>
<th>Species</th>
<th>CO₂*species</th>
</tr>
</thead>
<tbody>
<tr>
<td>Green</td>
<td>0.727</td>
<td>0.025*</td>
<td>0.341</td>
</tr>
<tr>
<td>C/N</td>
<td>0.743</td>
<td>0.142</td>
<td>0.575</td>
</tr>
<tr>
<td>Litter</td>
<td>0.334</td>
<td>&lt;0.001***</td>
<td>0.759</td>
</tr>
<tr>
<td>C/N</td>
<td>0.718</td>
<td>&lt;0.001***</td>
<td>0.759</td>
</tr>
<tr>
<td>Roots</td>
<td>0.875</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>C/N</td>
<td>0.688</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>Faeces</td>
<td>0.0870</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>summer C/N</td>
<td>0.112</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>Faeces</td>
<td>0.781</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>winter C/N</td>
<td>0.945</td>
<td>/</td>
<td>/</td>
</tr>
</tbody>
</table>

Figure 3.1: Disappearance rate (% of initial DM remaining) of (A) green leaf material and (B) leaf litter and roots of four plant species. Values are averaged over CO₂ treatments and are mean of 6 replicates ± s.e.
organic matter returns by grazers. Scenario 4 includes a possible decrease in faeces decomposition rate under elevated CO₂ based on the results observed in this study. To finish with scenario 5 combines scenarios 1, 2 and 4 and simulates overall decomposition rates in a grazed pasture under elevated CO₂. The decomposition rates used in the above scenarios were obtained by fitting average decomposition curves obtained in this experiment to a exponential decay model \( y = y_0 e^{-kt} \), where \( k \) is the decomposition rate.

3.3.7 Statistical analysis

The mass of green material and leaf litter remaining was analysed at each date by analysis of variance (ANOVA) using a split-split-plot model using the following hierarchy: CO₂ incubation as main plot, CO₂ origin as sub-plot and species as sub-sub-plot. The relative N loss of these two materials was analysed with the same model but at a single date. The model for the analysis of root material remaining mass and the root relative N loss was a split-plot with CO₂ incubation as main plot and CO₂ origin as sub-plot. The mass of faeces remaining at each sampling date and the relative N loss, in 2001, were analysed as the root material and sheep were used as replicates in 2000. All analyses were performed in Genstat v 6.1 (Genstat 2002).

3.4 Results

3.4.1 Initial chemical composition

The initial chemical composition of green leaf, litter and root material used in this study did not differ between CO₂ treatments but species strongly affected N concentration of leaf green material and litter (Table 3.1). Green leaves of \( P. \) dilatatum and \( T. \) subterraneum had higher N concentration than \( A. \) odorantum and \( H. \) radicata but the absolute difference was marginal. Leaf litter initial chemical composition was also strongly affected by species type (p< 0.001, Table 3.2) with \( T. \) subterraneum litter having a 2-fold greater N concentration compared to the three other species. \( Trifolium \) subterraneum litter C/N ratio was also significantly lower.
Figure 3.2: Effects of the CO₂ concentration during the production of the material (open shape: ambient and black bars: elevated) and during decomposition (circle: ambient and triangle: elevated) on the mass loss (% of remaining mass) of (A) faeces during summer and (B) during winter. Values are means of 3 replicates ± s.e.
compared to the other species. The average N concentration of the faeces used in the summer experiment tended to be lower under elevated CO$_2$ (-11%, p=0.087) and the non significance of this result was due to one sheep (p=0.005 when it is excluded). In contrast, the average N concentration of the faeces samples used in the winter experiment was unaffected by elevated CO$_2$, though it was about 30% lower than in the first experiment.

3.4.2. Decomposition rates

Decomposition rates of green leaf material and litter, as measured by DM loss from the incubation bags, were not significantly affected by the CO$_2$ origin treatment, or by the CO$_2$ incubation treatment. Decomposition rates of both green material and leaf litter were affected by species, however. At all sampling dates, the species effect on remaining mass was extremely significant for both types of material. After 6 months of incubation, _T. subterraneum_ and _H. radicata_ green leaves were nearly totally decomposed while about 5% of _A. odoratum_ and 11% of _P. dilatatum_ material remained (Fig. 3.1a). Litter material exhibited a similar species effect with _H. radicata_ and _T. subterraneum_ having the smallest remaining mass after 6 months of incubation (11% and 21% respectively) while about 50% of the grass litter still remained (Fig. 3.1b).

Although roots could not be separated into plant species, it was still of interest to compare rates of root and green or dead leaf decomposition. Again, neither the CO$_2$ origin treatment, nor the CO$_2$ incubation treatment altered root decomposition rates. However, decomposition of root material and green or dead leaf material followed different time courses (Fig. 3.1b). Decomposition rate of root material was initially similar to that of the litter of the fast decomposing species _T. subterraneum_ and _H. radicata_, but little change was observed over the last 4 months of incubation. About 25% of the initial root mass remained after 8 months, compared with 4% for dicot species litter and 40% for grass species litter.

The two measurements of faeces decomposition took place under contrasting climatic conditions. In the first experiment, conditions were dry, with volumetric soil
Figure 3.3: Effects of the CO₂ concentration during the production of the material (black bars: ambient and open bars: elevated) and during decomposition (empty bar: ambient and lined bar: elevated) on the relative N loss (remaining N / remaining mass) after 120 days of incubation of (A) green leaves and (B) leaf litter of four species (*Trifolium subterraneum* (Tri), *Hypochaeris radicata* (Hyra) and *Paspalum dilatatum* (Pasp) and *Anthoxantum odorantum* (Anth)) and (C) root material. Values are mean of three replicates ± s.e.

Figure 3.4: Effects of the CO₂ concentration during the production of the material (black bars: ambient and open bars: elevated) and during decomposition (empty bar: ambient and lined bar: elevated) on the relative N loss (remaining N / remaining mass) after 50 days of (A) faeces during summer and (B) faeces during winter.
moisture content to 15 cm depth averaging about 10%. In the second experiment, conducted in winter, soil moisture content averaged about 25%. Under the dry summer conditions remaining mass after 16 weeks was about 70% of the initial mass; during winter remaining mass was only 20% of the initial after 12 weeks (Fig. 3.2). In summer, the CO₂ origin dramatically altered decomposition rate (p<0.001 at all dates). Faeces produced under elevated CO₂ decomposed more slowly; with about 20% more of the initial mass remaining at a given sampling date (Fig. 3.2). Conversely, CO₂ concentration during incubation did not affect decomposition rates. In winter faeces decomposition rates were unaltered by any of the treatments (Fig 3.2).

3.4.3. N release during decomposition

The relative N loss of the different plant materials was calculated for the second retrieval date, after 4 months of incubation; no significant effect of the CO₂ origin or CO₂ incubation treatments was evident (Fig. 3.3). However, relative N loss of root material was about 1.1 indicating similar DM and N decomposition pattern whereas relative N loss for leaf litter of some species exceeded 1.5 at the same date (Fig. 3.3), indicating relatively slower N release during leaf litter decomposition.

Relative N loss for faecal material of the first experiment was calculated after 16 weeks, and after 6 weeks in the second experiment, since with faster decomposition rates in the second experiment, insufficient material remained at the last sampling date for analysis. In contrast to plant green material or litter, relative N loss of faecal material was lower than 1 (Fig. 3.4), being 0.95 and 0.88 for the summer and the winter experiments, respectively.
Table 3.3: Simulation of the effect of elevated CO₂ on the mass loss rates and N release rates at the pasture scale under different scenarios. Mass loss values are derived from individual material decomposition curves after fitting to an exponential decay model and averaged to take into account the relative importance of the different materials. N release values are calculated from mass loss values and relative N release values measured in this study. Scenario 1 simulates an increase in dicots in the pasture; scenario 2 simulates increase in root production and scenario 4 a decrease in faeces decomposition; scenarios 3 and 5 are combination of the latter.

<table>
<thead>
<tr>
<th>Scenario</th>
<th>Average mass loss [CO₂]</th>
<th>Average N release rate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ambient</td>
<td>Elevated</td>
</tr>
<tr>
<td>1) Change in botanical composition¹</td>
<td>5.50 10⁻³</td>
<td>6.10 10⁻³</td>
</tr>
<tr>
<td>2) Change in allocation to root growth²</td>
<td>5.50 10⁻³</td>
<td>5.70 10⁻³</td>
</tr>
<tr>
<td>3) Cut grassland¹,²</td>
<td>5.50 10⁻³</td>
<td>6.30 10⁻³</td>
</tr>
<tr>
<td>4) Decrease in faeces decomposition rates³</td>
<td>6.90 10⁻³</td>
<td>6.80 10⁻³</td>
</tr>
<tr>
<td>5) Grazed grassland¹,²,³</td>
<td>6.90 10⁻³</td>
<td>7.20 10⁻³</td>
</tr>
</tbody>
</table>

¹Dicots participation to herbage biomass is 15% and 30% under ambient and elevated CO₂, respectively.
²Root production is 30 and 50% of the herbage under ambient and elevated CO₂, respectively.
³Herbage utilisation rate is 50% of above ground production and herbage digestibility is 0.74.
3.5. Discussion

3.5.1. Effects of species changes under elevated CO$_2$ on leaf litter decomposition

Our results confirm earlier findings that elevated CO$_2$ does not alter the C/N ratio or decomposition rates of plant litter from individual species (Norby and Cotrufo, 1998; Ross et al., 2002). These previous studies concluded that indirect CO$_2$ effects on botanical composition and shifts in biomass allocation may be of greater importance with regards to the litter decomposition at the ecosystem scale. This view is consistent with our results. None of the three vegetal materials (green leaf, litter and root) used in this experiment exhibited higher C/N ratio or lower decomposition rates when produced under elevated CO$_2$ but species-specific differences were observed. Leaf litter from *T. subterraneum* and *H. radicata* decomposed more than two times faster than the litter of the two grasses included in this study. Cornelissen (1996) also showed significantly greater decomposition rates of dicots compared to grasses. In view of the substantial increase in the proportion of legumes and forbs under elevated CO$_2$ at this site (Edwards et al., 2001), it is clear from these results that elevated CO$_2$ must be associated with an overall increase in the decomposition rate of leaf litter at the pasture scale at this FACE site. Quantitatively, under the assumption of a 2-fold increase in the proportion of dicots under elevated CO$_2$ (Scenario 1), which is consistent with long term data from this site (P. Newton, unpublished data), the leaf litter mass loss and N release rates would increase by 10% at the sward scale (Table 3.3). Such an increase in decomposition rates driven by CO$_2$-induced shifts in botanical composition is probably not specific to this site. In temperate grassland an increasing abundance of dicots and legumes in particular, is a common response to elevated CO$_2$ and has been observed under a wide range of management conditions: in ryegrass/clover associations (Hebeisen et al., 1997; Jongen and Jones, 1998), natural grassland (Teyssonneyre et al., 2002) and in grazed temperate grassland (Newton et al., 2001; Edwards et al., 2001). In addition, not only does a high legume content induce a higher proportion of more rapidly decomposing litter but may also increase the average N concentration of the non-legume species and thus contribute to a faster decomposition of the latter (Hartwig et al., 2000).
The observed increase in the proportion of "fast-decomposing species" under elevated CO$_2$ leads to the question of whether the decomposition rate is a functional trait directly or indirectly selected for under elevated CO$_2$. Regardless of the response to elevated CO$_2$, low litter decomposability is linked with low leaf palatability (Cornelissen and Thomson, 1997; Moretto et al., 2001). Plants exhibiting low leaf palatability, a strategy used to limit or avoid defoliation by potential grazers, require large C investments in defence compounds. This decreases the amount of C able to be allocated in photosynthetically active tissues and therefore palatability is positively correlated with growth rate (Coley, 1988). When the plant response to elevated CO$_2$ is considered, there is much evidence that fast growing species (i.e. with a high growth rate) show stronger growth response to elevated CO$_2$ than slow growing species (Poorter, 1993; Poorter et al., 1996; Poorter and Navas, 2003). This leads to the conclusion that elevated CO$_2$ may be a selective agent for species with high decomposition rates. Berendse (1994) highlighted the importance of decomposition rates as a functional trait, suggesting that litter decomposition and the subsequent capacity to release nutrients was an active component of plant fitness. Indeed, plants appear to affect soil fertility in such a way that with time the substrate becomes more favourable for growth (Van Breemen, 1993). Under elevated CO$_2$, fast nutrient cycling may be an interesting feature because the higher C availability increases the relative need for other nutrients. Nevertheless, more comprehensive studies are needed to draw general conclusions about the importance of litter decomposition rate as an important trait determining species response to elevated CO$_2$.

3.5.2. Effects of CO$_2$-induced shift in biomass allocation on decomposition rates

Another level at which elevated CO$_2$ may alter decomposition rates at the ecosystem scale is through changes in biomass allocation and in particular a proportionally greater biomass allocation belowground. In another experiment on the same site it was shown that root growth and turnover were greatly increased under elevated CO$_2$ in summer and spring (Chapter 4) whereas aboveground herbage production was not affected. Strong CO$_2$ stimulation of root turnover in grasslands has been found elsewhere (Canadell et al., 1996; Fitter et al., 1996). In this study root decomposition was not found to differ greatly from leaf litter decomposition, at least.
during the first phase of decomposition (accounting for about two thirds of the initial mass). Our results suggest that roots might have a more recalcitrant fraction which induces low decomposition rates in the later stages of the process but it still seems that overall root decomposition is not intrinsically slower than leaf litter decomposition.

This contrasts with other studies reporting much slower root decomposition when compared to aboveground material. For example Gorissen and Cotrufo (2000) observed that roots of three grass species exhibited between 25% and 50% less mass loss than aboveground material after about 7 months of incubation under laboratory conditions. Nevertheless we consider that this overestimates dramatically the differences in decomposition rates between the two types of material for two reasons. Firstly the use of green leaves in this decomposition experiment artificially increases the contrast with root decomposition rate since green material decomposes much faster than true leaf litter (Fig. 3.1). In comparison, decomposition rates of live and dead root material may differ less than those of live and dead leaves because rather less translocation of substrates from roots occurs during root senescence (Gordon and Jackson, 2000). Secondly, previous studies clearly demonstrated the effects of litter position on decomposition rates. Dukes and Field (2000) showed that leaf litter decomposes faster when buried rather than placed at the soil surface. Therefore the comparison of leaf and root material decomposition under standard conditions does not answer the central question of the relative decomposition rate of root and leaf litter decomposition in situ, i.e. aboveground for leaf litter and belowground for senesced roots. Again using a simulation scenario based on data from this site, a CO₂-driven shift towards an increased proportion of root litter in the total litter return will not lead to decreased decomposition rates at the ecosystem scale (Scenario 2). On the contrary, we calculate it would lead to a small increase (about 4%) in the average decomposition rate (Table 3.3). In addition, given the low N immobilisation potential of decomposing root material compared to leaf litter (Fig. 3.2; Seastedt et al., 1992), there should be a further increase soil N availability (6.5%). Combining the CO₂ effects on botanical composition and shoot:root allocation from Scenario 3 we calculate that at elevated CO₂ there will be a 15% increase in mass loss and 17% increase in N mineralisation rate from litter (Table 3.3).
3.5.3. Effects of elevated CO$_2$ on ruminant faeces decomposition

In well managed grazed grasslands, up to 50% of the aboveground herbage production is ingested by herbivores (Parsons and Chapman, 2000) therefore an important part of the organic matter cycling occurs through urine and faeces returns (Parsons et al., 1991; Orr et al., 1995). In this study sheep faeces decomposition exhibited a strong negative CO$_2$ effect during the first of the two decomposition experiments (Fig. 3.2). Because animals eat green leaves and not senescing material, and because CO$_2$ effects on leaf chemical composition are frequently observed in green material, it might be anticipated that herbivore diets may be altered under elevated CO$_2$. Furthermore, through the grazing process, ruminants integrate both the CO$_2$-driven decrease in N concentration observed at the single plant scale and the changes in botanical composition; in this case a higher proportion of legumes and forbs. In a previous experiment, we observed that these two processes were counterbalanced, leading to a similar N intake by animals under ambient and elevated CO$_2$ (Chapter 2). Nevertheless elevated CO$_2$ altered the partitioning of N return between urine and faeces, resulting in a decrease in N in animal faeces, attributable to the increased digestibility and higher proportion of legumes under elevated CO$_2$ conditions (Chapter 2).

Nevertheless, the decrease in faeces decomposition rate was only observed in the summer experiment. Under dry conditions, a thick crust forms over the surface of excreted faeces (Holter, 1979) that dramatically limits potential exchanges with the soil surface. Furthermore, under these dry conditions the soil fauna, in particular the earthworm population that plays a major role in dung decomposition (Hirschberger and Bauer, 1994), is mainly inactive, at least in the top soil (Yeates, 1976). The two most common earthworm species in New Zealand pastures are *Allobophora calliginosa* and *Lumbricus rubellus* and both of them develop inactive forms during the dry season. Decomposition that does occur in drier summer conditions is most likely to be caused by microbial and/or fungal activity and thus could be expected to be affected by variation in the initial C/N ratio of the material. By contrast, during winter, constant rainfall facilitates physical breakdown of the faeces and a fully active invertebrate population leads to fast decomposition. It seems likely that under these conditions, decomposition rate would be less dependent on initial quality of the
material. In a model ecosystem where dung decomposition is assumed to be the only process affected by elevated CO₂ (Scenario 4) both the ecosystem decomposition rate and the N mineralisation rate would be slightly decreased (by 2 and 5% respectively, Table 3.3). When all possible effects of elevated CO₂ are combined (Scenario 5) overall average decomposition rates are not altered by elevated CO₂ (Table 3.3).

3.6. Conclusion

This study confirmed that direct effects of elevated CO₂ on litter quality at the single plant scale could be discounted as a potential mechanism affecting organic matter decomposition in grasslands. However, secondary effects of elevated CO₂ at the community level through an increase in proportion of the legumes and other dicots, tended to increase litter decomposition rate. When organic matter return through herbivore faeces is not considered, these indirect CO₂ effects would lead to an increase of about 15% in decomposition rates. Under grazing, elevated CO₂ did not affect estimated decomposition rate. But importantly, grazing increased decomposition rates by more than 15% and N release rates by 60% when compared with grasslands without animal returns, probably enhancing potential N losses in particular under elevated CO₂ (Chapter 2). As a consequence, it appears that the effects of elevated CO₂ on decomposition processes in grazed grasslands act at different ecosystem scales that need to be included in future studies in order to predict more accurately whether the organic matter in grassland soils will accumulate or not under increasing levels of atmospheric CO₂.

3.7. Acknowledgements

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3.8. References


Gorissen A, Cotrufo MF (2000) Decomposition of leaf and root tissue of three perennial grass species grown at two levels of atmospheric CO$_2$ and N supply. *Plant and Soil, 224*, 75-84.


Chapter 4: Increased Quantity and Quality of Coarse Soil Organic Matter Fraction at Elevated CO$_2$ in a Grazed Grassland Are a Consequence of Enhanced Root Growth and Turnover

Plate 4.1: View of an "ingrowth core" pipe buried in the field with a 45° angle
Chapter 4: Increased Quantity and Quality of Coarse Soil Organic Matter Fraction at Elevated CO$_2$ in a Grazed Grassland are a Consequence of Enhanced Root Growth and Turnover

Allard V, Newton PCD, Soussana, J-F., Carran, RA, Matthew C, Lieffering M.

4.1. Abstract

The aims of this study were, firstly to determine whether elevated atmospheric CO$_2$ modified plant organic matter (OM) fluxes to the soil and secondly to assess whether any changes in the fluxes affected OM accumulation in the soil. A grazed temperate grassland was exposed for four years to elevated atmospheric CO$_2$ (475 µl l$^{-1}$) using a Free Air CO$_2$ Enrichment (FACE) facility. Neither aboveground herbage biomass nor leaf litter production were affected by elevated CO$_2$. In comparison, roots growth rate and turnover were clearly stimulated by elevated CO$_2$ in particular under relatively dry soil conditions. Consequently, significantly more plant material was returned to the soil under elevated CO$_2$. This was translated into an accumulation of coarse (> 1 mm) particulate organic matter (POM) under elevated CO$_2$ but not of finer POM fractions. In addition, the accumulating POM exhibited a lower C/N ratio, which was attributed to the higher proportion of legumes in the pasture under elevated CO$_2$. Only small changes were detected in the size and activity of the soil microbial biomass in response to the POM accumulation, suggesting that higher organic substrate availability did not stimulate microbial growth and activity despite the apparent higher quality of accumulating POM. As a result, elevated CO$_2$ may well lead to an accumulation of OM in grazed grassland soil in the long term.
4.2. Introduction

As the concentration of CO$_2$ in the atmosphere increases (Keeling et al., 1995), attention has focused on the possibility to increase carbon (C) storage capacity on land. In this context, grassland responses to elevated CO$_2$ are of particular importance since grasslands account for 25% of the terrestrial land surface and 10% of global C stores (Schimel, 1995). Unlike tropical forests, where vegetation is the primary source of C storage, most of the grassland C stocks are in the soil. Therefore possible accumulation of OM in grasslands soils under elevated CO$_2$ may be of central importance in mitigating atmospheric CO$_2$ increase (Scurlock and Hall, 1998).

At least three processes have been identified as mechanisms that could lead to increased grasslands soil C storage under elevated CO$_2$. First, much attention has been paid to the potential alteration of litter decomposition rates that were expected to be lower under elevated CO$_2$ (Strain and Bazzaz, 1983). Nitrogen (N) concentration often decreases in plants that have been grown under elevated CO$_2$ (Cotrufo et al., 1998; Chapter 3), but due to resorption processes during senescence, a decrease in N concentration is generally not observed in plant litter (Norby and Cotrufo, 1998). Consequently, litter quality and hence litter decomposition per se are commonly unaffected by elevated CO$_2$ (Hirschel et al., 1997; van Vuuren et al., 2000).

Second, increased root exudation of easily decomposable C has often been reported in elevated CO$_2$ studies (Paterson et al., 1996; van Ginkel et al., 2000). Exudation is difficult to quantify but estimates range from 10 to 40% of net assimilated C in annual crops (Whipps, 1984; van Veen et al., 1991). Because the highly fermentable C of root exudates generates a high microbial activity in the rhizosphere (Hale and Moore, 1979), even small increases in root exudate quantity may alter soil OM dynamics through changes in microbial trophic preferences. It has been suggested (Lekkerkerk et al., 1990) that microorganisms may turn to using more easily fermentable C substrate from roots exudates when this is available, rather than more recalcitrant old OM (Cardon et al., 2001; van Veen et al., 1991).

Third, in addition to potential modifications of OM quality, elevated CO$_2$ may also induce greater C fluxes from the growing plants to the soil through leaf (Navas et
al., 1995; Niklaus et al., 2001) and root litter deposition in particular. Belowground plant-soil fluxes have indeed been observed to exhibit stronger responses to elevated CO₂ as shown by lower shoot:root ratio (Cotrufo and Gorissen, 1997; van Ginkel et al., 1997) and increased root turnover (Canadell et al., 1996; Fitter et al., 1996) although these results are not ubiquitous (Niklaus et al., 2001).

Because of the large background levels of organic C in grassland soils, changes in the size of soil C compartments are difficult to detect through direct measurements (Ross et al., 1996). The coarse particulate POM compartment (>1 mm) represents the first stage in a continuum of decomposition in grassland ecosystems that culminates in stabilised soil OM (Mellilo et al., 1989) consequently we might expect to see any changes in response to elevated CO₂ in short term experiments (relative to the duration of the complete decomposition process) expressed in the coarse POM pool. A method for separating newly deposited OM compartments is therefore needed to study belowground OM accumulation under elevated CO₂.

In this study we evaluate changes in POM pools and their possible origin by measuring aboveground herbage biomass, leaf litter and root C fluxes in a grassland grazed by large ruminants and exposed to elevated CO₂ (475 μl l⁻¹) for about four years. We are not aware of other studies that have measured the effects of elevated CO₂ on these fluxes under grazing despite the fact that grazing is the most common management for grasslands globally and that there is an expectation that grazed systems will respond differently to elevated CO₂ from those managed by infrequent cutting (Newton et al., 2001). In particular, the frequent removal of herbage aboveground biomass by the ruminants may affect leaf litter production and its interactions with elevated CO₂ as well as root growth and turnover (Matthew et al., 1991).
4.3. Materials and methods

4.3.1. Experimental site

The study was carried out between December 2000 and November 2001 in a temperate pasture on the west coast of the North Island of New Zealand (40°14'S, 175°16'E). The pasture had been under permanent grazing by sheep, cattle and goats since at least 1940. The mean annual rainfall at the site was 875 mm. More details about the site botanical and physical characteristics can be found in Edwards et al. (2001). The experimental facility consisted of six FACE (Free Air CO2 Enrichment) rings (McLeod and Long, 1999) each 12 m in diameter; the rings were paired into three blocks based on initial soil and botanical characteristics. In each block, one ring was fumigated with CO2 to a target value of 475 µl l−1 CO2 (elevated CO2 treatment) during the photoperiod, the second being left at ambient atmospheric CO2 concentration (ambient CO2 treatment). Enrichment began on 1 October 1997.

4.3.2. Experimental areas

All plant measurements were made during two separate periods of four months in 2000-2001. The first period started in December 2000 and ended in May 2001 (summer-autumn) and the second period extended between July and November 2001 (winter-spring). In December 2000, 12 (2 × 6 species) circular areas (Ø=200 mm) were selected in each ring based on the dominance of each of these six plant species: *Lolium perenne* L., *Agrostis capillaris* L., *Anthoxanthum odoratum* L. (all C3 grasses), *Trifolium repens* L. (legume), *Paspalum dilatatum* Poir. (C4 grass) and *Hypochoeris radicata* L. (forb). New circular areas were selected in the ambient and elevated CO2 rings in June 2001 for the second period. Because *T. repens* was absent from the rings in the second period the selected legume was *Trifolium subterraneum* but all the other species remained the same. For both experiments, the selection of the experimental areas was immediately followed by a grazing of the rings by adult sheep over a period of three days. Similarly, two months after the beginning of each experiment, the rings were grazed again. During both periods the botanical
Figure 4.1: Schematic representation of an "ingrowth core" pipe
composition of some of the sampling areas changed over time. No species effect
could thus be tested for and the botanical composition in the sampled areas was
considered to be broadly representative of the entire rings. Indeed no significant
differences were observed between the botanical composition of the whole rings and
of the combined area made of the 12 experimental areas (data not shown).

4.3.3. Root measurements

In both experiments, after the initial grazing, soil cores (Ø=35 mm,
length=250 mm), were taken at a 45° angle from the periphery of each sample area
towards its centre. Roots were removed from the cores by sieving (mesh size=1 mm),
oven dried (60°C, to constant mass) and weighed to provide an estimate of the root
standing biomass at the beginning of the experiment. Root growth rate was assessed
using an ingrowth core technique. In each hole resulting from the initial soil coring, a
PVC pipe (Ø=35 mm, length=250 mm) was installed. Windows allowing roots
penetration were made by removing 150 mm longitudinal sections of PVC from the
middle of each pipe leaving only two 1 cm wide PVC strips to preserve the structure
(Fig. 4.1). This design was used to allow precise re-positioning during the following
samplings and to facilitate core removal. The pipes were then filled immediately with
sieved soil originating from the same ring and from which roots and debris had been
removed using a 1 mm sieve. Soil compaction during refilling was standardized by
applying the same filling methods at all samplings. At harvest times (experiment 1: 25
January, 20 February, 27 March and 1 May 2001; experiment 2: 3 August, 30 August,
5 October and 5 November 2001.), the PVC pipes and the soil they contained were
removed with a metal tube with a diameter slightly larger than the PVC pipes. Soil
and roots were extracted and the pipe was immediately replaced and refilled with soil
free of roots. The root samples were cleaned on a sieve (mesh size = 1 mm), oven
dried (60°C, to constant mass) and weighed to give the value of root growth during
the burial period. Root turnover was estimated by dividing the initial standing root
biomass by the average of root production over the period (four successive
samplings). We are aware that this estimate is strongly dependent on the initial
standing root biomass value that could vary over the four months of the experiment.
Therefore this estimate of turnover should be regarded as a tool to compare the CO₂ treatments and the actual values should be considered with care.

4.3.4. Aboveground measurements

After two and four months of the root growth measurement period, aboveground biomass in the same areas as that used for root growth, was cut to 2 cm above ground level with powered hand shears. A sub-sample was taken for dissection into plant species and dry weights of these components and of the total sample were taken after drying to constant mass at 60°C. During the summer-autumn measurement period, dead leaf material deposited on the soil surface was regularly removed in the selected 200 mm diameter areas using a vacuum cleaner. Material stuck in dense vegetation patches was removed by hand. The first litter sampling occurred on 1 March then on four occasions at intervals of 15 days. The material removed was separated from adherent soil, oven dried (60°C, to constant mass) and weighed. The first sampling was used as an estimate of standing litter biomass and the subsequent samplings used to calculate the average leaf litter deposition rate.

4.3.5. Soil compartments analysis

In November 2001, at the end of the second measurement period, six soil cores (Ø=90 mm, depth=250 mm were sampled from each FACE ring. Aboveground vegetation and litter were removed and the cores were shipped to France (INRA Clermont-Ferrand, Agronomy Unit) and kept in a cool room (4°C) before further analysis. In January 2002, the soil cores were cut longitudinally into equal quarters. One of these quarters was sieved (mesh size=1mm) and remaining roots and coarse litter fractions carefully removed. Sieved soil was used for microbial biomass (MBM) and extractible organic matter (EOM) measurements (see below) and total soil C and N concentrations. From the remaining three quarters of undisturbed soil, two POM fractions ("coarse": >1mm and "fine": 0.2-1mm) were obtained by wet sieving. The soil samples (about 1kg) were rotationally shaken in water in a plastic cylinder.
Figure 4.2: Maximum and minimum air temperature (°C) (dotted lines) and weekly rainfall (mm week$^{-1}$) (bars) at the experimental site over the course of the experiment. The two horizontal dotted lines figure the two periods of root growth measurements.

Figure 4.3: Volumetric soil water content in the experimental plots (% moisture) under ambient (solid circles) and elevated CO$_2$ (open circles) over the course of the experiment. Data are mean of three replicates per treatment.
After stirring for 30 min, the samples were sieved through mesh sizes of 1, 0.2 and 0.05 mm. The residues over 1 mm were collected on the first sieve and dispersed in water to eliminate any adherent soil. The finer part of the POM (0.2-1 mm) collected on the second sieve was separated from sand particles by densimetry in water. The smallest POM fraction (0.05-0.2 mm) was also separated from dense particles in water but the difficulty of distinguishing between POM and the pure mineral fraction forced us to exclude this compartment from the study. The POM fractions were then oven dried (75°C, to constant mass) and finely ground using a ball mill (Retsch, Germany). The elemental C and N concentration of the two POM fractions and of total soil were determined using an elemental analyser (Carlo Erba, Italy) in INRA-Nancy, France.

Microbial biomass was extracted following the fumigation-extraction procedure described by Vance et al. (1987). Four replicates per soil sample were fumigated and two were unfumigated to compensate for the higher variability of C and N contents in fumigated extracts (G. Alvarez, personal communication). Organic C and N in the extracts were analysed with an automated TOC/TN analyser (Skalar Formacs, Netherlands). Microbial C biomass was calculated as $C_{mic} = (\text{C}_{fumigated} - \text{C}_{unfumigated}) / k_{EC}$, where $k_{EC}$ is the extraction yield for microbial C ($k_{EC} = 0.42$, Ross and Tate, 1993). Microbial N biomass was calculated as $N_{mic} = (\text{N}_{fumigated} - \text{N}_{unfumigated}) / k_{EN}$, where $k_{EN}$ is the extraction yield for microbial N ($k_{EN} = 0.45$). These coefficients are not affected by elevated CO₂ (Ross et al., 1993). EOM was extracted from the soil samples by autoclaving the equivalent of 10 g dry soil using a modification of the method described in Lemaitre et al. (1995) (P. Loiseau, personal communication). The soil was first mixed with 40 ml distilled water for 1 min and then autoclaved (120°C, 2h). The extracts were filtered (Whatman No. 42 paper) before being analysed for organic C and N with an automated TOC/TN analyser (Skalar Formacs, Netherlands).
Figure 4.4: Aboveground biomass (g DM m$^{-2}$) under ambient (solid bars) and elevated CO$_2$ (open bars) at the 4 successive sampling dates. Results are mean of three replicates per treatment ± standard deviation.

Table 4.1: Mean leaf litter standing biomass, deposition rate and residence time under ambient and elevated CO$_2$. Leaf litter deposition rates are means of 4 successive sampling times. Standard deviations are indicated.

<table>
<thead>
<tr>
<th></th>
<th>Ambient CO$_2$</th>
<th>Elevated CO$_2$</th>
<th>CO$_2$ effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leaf litter standing biomass (g DM m$^{-2}$)</td>
<td>154.2±28.3</td>
<td>126.6±8.3</td>
<td>$p=0.508$</td>
</tr>
<tr>
<td>Leaf litter deposition rate (g DM m$^{-2}$ day$^{-1}$)</td>
<td>3±0.4</td>
<td>2.1±0.2</td>
<td>$p=0.116$</td>
</tr>
<tr>
<td>Residence time (day)</td>
<td>49.8±3.2</td>
<td>61±9.1</td>
<td>$p=0.42$</td>
</tr>
<tr>
<td></td>
<td>(standing biomass / deposition rate)</td>
<td></td>
<td></td>
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</table>
4.3.6. Soil water content

Volumetric soil water content in the rings was measured weekly by time domain reflectometry (TDR) (Trase System, Soil Moisture Equipment Corporation, Santa Barbara California) using 15 cm probes. Mean values were calculated from 16 measurements taken in each FACE ring using a stratified random sampling method with four readings taken in each quadrant.

4.3.7. Statistical analysis

Analysis of variance (ANOVA) in Genstat (Genstat, 2002) was used to compare CO₂ effects on the different measured variables. The model used was a randomized block with three replicates (rings) per CO₂ treatment. Sub-samples within each ring were averaged prior to analysis. Root growth data were analysed using a repeated measures model using the four successive sampling of each experiment as repeated measures. Similarly, soil water content was analysed as repeated measures. Data that did not satisfy the homogeneity of variances assumption required for an ANOVA were log transformed prior to the analysis.

4.4. Results

4.4.1. Climate and soil moisture

The first period of root growth measurements (18 December-1 May) was characterised by typical weather for this part of the year with 210 mm of rainfall over the 4 months of study and an average daily maximum temperature of 21.5°C (Fig. 4.2). The second experimental period (2 July- 5 November) had an average daily maximum temperature of 15.3°C and total rainfall over the period was 488 mm (Fig. 4.2) which was considerably higher than the long-term (50 years) average of 372 mm. Volumetric soil moisture content was measured weekly in the rings (Fig. 4.3) and was not affected by elevated CO₂ (p=0.507). The first part of the root growth experiment
Table 4.2: Root standing biomass, growth and residence time under ambient and elevated CO₂ for the two measurement periods. Results are means of three replicates per treatment ± standard deviation except for root growth results that are mean of 4 successive sampling and three replicates per sampling. P values are indicated.

<table>
<thead>
<tr>
<th></th>
<th>Period 1 (19 Dec - 1 May)</th>
<th>Period 2 (2 Jul - 5 Nov)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ambient CO₂</td>
<td>Elevated CO₂</td>
</tr>
<tr>
<td>Root standing biomass</td>
<td>122.1±17.5</td>
<td>81.4±23.5</td>
</tr>
<tr>
<td>(g DM m⁻²)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Root growth rate</td>
<td>0.9±0.2</td>
<td>1.4±0.1</td>
</tr>
<tr>
<td>(g DM m⁻² day⁻¹)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Root residence time</td>
<td>173.6±20.4</td>
<td>62.5±18.7</td>
</tr>
<tr>
<td>(day) (standing biomass/ growth)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 4.5: Root growth rate (g DM m⁻² day⁻¹) under ambient (solid circle) and elevated CO₂ (open circle) for the 2 measurement periods. Values are mean of three replicates per treatment ± standard deviation.
Chapter 4

(December-May) took place under low soil water content (9.1% on average) and the second part under relatively high soil water content (25.1% on average) (Fig. 4.3).

4.4.2. Aboveground biomass and litter deposition

Elevated CO₂ did not significantly affect the herbage biomass at any of the four harvests (Fig. 4.4). Leaf litter standing biomass was not significantly different between CO₂ treatments, the average mass of litter being 140 g dry matter (DM) m⁻² (Table 4.1). Also, neither the mean rate of litter deposition nor the estimate of the residence time of the litter was significantly altered by the CO₂ treatment (Table 4.1).

4.4.3. Root standing biomass, growth rate and turnover

Mean root standing biomass tended to be lower under elevated CO₂ at the December 2000 harvest (-33.6%, p=0.06) (Table 4.2) and showed a similar trend in July 2001 (-20.3%) but again it was not statistically significant. The difference between seasons in root standing biomass was very marked, with the average biomass measured in winter-spring being less than half of that measured in summer-autumn. Root growth rate was significantly enhanced by elevated CO₂ during the summer-autumn experiment (Fig. 4.5) (p=0.026) with a consistent CO₂ stimulation averaging 56.8%. During the winter-spring period there was no consistent effect of elevated CO₂; on average, root growth was 10.5% greater over this period, largely due to significantly higher growth in October (Fig. 4.5). As a result of the lower root standing biomass and the higher root growth under elevated CO₂, root residence time was significantly lower under elevated CO₂ during the summer-autumn period (-64%, p<0.001). A similar trend occurred during the winter-spring period (-23%) but this trend was not significant.
Figure 4.6: Mass, C and N concentration, C and N pools per kg of soil and C/N ratio in organic matter fractions and total soil under ambient (solid bars) and elevated CO$_2$ (open bars). Values are mean of three replicates per treatment ± standard deviation. * and ** denote significant CO$_2$ effect at the 0.05 and 0.01 levels respectively.
4.4.4. Soil C and N

The mass of the coarser POM fraction (> 1 mm) isolated by OM fractionation was 24.5% (p=0.024) greater under elevated CO₂ (from 3.1 to 3.9 g DM kg⁻¹ soil) (Fig. 4.6). This fraction had a similar increase in C mass (p=0.031) since the C concentration was not affected by elevated CO₂ (Fig. 4.6). On the other hand the mean N mass increase under elevated CO₂ in this pool was more marked (+45%, p=0.009) due to the 20% greater mean N concentration (p=0.026). Consequently the C/N ratio of the >1mm POM fraction dropped significantly from 37 to 31 g C g⁻¹ N (p=0.033), on average. The smaller POM fraction (0.2-1mm) exhibited similar trends (Fig. 4.6) but none of these were statistically significant. Under elevated CO₂, on average, the mass of this smaller fraction was 14% higher and the C/N ratio 7.9% lower. Total soil C and N was not significantly different under elevated CO₂ although there were trends consistent with the differences in POM pools i.e. increased C mass (+7%) and decreased C/N ratio (-1.7%). The C/N ratios of the soil OM fractions decreased with fraction size (Fig. 4.7).

4.4.5. Soil microbial biomass (MBM) and Extractable Organic Matter (EOM)

Soil microbial C and N were not affected by elevated CO₂ (Table 4.3). Nevertheless a trend showing a slightly lower MBM C/N ratio was observed under elevated CO₂ (p=0.055). Extractable organic C exhibited a significant increase under elevated CO₂ but of limited amplitude (about 3%). Extractible organic N and C/N were not affected by the elevated CO₂ treatment (Table 4.3).

4.5. Discussion

4.5.1. Plant organic matter fluxes

As shown by the lack of a CO₂ effect on herbage biomass and leaf litter deposition four years after the beginning of the CO₂ enrichment (Table 4.1), aboveground OM inputs to the soil were not affected by elevated CO₂ in our study.
Figure 4.7: Effect of atmospheric CO₂ concentration on the C/N ratio of the organic matter fractions and total soil.

Table 4.3: Microbial carbon (MBM-C) nitrogen (MBM-N) and C/N ratio (MBM-C/N) and extractible organic C (EOM-C), N (EOM-N) content and C/N ratio (EOM-C/N) in the top soil (0-25 cm) under ambient and elevated CO₂. Values are means of three replicates per treatment ± standard deviation. P values are indicated.

<table>
<thead>
<tr>
<th></th>
<th>Ambient CO₂</th>
<th>Elevated CO₂</th>
<th>CO₂ effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MBM-C</strong> (mg kg⁻¹ soil)</td>
<td>492.2±52.2</td>
<td>483.4±74.3</td>
<td>p=0.868</td>
</tr>
<tr>
<td><strong>MBM-N</strong> (mg kg⁻¹ soil)</td>
<td>75.1±9.4</td>
<td>82.1±15.1</td>
<td>p=0.346</td>
</tr>
<tr>
<td><strong>MBM-C/N</strong> (gC g⁻¹ N)</td>
<td>6.8±0.7</td>
<td>6.2±0.6</td>
<td>p=0.055</td>
</tr>
<tr>
<td><strong>EOM-C</strong> (mg kg⁻¹ soil)</td>
<td>5322.9±265.3</td>
<td>5491.8±240.6</td>
<td>p=0.046</td>
</tr>
<tr>
<td><strong>EOM-N</strong> (mg kg⁻¹ soil)</td>
<td>637.8±35.7</td>
<td>665.3±28</td>
<td>p=0.220</td>
</tr>
<tr>
<td><strong>EOM-C/N</strong> (g C g⁻¹ N)</td>
<td>8.4±0.3</td>
<td>8.2±0.2</td>
<td>p=0.435</td>
</tr>
</tbody>
</table>
Long-term measurements of aboveground biomass on this experimental site are consistent with this result and show no consistent increase of herbage production under elevated CO$_2$ (Edwards et al., 2001). Furthermore, no difference in tissue turnover (leaf birth and death rates) under elevated CO$_2$ was measured in growth rooms on turves of the same soil type (Clark et al., 1995) and in the FACE rings (H. Clark, unpublished data) under elevated CO$_2$. Elevated CO$_2$ effects on aboveground production in grasslands vary greatly between studies reporting both positive and negative effects (Owensby et al., 1999; Morgan et al., 2001; Shaw et al., 2002) and increases in litter production usually exceeds any increase in standing biomass (Niklaus et al., 2001; Navas et al., 1995). Nevertheless, these previous experiments reporting an increase in leaf litter production under elevated CO$_2$ are based on grasslands with very lenient defoliation regime. The pasture used in the study presented here is submitted to a relatively high utilisation by ruminants and therefore, this probably minimises the potential for litter accumulation.

In contrast to the lack of response of aboveground OM fluxes under elevated CO$_2$, we observed a strong positive response of belowground fluxes under elevated CO$_2$. There have been several descriptions of increased root standing mass in grasslands under elevated CO$_2$ (Casella and Soussana, 1997; Loiseau and Soussana, 1999; Newton et al., 1996) but it is not ubiquitous (Navas et al., 1995; Stöcklin et al., 1998). Larger root standing biomass however, does not necessarily tell us much about the flux of C belowground (Canadell et al., 1996) as it is root turnover that is the major mechanism of C transfer to the soil (van Veen et al., 1991). There are few data on root turnover in grassland soils in the literature and these again do not show a consistent CO$_2$ response. Fitter et al. (1996) observed concomitant higher root birth rates and shorter root longevity under elevated CO$_2$. Fitter et al. (1997) found similar results but only for one of the two soil types included in their study. However, Arnone et al. (2000) reported no detectable change in root production, mortality and root length density under elevated CO$_2$. In our study we observed both a higher root production rate as measured with ingrowth cores (Fig. 4.5) and a lower standing root biomass under elevated CO$_2$ (Table 4.2). Consequently, the estimate of root residence time was dramatically decreased, at least during the summer period of the experiment, implying a markedly higher flux of OM to the soil pool from roots under elevated CO$_2$ (Table 4.2). As stated in the Materials and Methods section the actual values of
root turnover estimates should be considered with care since the standing root biomass value was only measured once and was very probably not representative of root standing biomass all over the course of the experiment.

Amone et al. (2000) hypothesised that growth conditions may interfere with belowground plant response since high response systems were mostly identified in glasshouse or growth chambers whereas in field studies it was difficult to detect any consistent increase in root system size under elevated CO₂. This was clearly not the case here. Increased root production under elevated CO₂ has also been linked with enhanced soil moisture content (Leadley et al., 1999) but this was not the case in our experiment. We found increased growth at elevated CO₂ during the period of low soil moisture but no difference in moisture content between the CO₂ treatments (Fig. 4.3). This effect has been observed previously in turves of the same soil type growing in growth rooms; in this case, root length density was significantly greater under elevated CO₂ when the turves were grown in periods of low soil moisture (Newton et al., 1996). In that study, C assimilation was sustained longer into the moisture deficit period at elevated CO₂ with a clear trend showing increased biomass allocation belowground rather than aboveground (Newton et al., 1996). Other indirect CO₂ effects may be relevant to the increased root turnover observed under elevated CO₂. In particular, Yeates et al. (submitted) have measured significantly greater abundance (4.3-fold) of the root-feeding nematode Longidorus elongatus in the elevated CO₂ soil of the same experiment and the possibility of interactions between elevated CO₂, root herbivory and root turnover need further investigation.

4.5.2. Soil organic matter

After fractionation of the soil OM, we observed a significant increase in the coarser POM fraction at elevated CO₂ and a similar but non-significant trend in the 0.2-1 mm fraction (Fig. 4.6). I suggest that the greater POM is due to the increased fluxes of OM from the plant root pool. In this study, root turnover has been identified in our system as the flux that exhibited the strongest response to elevated CO₂, and consequently, we suggest that the accumulation of soil POM is in large part a direct consequence of increased root turnover. Greater amounts of POM under elevated CO₂
have been reported by Loiseau and Soussana (1999) who found a 40% increase in macro-organic matter in ryegrass swards exposed to two and a half years of CO₂ enrichment and by Gill et al. (2002) who found a positive relationship between POM and CO₂ concentration (over a gradient from 200 to 550 ppm) after three years of enrichment. In contrast, Niklaus et al. (2001) did not observe any change in the POM fraction in a 6 years open-top chamber experiment in a calcareous grassland under elevated CO₂.

Regardless of the CO₂ treatment, the C/N ratio declined with decreasing OM fraction size. This is a general feature in grassland soil (Guggenberger et al., 1994) due to a release of CO₂ during the stabilisation of the soil OM. Of particular note was the decrease in the C/N ratio of the POM we observed under elevated CO₂ (Fig. 4.6). This result is in contradiction with the findings of Loiseau and Soussana (1999) and Gill et al. (2002) who both showed increased C/N ratios at elevated CO₂. We suggest that the reason for this difference in results lies in the multispecific composition of the sward and the potential this provides for secondary effects to occur through changes in botanical composition. In particular, the higher proportion of legumes under elevated CO₂ (Newton et al., 2001) may explain the higher quality of the recent soil POM pools through a higher N content of legume root material compared to non-legume species. The original idea was to study decomposition rates of single species but root material could not be sampled individually, thus this hypothesis could not be explicitly tested.

The CO₂-induced difference in POM C/N ratios was significant in the coarser fraction (>1mm) but not in the small OM fraction (0.2-1mm) and the total soil (Fig. 4.7). That can be expected since the latter is made of older OM and thus probably contains more material produced before the beginning of the CO₂ fumigation. Therefore, over time, we might expect the ratio to decrease further and the CO₂ effects to be transmitted in the smaller fractions as shown in (Fig. 4.7).
4.5.3. Microbial biomass and dynamic

In spite of the higher POM availability, microbial biomass did not increase under elevated CO\textsubscript{2} (Table 4.3). It confirms results from a long-term study on the same site (Ross \textit{et al.} submitted) showing no CO\textsubscript{2} effect on microbial biomass since the beginning of the CO\textsubscript{2} fumigation. Extractible organic matter (EOM), a compartment that is supposed to integrate microbial dynamics over a relatively long period (Lemaitre \textit{et al.}, 1995) was only slightly increased under elevated CO\textsubscript{2}. The lack of a significant response in terms of microbial biomass to the increased availability of substrate for microbial growth under elevated CO\textsubscript{2} suggests that under these experimental conditions, C is not a major limitation to microbial growth. However the C/N ratio of the microbial biomass decreased significantly under elevated CO\textsubscript{2} suggesting possible modifications of the relative abundance of microbial communities in response to altered POM quality and/or quantity.

4.6. Conclusion

Despite a relatively low level of CO\textsubscript{2} enrichment compared to other elevated CO\textsubscript{2} studies (i.e. 475 μL l\textsuperscript{-1}), an accumulation of OM was observed in the coarser POM fraction and it appears that finer pools may likely express similar changes in the longer term. We suggest that the accumulation of coarse POM was a consequence of increased root turnover at elevated CO\textsubscript{2}. It is unclear whether the increase in root turnover was a direct effect of elevated CO\textsubscript{2} on plant functioning or an indirect effect of elevated CO\textsubscript{2} on soil processes such as root herbivory by nematodes. The POM at elevated CO\textsubscript{2} had a lower C/N ratio which I suggest was due to a CO\textsubscript{2}-induced effect on botanical composition.

Our results differ from some other published studies e.g. the lack of a soil moisture effect and no difference in the amount of leaf litter input. In most cases these differences might be expected given the experimental system; in particular, that our experiment was grazed by herbivores and that defoliation was at frequent intervals through the year rather than infrequent as done in several other studies. Another major difference from other studies is the lower C/N ratio of the accumulating POM under
elevated CO₂. A higher C/N ratio of newly deposited OM under elevated CO₂ was hypothesised to be responsible for shifts in microbial trophic preferences towards metabolisation of older soil OM in order to fulfil their nutritional requirements (Cardon et al., 2001). Consequently organic C storage under elevated CO₂ was relatively small since the increase in POM-C storage was buffered by the decrease in older organic C (Gill et al., 2002). Our results suggest that in grazed pastures with high plant species diversity we might expect extra C sequestration in soil OM mainly through an increase of C input rather than decreasing quality of accumulating OM.

4.7. Acknowledgements

Financial support from the New Zealand Foundation for Research, Science and Technology (Contract C10X205) is gratefully acknowledged. V. Allard was funded by a Massey University Doctoral Scholarship. His stay in New Zealand was also supported by a grant from the Scientific and Cultural service of the French Embassy in New Zealand. We thank Y. Gray, S. Brock, S. Dunn, E. Lawrence and T. Rayner for their technical assistance during field work and R. Delpy for his skilful help with OM fractionation. We also want to thank P. Loiseau for useful discussions and H. Clark for comments on the manuscript.

4.8. References


Chapter 4


Chapter 5: Direct and Indirect Effects of Elevated CO₂ on *Lolium Perenne* Carbon Allocation and Nitrogen Yield
Chapter 5: Direct and Indirect Effects of Elevated CO$_2$ on *Lolium perenne*

Carbon Allocation and Nitrogen Yield.

Allard V, Robin C, Soussana J-F, Newton PCD

5.1. Abstract

It is still unclear whether elevated CO$_2$ increases plant root exudation and affects soil microbial biomass by providing a greater quantity of readily decomposable carbon (C). Also the effects of elevated CO$_2$ on the C and nitrogen (N) contained in old soil organic matter pools is unresolved. In this study we compared the short- and long-term effects of elevated CO$_2$ on C and N pool and fluxes by growing ryegrass (*Lolium perenne*) plants on soil monoliths originating from the New Zealand FACE site (ambient and enriched soil) and under low and elevated concentration of atmospheric CO$_2$ in a glasshouse. Using $^{14}$CO$_2$ pulse labelling, we studied the effects of elevated CO$_2$ on C allocation within the plant-soil system. The results showed that under elevated CO$_2$ more root-derived C was found in the soil and in the microbial biomass. The increased availability of substrate enhanced soil microbial growth and acted as a "priming effect", enhancing native soil organic matter decomposition regardless of the mineral N supply. Despite indications of faster N cycling in soil under elevated CO$_2$, N availability stayed unchanged. Soil from elevated CO$_2$ rings exhibited higher N cycling rates but again it did not affect plant N uptake. Although there are difficulties in extrapolating glasshouse experiment results to the field, we conclude that the accumulation of coarse organic matter observed in the field under elevated CO$_2$ was probably not created by an imbalance between C and N but was likely to be due to more complex phenomena involving soil mesofauna and/or other nutrients limitations.
5.2. Introduction

Plant-soil interactions in the context of elevated atmospheric CO$_2$ concentration (Keeling et al., 1995) have received much attention during the past decade for two main reasons. First is the question of whether the initial fertilisation response that most ecosystems express under elevated CO$_2$ in the short-term, is sustainable in the long-term or whether other nutrients and N in particular will limit plant response to increased C availability (Diaz et al., 1993). Second is the question of whether the direct effects of elevated CO$_2$ on plants will alter soil C stocks and thus exacerbate or buffer the increase in atmospheric CO$_2$ (Smith et al., 2000). As C and N dynamics are strongly linked in soils through soil organic matter decomposition, both questions depend on potential changes in organic matter fluxes from plants under elevated CO$_2$ and on the fate of old organic matter pools. There have been numerous studies assessing CO$_2$ effects on plant litter production and decomposition both directly through changes in litter quality at the species scale (Cotrufo et al., 1994; Hirschel et al., 1997) or indirectly through modification of botanical composition (Kemp et al., 1994; Chapter 3) and biomass allocation (Gorissen and Cotrufo, 2000; Chapter 3). In the particular context of grasslands, a large proportion of organic matter returns from plant to soil occurs through animal faeces and urine and this “pathway” can also be affected by elevated CO$_2$ (Chapter 2).

Root exudation of easily decomposable C is a third pathway of transferring organic matter from plant to soil. Although exudation is probably minor compared to other fluxes (for details about the methodological difficulties involved in exudation quantification see Todorovic et al., 2001), its chemical nature may be more important for ecosystem functioning than its volume (Cardon, 1996). Indeed, the C contained in root exudates is more easily accessible than polymerised C of old organic matter and has the potential to enhance the growth of soils microorganisms that are usually C limited (Smith and Paul, 1990). In relation to elevated CO$_2$, both the frequently observed increase in root biomass (Rogers et al., 1994; Chapter 4) and the potential increase in the amount of C exuded per se (Lekkerkerk et al., 1990; Cheng and Johnson, 1998) might enhance microbial activity in soils and subsequently modify N availability for plants.
Prior to this study, we observed that elevated CO$_2$ altered some organic matter fluxes from plant to soil in a grazed pasture Free Air CO$_2$ Experiment (FACE), in particular, we measured a significant increase in the amount of root material entering the soil organic matter pool. After five years of CO$_2$ enrichment, this led to an accumulation of coarse organic matter in the soil (Chapter 4). The main objective of this study was to study the potential interactions between altered soil organic matter pools in enriched soil and the short-term effect of elevated CO$_2$ on plant root exudation on soil N availability. We tested whether elevated CO$_2$ increased C exudation by ryegrass roots and if the effect of these short-term CO$_2$ effects on soil microbial activity and old soil organic matter decomposition altered N availability. A cross-over design between soil, from ambient or enriched FACE rings, and atmosphere (plants grown under ambient or enriched CO$_2$ concentration) allowed us to study potential interactions between altered exudation processes and the changes in organic matter content observed in the field.

5.3. Materials and methods

5.3.1. Soil, plant and growth conditions

In November 2001, 48 soil monoliths (9 cm in diameter, 25 cm deep) were taken from the New Zealand grazed pasture FACE facility and kept in individual PVC cylinders. More details about the FACE site characteristics and management can be found in Chapter 1. Eight monoliths were taken from each ambient and elevated CO$_2$ (475 μl CO$_2$ l$^{-1}$) ring. Monoliths originating from ambient rings and elevated CO$_2$ rings are hereafter referred to as S- and S+ respectively. All aboveground vegetation was removed manually before the monoliths were sent by air to France where they were kept in a cold room (4°C) until utilisation. In order to allow good water drainage, 5 cm of soil was removed from the bottom of each monolith and replaced with gravel held in place with plastic mesh.

In April 2002, the soil monoliths were placed in two glasshouses at INRA (Champenoux, France) in which the CO$_2$ level was controlled. One glasshouse had
Figure 5.1: CO₂ concentrations (ambient: black circles and elevated: open circles) and daily average air temperature in the ambient and elevated CO₂ glasshouse over the course of the experiment. The data recording device was out of order during the second week of May but the CO₂ concentration was still regulated at that time.
elevated CO₂ levels (target = 700 ppm), the other was kept at ambient CO₂ concentration. Due to a minor air leak between the two glasshouses the CO₂ level in the ambient chamber was higher than ambient air (460 µl l⁻¹). Nevertheless the difference between the two chambers was steady and on average 230 µl l⁻¹ (Fig. 5.1).

One half of the monoliths were placed in the ambient glasshouse (hereafter referred as A-) and the remaining placed in the enriched glasshouse (A+). In order to exclude any potential ring effect for the soil treatment, care was taken to allocate a similar number of monoliths originating from a given FACE ring to each glasshouse. Three seedlings of *Lolium perenne* (cv. Aragon, Cebeco Handelsraad, NL) were transplanted into each monolith (April 1) and after three weeks (most plants at the three leaf stage). The plants were thinned to leave the most vigorous plant.

5.3.2. Plant management

One half of the monoliths of each of the soil-atmosphere combinations was fertilised with 100 kg N ha⁻¹ (N+) and the other half with 5 kg N ha⁻¹ (N-) in order to simulate plant growth without N fertilisation but nevertheless allowing ¹⁵N labelling. The fertiliser was supplied as ¹⁵NH₄¹⁵NO₃ at 5 and 19 atom% ¹⁵N excess for N+ and N- treatments respectively in three split applications: 25% at the removal of the extra seedlings (May 7) and then 37.5% after 2 weeks (May 23) and 5 weeks (June 18) of growth. All plants were irrigated to provide non-limiting amounts of water. They were watered from the top of the monolith according to the climatic conditions and plant size (twice weekly at the beginning of the experiment and twice daily during the last two weeks) in order to keep the soil surface moist and water present at the bottom of the monoliths. The temperature in the chambers was constantly monitored and no difference was measured between the two chambers (Fig. 5.1).

5.3.3. ¹⁴C plant labelling

The day before labelling the plants were removed from the CO₂ controlled glasshouse and brought to ENSAIA, Nancy where the ¹⁴C labelling facility was
Figure 5.2: Schematic view of a plant before $^{14}$C labelling. The plant in its PVC pot is placed within a PVC container closed with a PVC lip. The space between the lip central hole and plant stems was filled with hydrophilic cotton and air tightness was insured by silicone.
installed. Fig. 5.2 describes the preparation of the labelling containers: a plastic tube (3 cm diameter, 4 cm length) was installed at the base of the plant and was filled with liquid silicone (RTV 585, Rhône Poulenc, France) to ensure an airtight seal between the shoot and the root-soil compartment. Each plant with its pot was then placed inside a PVC cylindrical container (12 cm diameter, 40 cm length) with a PVC lip that was screwed at the top of the container. The space between the plastic tube surrounding the base of the plant and the lip was filled with silicone as above. The air tightness of the containers was checked by immersion in water. Because the labelling had to take place under an atmospheric CO₂ concentration similar to the one during plant growth in order not to disturb the A treatment, A+ and A- plants could not be labelled simultaneously. The labelling of the plants took place on June 27 (A-) and July 1 (A+). The day before the labelling, the 24 plants from each of the CO₂ treatments with their containers were put in the assimilation chamber where temperature and light were controlled. Labelling began six hours after the beginning of the photoperiod. For labelling, the ^1₄CO₂ was generated in a flask outside of the chamber by addition of acid to NaH[^1₄]CO₃; respectively 780 and 1410 μCi of ^1₄C were assimilated during the A- and A+ labelling. After two hours of exposure to the labelled CO₂, the remaining ^1₄C was removed from the chamber and the plants were kept in the chamber for a further 48 hours. During this period the soil compartment of each pot was constantly flushed with CO₂ free air and the output air was trapped in NaOH (2M, 200 ml) traps in order to quantify rhizomicrobial respiration.

5.3.4. Plant and soil sampling

Plants were sampled 48 hours after the beginning of the ^1₄C labelling. Aboveground tissues were separated into laminae and sheaths and were immediately dipped into liquid N and stored at -30°C until freeze drying. After removal from the PVC pot, the soil core was carefully broken and all the soil that was not adhering to the root system was considered as bulk soil. The root system was gently shaken and the soil that was still adhering was considered as rhizospheric soil. Sub-samples of both soil types were carefully cleaned from remaining fine roots prior to further analysis. Roots were processed as the other plant tissues (see above) before analysis.
Plant tissues and a sub-sample of soil fractions were dried (freeze dried for plant tissues), weighed and ground with a rotational ball mill (Retsch, Germany) before being analysed for C and N concentration with an elemental C-N-S analyser (Carlo Erba, Italy).

Microbial biomass was determined in the two soil compartments by fumigation-extraction (Vance et al., 1987) using fresh soil (equivalent to 10g of dry soil) and 40 ml of 0.1M K$_2$SO$_4$ for the extraction. Both fumigated and non fumigated samples were replicated twice and the values averaged prior to statistical analyses. C and N concentration in the extracts were measured by combustion with a total organic C (TOC) and total N analyser (Shimadzu, Japan) and microbial biomass C and N were calculated as a difference between fumigated and non fumigated samples using extraction factors of $k_{EC}=0.42$ and $k_{EN}=0.45$ for C and N respectively (Ross and Tate, 1993). Total soil and root respiration during the 48 hours of the chasing period were measured by quantifying inorganic C in the NaOH traps with the TOC analyser.

5.3.5. **$^{14}$C analysis**

$^{14}$C in plant material was determined with a $^{14}$C analyser (Radiomatic series A 500, Packard, USA) which was installed in series following the elemental C-N-S analyser. Soil $^{14}$C activity was too low to be analysed with the previous method; consequently a sample oxidizer (Packard, USA) was used and the CO$_2$ contained in the combustion products was trapped in a liquid scintillation cocktail (Carbosorb and Permafluor, Packard, USA) and counted with a Packard Tricarb liquid scintillation spectrometer. Liquid samples were counted for $^{14}$C with the liquid scintillation spectrometer in different scintillation cocktails: radioactivity of the NaOH traps was analysed in Ultima Gold whereas extracts from fumigation extractions were analysed in Ultima Flo (Packard, USA) to eliminate error caused by a precipitate occurring in the extract/Ultima gold mixture with the concentrated K$_2$SO$_4$ solution.
Figure 5.3: Effects of the treatment on root, stem, leaf and plant dry matter (g DM per plant). Values are average of 6 replicates ± SE. Bars are grouped by fertiliser N (N- or N+), soil type (ambient soil, S- or enriched soil, S+) and atmospheric CO₂ concentration (ambient, A- or enriched, A+).
5.3.6. $^{15}$N analysis and calculations

Plant and soil samples analysed for $^{15}$N isotopic excess were processed in Nancy and sent to the University of California, Davis isotope laboratory to be analysed with a mass spectrometer (PDZ Europa, UK).

$^{15}$N results are presented using notations derived from Loiseau and Soussana (2000). The N derived from fertiliser (NF) in plant was calculated after sampling as:

$$NF_{plant} = \frac{E^{15}N_{plant}}{E^{15}N_{fert}} N_{plant}$$

where $E^{15}N_{plant}$ and $E^{15}N_{fert}$ are isotopic $^{15}$N excess in the plant and the fertiliser respectively and $N_{plant}$ is the total amount of N in the plant. The N derived from soil (NS) in the plant was calculated as the difference between total N and NF:

$$NS_{plant} = N_{plant} - NF_{plant}$$

And the real recovery (RR) of the fertiliser was calculated as:

$$RR_{plant} = \frac{NF_{plant}}{N_{fert}}$$

where $N_{fert}$ is the amount of N in the fertiliser.

5.3.7. Statistical analysis

All the measured variables were analysed by analysis of variance (ANOVA) as a totally randomized factorial ANOVA ($A \times S \times N$) with 6 replicates. Pots were considered to be the experimental units though we are aware that the A treatment (CO$_2$ concentration of the atmosphere of growth) was not replicated sensu stricto due to the use of two glasshouses only. Because these glasshouses differed only by their CO$_2$ concentration it was considered that pots inside each glasshouse could be used as replicates. When the measure of a given variable was duplicated for experimental reasons (for example microbial biomass), the values were averaged per pot prior to analysis. All analyses were done with Genstat v 6.1 (Genstat, 2002).
Figure 5.4: Effects of the treatment on root, stem and leaf N concentration (%) and total plant N content (mg N per plant). Values are average of 6 replicates ± SE. Bars are grouped by fertiliser N (low, N- or high, N+), soil type (ambient soil, S- or enriched soil, S+) and atmospheric CO2 concentration (ambient, A- or enriched, A+). P values are given for the main effects.
5.4. Results

5.4.1. Dry matter production

Elevated CO₂ concentration during plant growth strongly increased total plant biomass at the end of the experiment. A+ plants had on average 24% more total biomass than A- plants (Fig. 5.3). This positive CO₂ effect mainly affected plants that were supplied with high N fertilisation as shown by the significant interaction between A and N treatment (p=0.02) on total plant biomass. Under N+, elevated atmospheric CO₂ induced a 42% increase in plant biomass whereas this increase was only about 3% under low N supply. Soil origin had no significant effect on total plant biomass despite a trend showing a negative effect of S+ soil (-10%).

When individual tissues were considered, the increased biomass under elevated atmospheric CO₂ only affected sheaths (+55%) and laminae (+36%) while root biomass increased under N+ but not under N- (Fig. 5.3). Similarly the enhanced biomass under high N fertilisation affected aboveground tissues more than roots. Again, the biomass response of laminae and sheaths to elevated atmospheric CO₂ was stronger under high N. Root biomass response to elevated CO₂ was even negative under low N whereas it was enhanced under high N.

5.4.2. N yield and concentration

Elevated atmospheric CO₂ had no significant effect on the total amount of N in plant biomass but it strongly modified N concentration in plant tissues: N concentration decreased by 16, 20 and 36% in root, sheaths and laminae respectively when plants were grown under elevated CO₂ (Fig. 5.4). Laminae N concentration was more reduced by elevated CO₂ under high N fertilisation. The origin of the soil only affected laminae N concentration but did not affect total plant N yield. Plants grown in soil originating from FACE rings exhibited a 22% higher lamina N concentration than plants grown in soil from ambient rings (p=0.032). High N fertilisation significantly increased N concentration in plant tissues by 26%, 36% and 49% in root, sheaths and laminae respectively. Coupled with the N driven increases in plant
Table 5.1: Nitrogen derived from fertiliser (NF in mg N per plant), nitrogen derived from soil (NS mg N per plant) and real recovery of the fertiliser (RR in %) under ambient (A-) and elevated (A+) atmospheric CO₂ concentration, on soil originating from ambient (S-) and elevated (S+) FACE rings and under low (N-) or high (N+) nitrogen fertilisation. Values are mean of six replicates ± s.d. P values of the main effects are given at the 5% level. There was no significant interaction between main effects.

<table>
<thead>
<tr>
<th>N</th>
<th>Atm</th>
<th>Soil</th>
<th>NF</th>
<th>NS</th>
<th>RR</th>
</tr>
</thead>
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<tr>
<td></td>
<td></td>
<td>S-</td>
<td>2.9±0.5</td>
<td>74.5±22.5</td>
<td>96.2±16.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S+</td>
<td>2.4±0.2</td>
<td>66.9±8</td>
<td>79.8±6.1</td>
</tr>
<tr>
<td>N-</td>
<td>A-</td>
<td>S-</td>
<td>2.3±0.3</td>
<td>67.2±23.2</td>
<td>77.8±9.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S+</td>
<td>2±0.2</td>
<td>60.4±16.2</td>
<td>67.8±5.1</td>
</tr>
<tr>
<td>N-</td>
<td>A+</td>
<td>S-</td>
<td>64±11.2</td>
<td>66.8±14.7</td>
<td>100.1±17.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S+</td>
<td>56.4±7.1</td>
<td>66.1±17.7</td>
<td>88.3±11.2</td>
</tr>
<tr>
<td>N+</td>
<td>A-</td>
<td>S-</td>
<td>60.1±5.7</td>
<td>62.4±8.1</td>
<td>94.1±9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S+</td>
<td>54.1±3.5</td>
<td>70.4±10.4</td>
<td>84.7±5.5</td>
</tr>
</tbody>
</table>

Effects

<p>| | | | |</p>
<table>
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<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>p&lt;0.001</td>
<td>n.s.</td>
<td>p&lt;0.001</td>
</tr>
<tr>
<td>A</td>
<td>p=0.002</td>
<td>n.s.</td>
<td>p=0.003</td>
</tr>
<tr>
<td>S</td>
<td>p&lt;0.001</td>
<td>n.s.</td>
<td>p&lt;0.001</td>
</tr>
</tbody>
</table>
biomass described above, this resulted in an 80% increase in total plant N uptake for N+ plants (p<0.001).

5.4.3. Origin of plant N

Elevated atmospheric CO₂ significantly decreased the amount of plant N derived from fertiliser (NF) by about 6% (p=0.002) but this decrease was stronger at low N (-17%) than at high N (-5%) as shown by the significant interaction between N and CO₂ (p=0.047) (Table 5.1). Plant N derived from soil (NS) was not affected by elevated CO₂ despite a trend showing an apparent decrease. The proportion of plant N deriving from fertiliser (% NF) was not affected by elevated CO₂. Soil from FACE rings also decreased plant NF (-11 %, p<0.001) and left NS unaffected. Consequently, %NF significantly decreased when plant were grown on FACE soil (-8%). Obviously high N fertilisation increased plant NF and %NF dramatically. Interestingly NS was not affected by N fertilisation.

5.4.4. ¹⁴C allocation within the plant-soil system

¹⁴C allocation within the plant was clearly affected by the treatments (Table 5.2). Elevated atmospheric CO₂ increased by 15% the percentage of assimilated CO₂ found in the sheath compartment and concomitantly decreased ¹⁴C allocation to the laminae. The origin of the soil had similar effects on ¹⁴C at the plant scale with S+ soil increasing allocation to sheaths and decreasing allocation to laminae by respectively 9 and 17%. On the other hand N fertilisation significantly increased allocation to laminae with N+ plants showing a 30% increase in the fraction of assimilated ¹⁴C found in laminae. Most of the assimilated ¹⁴C remained in the plant 48 hours after the labelling (about 85%) and this was not affected significantly by the treatments. The proportion of ¹⁴C that was respired by the roots and soil microbial biomass during the 48h chasing period was about 10% of the assimilated ¹⁴C and stayed unaffected by the treatments. High N fertilisation decreased the amount of ¹⁴C present in the soil (-31%, p=0.009) and a strong trend showed a similar effect on the proportion of ¹⁴C found in the microbial biomass (-30%, p=0.051). Elevated
Table 5.2: Fraction of total radioactivity recovered per pot in each compartment and relative specific activity of each compartment under ambient (A-) and elevated (A+) atmospheric CO₂ concentration, on soil originating from ambient (S-) and elevated (S+) FACE rings and under low (N-) or high (N+) nitrogen fertilisation. Values are mean of six replicates ± s.d. P values of the main effects are given at the 5% level. Values in italics are significant at the 10% level only. Only the statistically significant interactions are presented here.

<table>
<thead>
<tr>
<th>Fertilisation</th>
<th>Atmosphere</th>
<th>N-</th>
<th>N+</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Soil</td>
<td>S-</td>
<td>S+</td>
</tr>
<tr>
<td>% of total radioactivity recovered</td>
<td>Plant</td>
<td>84.8±6.3</td>
<td>83.1±2.8</td>
</tr>
<tr>
<td>Laminae</td>
<td>10.9±4.3</td>
<td>10.7±2.4</td>
<td>10.9±4.4</td>
</tr>
<tr>
<td>Sheaths</td>
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<td>35.5±6.9</td>
</tr>
<tr>
<td>Roots</td>
<td>40.8±17.9</td>
<td>38.5±6.9</td>
<td>37.5±7.7</td>
</tr>
<tr>
<td>Rhiz. respiration</td>
<td>11.1±4.6</td>
<td>11.9±2.8</td>
<td>11.7</td>
</tr>
<tr>
<td>Soil</td>
<td>4±1.8</td>
<td>5±1.4</td>
<td>5.1±3.3</td>
</tr>
<tr>
<td>Microbial BM</td>
<td>0.4±0.3</td>
<td>0.6±0.2</td>
<td>0.7±0.4</td>
</tr>
<tr>
<td>Relative Specific Activity</td>
<td>Plant</td>
<td>48.5±1.4</td>
<td>47.2±0.9</td>
</tr>
<tr>
<td>Laminae</td>
<td>26.9±1.7</td>
<td>25.2±0.8</td>
<td>24.6±1.6</td>
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<tr>
<td>Sheaths</td>
<td>94.5±2.3</td>
<td>79.9±1.8</td>
<td>83.8±5.1</td>
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<tr>
<td>Roots</td>
<td>39.7±2.9</td>
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<tr>
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<td>196.4±24.9</td>
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<td>0.3±0.04</td>
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<tr>
<td>Microbial BM</td>
<td>3±0.2</td>
<td>3.5±0.1</td>
<td>4.4±0.4</td>
</tr>
</tbody>
</table>
atmospheric CO₂ operated in the opposite direction with an increased proportion of
¹⁴C found in the soil compartments, in particular in the microbial biomass (+62%,
p=0.001). The origin of the soil did not affect significantly ¹⁴C recovery within these
compartments but there was a strong trend for increased allocation of ¹⁴C towards the
microbial biomass in S+ soil (+20%, p=0.067).

When ¹⁴C distribution is considered in terms of relative specific activity (RSA), it takes into account both the size of the compartment and its metabolic
activity. High N fertilisation decreased RSA of all compartments (Table 5.2) although
the effect on laminae and root were not statistically significant. Elevated atmospheric
CO₂ decreased RSA in the plant compartment but strongly increased RSA in soil
(+29%) and microbial biomass (+33%). There was a 24% increase in soil RSA when
plants were grown in S+ soil but this difference was non-significant.

5.5. Discussion

5.5.1. Elevated atmospheric CO₂ increases root exudation

Numerous studies have shown that increases in plant aboveground biomass
under elevated CO₂ are often smaller than expected based on the stimulation of C
assimilation (Chapter 1). It was therefore hypothesised that a great part of the extra C
was allocated belowground. Previous studies in the New Zealand pasture FACE have
shown greater C allocation towards roots and soil. Despite a greater photosynthetic
activity (von Caemmerer et al., 2001) in some abundant species herbage production
stayed unaffected by elevated CO₂ (Newton et al., 2001). However, in this thesis it
has been shown that root production and turnover are greatly stimulated under FACE
conditions (Chapter 4). In addition, both mycorrhizal infection rate (Rillig, M.
personal communication) and nematode abundance (Yeates et al., 2003) were
enhanced under elevated CO₂ showing that other soil C sinks were also greater under
elevated CO₂. In the present study the greater proportion of assimilated ¹⁴C found in
the soil and the microbial biomass when L. perenne plants were submitted to elevated
CO₂ gives a clear indication of enhanced root exudation. This is a common
observation (Cheng and Johnson, 1998; van Ginkel et al., 2000 but see Hodge and
Table 5.3: Microbial carbon biomass (MB-C in mg C g\(^{-1}\) soil), C/N ratio of the microbial biomass (MB-C/N) and rhizomicrobial respiration (mg C day\(^{-1}\) pot\(^{-1}\)) under ambient (A-) and elevated (A+) atmospheric CO\(_2\) concentration, on soil originating from ambient (S-) and elevated (S+) FACE rings and under low (N-) or high (N+) nitrogen fertilisation. Values are mean of six replicates ± s.d. P values of the main effects are given at the 5% level. Values in italic are significant at the 10% level only. No interactions between treatments were measured.

<table>
<thead>
<tr>
<th>N</th>
<th>Atm</th>
<th>Soil</th>
<th>MB-C  (mgC g(^{-1}) soil)</th>
<th>MB-C/N</th>
<th>Respiration (mgC day(^{-1}) pot(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-</td>
<td>A-</td>
<td>S-</td>
<td>339.7±7.6</td>
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<td>68.25±10.5</td>
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<tr>
<td></td>
<td></td>
<td>S+</td>
<td>346.7±6</td>
<td>6±5.2</td>
<td>67.45±19.9</td>
</tr>
<tr>
<td></td>
<td>A+</td>
<td>S-</td>
<td>369.1±7.4</td>
<td>7.4±0.9</td>
<td>81.9±33.6</td>
</tr>
<tr>
<td></td>
<td></td>
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<td>82.3±36.75</td>
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<td>N+</td>
<td>A-</td>
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<td>102.75±19.7</td>
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<td></td>
<td></td>
<td>S+</td>
<td>295.9±7.6</td>
<td>7.6±2.3</td>
<td>70.9±15.3</td>
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<tr>
<td></td>
<td>A+</td>
<td>S-</td>
<td>438.1±7.9</td>
<td>7.9±3</td>
<td>98.95±26</td>
</tr>
<tr>
<td></td>
<td></td>
<td>S+</td>
<td>446.1±11.2</td>
<td>11.2±5.2</td>
<td>112.15±49.45</td>
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Effects

<table>
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<th>N</th>
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<tr>
<td></td>
<td>n.s.</td>
<td></td>
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<tr>
<td>A</td>
<td>p=0.003</td>
<td>n.s.</td>
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<td>p=0.057</td>
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<td>S</td>
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Millard, 1998) but the causes of this CO₂ driven increase in root exudation are still unclear. Is this phenomenon primarily caused by a proportional growth response of roots (van Ginkel *et al.*, 2000; Mikan *et al.*, 2000) or is exudation per unit root area also increased (Lekkerkerk *et al.*, 1990; Cheng and Johnson, 1998)? The results presented here support the second hypothesis since the apparent increased exudation under elevated CO₂ occurred without any change in root biomass (Fig. 5.3). This implies that exudation *per se* was affected by elevated CO₂, but extrapolation of this to what would happen under field conditions would be dangerous. There are good indications that root growth was at least partially limited by pot size. Previous studies describing a relative lower stimulation of root growth compared to shoot growth under elevated CO₂ are rare in the literature hence it can be suspected that belowground growth was constrained in one way or another. We do not know to what extent this may have affected the root exudation process but because C acquisition by roots is controlled simultaneously by leaves’ capacity to assimilate C and by the roots’ demand for C (Farrar and Jones, 2000), it seems intuitively reasonable to expect that an experimental artefact limiting the sink strength for C may have modified the exudation process.

5.5.2. Increased availability of easily decomposable C induced higher microbial growth

Whether the CO₂ stimulation of exudation is greater or proportional to the plant biomass increase is nevertheless only of relative importance here. Indeed, the main question remains whether a greater availability of easily decomposable C will or will not affect soil microbial biomass that is usually C limited. Several studies have observed a strong enhancement of root derived C allocation towards microbial biomass under elevated CO₂ (Mikan *et al.*, 2000; van Ginkel *et al.*, 2000) and a stimulation of microbial biomass due to an increase in available C substrate (Zak *et al.*, 1993; Cotrufo and Gorissen, 1997). In our study, the fraction of assimilated ¹⁴C and the RSA in the microbial biomass significantly increased under elevated CO₂. Not only did the increased availability of C substrate under elevated CO₂ stimulate the size and activity of the microbial biomass pool but there was also an increase in the
apparent preference of soil microorganisms for exudates (as shown by the higher RSA, Table 5.2).

5.5.3. Effects on native soil organic matter decomposition; occurrence of a priming effect

Belowground respiration (i.e. by soil microorganisms and roots) was increased under elevated CO₂ whereas root biomass was not affected. In consequence, it is reasonable to consider that the enhancement of belowground respiration was mainly caused by an increase in soil respiration. As described in Cheng and Johnson (1998) two mechanisms could be the cause of this phenomenon. First, as observed in this study, root exudation was higher under elevated CO₂ and the subsequent metabolism of this material by soil microorganisms would lead to an increase in soil respiration. If this process was quantitatively important, it should increase the intensity of ^14C labelling of soil CO₂; this was not the case here (Table 5.2). This implies that exudates use by soil microorganisms only represented a small part of the observed increase in soil respiration and that mineralisation of native soil organic matter was increased: this is the so called “priming effect” (Bingeman et al., 1953).

Past studies describing the effect of increased exudation on SOM decomposition have shown both stimulatory (Zak et al., 1993) or suppressive (Cardon et al., 2001) effects of root derived exudates on SOM decomposition. An hypothesis (Lekkerkerk et al., 1990; Cardon, 1996) identifying N as a main driver of the response of microorganisms to increased exudates predicts that under sufficient N availability, soil microorganisms will preferentially metabolize root exudates that have a high C:N ratio, but that are easily available, instead of decomposing N-rich native SOM in which the C is less accessible. However, several studies have contradicted this prediction by showing an opposite effect; the CO₂-induced “priming effect” being stronger under high N (Hungate et al., 1997; Cheng and Johnson, 1998). In our study the absence of interaction between N and atmospheric CO₂ on rhizosphere respiration (Table 5.3) suggests that the “priming effect” was independent of N availability. Fontaine et al. (2003) recently proposed a conceptual model in which the “priming effect” resulted from the competition for energy and nutrient acquisition between two
groups of microorganisms: those feeding on mainly easily accessible C and those specialized in metabolising native SOM. Under this assumption, the "priming effect" might well be dependent on complex controls, in particular if the structure of microbial populations is modified by elevated CO$_2$ (Montealegre *et al.*, 2002) or if soil mesofauna feeding on soil microbes are also affected by elevated CO$_2$ (Yeates *et al.*, 2003). Therefore a simple model considering soil microorganisms as a single entity responding to C and N availability is probably over-simplistic and can lead to the apparent contradictions described above.

5.5.4. *Did the increased SOM mineralisation affect N availability for plants?*

The occurrence of a "priming effect" caused by a CO$_2$-induced enhancement of the microbial biomass could potentially affect N availability in two opposite ways. Firstly, increased microbial growth under elevated CO$_2$ could increase microbial need for N; at a given moment in time more N would therefore be immobilised and not accessible by plants (Diaz *et al.*, 1993). Secondly, the increased metabolisation of native SOM due to the "priming effect" described above could increase soil mineral N (Zak *et al.*, 1993) since stabilised SOM is usually richer in N. Our results give no evidence that N availability for the plants was modified under elevated atmospheric CO$_2$ (Fig. 5.4). Although N concentration of the different plant organs was lower under elevated CO$_2$, in particular in laminae, total plant N yield was the same. Nevertheless an unaltered plants N yield gives no indication of the origin of the N and potential effects of the treatments on N dynamics in the soil. $^{15}$N data gives clear indications of altered N cycling. NF was clearly decreased by elevated atmospheric CO$_2$ while NS stayed unaltered (Table 5.1). Similarly, the concomitant decrease of fertiliser recovery rate (RR) and unaltered plant N yield under elevated CO$_2$, shows that for a given amount of N uptake by plants, a greater proportion originated from soil rather than from the fertiliser. Under the assumption that both fertiliser and soil mineral N are perfectly mixed in the soil (e.g. plants do not preferentially take up one or another), this indicates an increased dilution of fertiliser N by soil N because of the enhanced native SOM decomposition. Therefore microbial N immobilization and SOM N mineralisation seems to counterbalance each other leaving N availability unchanged for plant growth.
Chapter 5

5.5.5. Long-term CO₂ effect: origin of the soil

In a previous experiment in the New Zealand pasture FACE site (Chapter 4) we have shown that soil originating from elevated CO₂ gave clear indications of accumulating particulate organic matter (POM) and that this accumulating POM had a lower C/N ratio than the POM in the ambient soil. We attributed the accumulation of POM mainly to an increase in root growth and turnover under elevated CO₂ (Chapter 4) and the decrease in C/N ratio to the increased proportion of legumes in the pasture (Edwards et al., 2001). On this basis, we hypothesised that such a modification of the SOM quantity and quality could potentially alter plant N nutrition and control the short term effects of elevated CO₂ on soil dynamics that occurs through increased exudation. As expected, due to the lower C/N ratio of the POM, soil coming from FACE rings provided more N for a given amount of mineralized soil organic matter as shown by the higher dilution of fertiliser N by soil N (Table 5.1). But the origin of the soil had no significant effect on plant N yield showing that N availability stayed most probably unaltered. Because we did not observe a significant increase of the microbial biomass nor any change in its C/N ratio, we can assume that there was no increase in the gross microbial N immobilisation. Then why did the hypothesised increase in gross N mineralisation from SOM not induce an increase in mineral N availability? In this study we did not measure ^15N in soil organic matter compartments. Loiséau and Soussana (2000) using a similar labelling technique have shown that a significant amount of fertiliser N was immobilised in soil organic matter pools shortly after fertilisation, thus limiting transfers of ^15N through litter decomposition. In our study the greater amount of SOM in FACE soil might well have created a greater sink for soil N. Therefore the greater gross N mineralisation observed may have been counterbalanced by a greater gross N immobilisation by SOM.

5.6. Conclusion

Our results give clear indications that elevated CO₂ effects on root exudation have a negative effect on C sequestration due to the existence of a priming effect on native SOM decomposition. This phenomenon also explains the faster soil N turnover that allows a sustained N availability for increased plant growth. It also appears that
the accumulation of organic matter observed in the field in elevated CO$_2$ rings does not induce slower N cycling under the conditions of our experiment. This strongly suggests that the observed accumulation of SOM under elevated CO$_2$ is not driven by simple C and N limitations.

5.7. Acknowledgements

I thank P. Gross for the glasshouse maintenance, P. Marchal for help with the analysis and the numerous students and staff who helped with plant sampling after the $^{14}$C labelling.

5.8. References


Chapter 6: General Discussion
6.1. Introduction

In the conclusion of a recently published review, Körner (2000) emphasised that CO₂ experiments should avoid over-simplification, in other words should aim for realism in their design: “By “realism” I mean accounting for the complexity of interactions in real life. Since there is no doubt that plant species respond in rather different ways depending on their age, neighbours, microbial partners, soil resources and atmospheric condition, the experimental negation of these interactions and dependencies is wasteful – or, even worse, creates a biased picture of the world.” In elevated CO₂ experiments conducted in grassland to date, one source of such behavioural variation that has been overlooked is the potential influence of grazing and the need for information about elevated CO₂ effects on grazed grasslands was also noted by Newton et al. (2001), who wrote: “...it is by no means certain that we can extrapolate the results of CO₂ enrichment experiments from cut to grazed situations. In fact, what is required is not only a greater understanding of how changes in pasture might alter animal performance at elevated CO₂ but also, more importantly, a clearer picture of the CO₂ response of ecosystems in which grazing animals are an integral part”.

The overall objective of this thesis is derived from the concepts expressed in these two statements. To assess the effects of elevated CO₂ on organic matter fluxes in a grazed grassland and the possible influence of these fluxes on soil organic matter accumulation and N availability it was thus necessary not only to consider species diversity and their relative response to elevated CO₂, but also the ecosystem fauna both aboveground (i.e. sheep) and belowground. In this general discussion I will summarize the results described in the previous chapters and draw some general conclusions. The two main themes are organic matter accumulation in the soil under elevated CO₂ in relation to the organic matter fluxes and the implications of grazing on nitrogen (N) cycling.
Figure 6.1: Possible pathways through which elevated CO₂ may affect plant-soil organic matter fluxes.
6.2. Effects of elevated CO$_2$ on plant-soil organic matter fluxes and organic matter accumulation in the soil

Classically, three main mechanisms are proposed in the elevated CO$_2$ literature as possible explanations for a modification of organic matter pools in soil under elevated CO$_2$ (Figure 6.1). Firstly, plant litter decomposition rates could be lower under elevated CO$_2$ both through direct effects of elevated CO$_2$ (decreased decomposability at the species scale arising from higher C:N ratio) or indirect effects (shifts in botanical composition or biomass allocation). Secondly, due to the increase in C assimilation under elevated CO$_2$ the size of the organic matter fluxes could be greater. Thirdly, increased root exudation of readily decomposable C under elevated CO$_2$ could modify the trophic preference of soil microorganisms and alter soil organic matter decomposition. I will now briefly review the results presented in the previous chapters about these three effects pathways.

6.2.1. Rates of organic matter decomposition

The data presented in Chapter 3 clearly showed leaf litter C:N ratio and decomposition rates, when considered at the single species level, were not affected by elevated CO$_2$. This result is now globally accepted (Norby and Cotrufo, 1998) after a long period of conflicting results showing both lower (Couteau et al., 1991; Cotrufo and Ineson, 1996) or unaltered litter decomposition rates under elevated CO$_2$ (Franck et al., 1997; Dukes and Field, 2000). These divergences very probably have methodological origins. In particular due to the use of green leaves in decomposition experiments, instead of naturally senesced litter, can cause an experimental bias since green leaves usually exhibit a major change of their chemical composition under elevated CO$_2$ (Cotrufo et al., 1998).

The rejection of the “litter hypothesis” (as formulated by Strain and Bazzaz, 1983) does not diminish the importance of indirect effects of elevated CO$_2$ at the community level that may affect overall decomposition rates. The results presented in Chapter 3, identify effects of elevated CO$_2$ on botanical composition as an important driver of decomposition processes. Two of the four species used in this experiment,
Trifolium subterraneum and Hypochaeris radicata exhibited greater decomposition rates than the two grasses. These two species with a higher decomposition rate are also two species that tend to increase in proportion in the pasture under elevated CO₂. It was therefore concluded that, overall, elevated CO₂ should lead to an increase in leaf litter decomposition rate at the study site. Another secondary effect of elevated CO₂ that affected decomposition rates was the increase in biomass allocation towards root growth and turnover (Chapter 4). Whereas above ground production was not changed by elevated CO₂, root growth was stimulated by nearly 60% in summer-autumn and about 10% in winter-spring. Root turnover was also greater under elevated CO₂ and the result of these effects was a large increase in the proportion of litter originating from roots under elevated CO₂. A more comprehensive study involving species-specific root litter material would be required to quantitatively define, at the pasture scale, the global implications of this shift in biomass allocation. Nevertheless, the results obtained in Chapter 3 where decomposition of mixed-species root material was tested, show that a greater proportion of root litter in the total litter pool is not only unlikely to decrease the overall decomposition rate but probably leads to a slight increase. Overall, decomposition rates per se of plant litter at the study site were therefore increased under elevated CO₂. The only component of the plant-soil organic matter returns to the soil that showed a decrease in decomposition rate was ruminant faecal material under dry conditions.

6.2.2. Fluxes of organic matter under elevated CO₂

To determine if elevated CO₂ affects organic matter accumulation in soil the rates of decomposition must not be considered in isolation. Changes in flux sizes are also of importance. Leaf litter deposition was found to be unaltered under elevated CO₂ (Chapter 4) but root production and turnover were higher under elevated CO₂, hence root litter deposition was greatly increased. This has implications for the relative proportion of root litter relative to leaf litter as discussed above and also for the total return of organic matter to the soil. In this study there were clear indications that the total amount of plant organic matter returning to the soil is greatly increased under elevated CO₂. This result is not a consistent outcome reported in the literature and some discussion of the reasons for this is necessary. Even with the commonly
observed disparity between C assimilation and C accumulation (see Chapter 1) and the consequence that plants may dissipate extra C fixed under elevated CO₂ via enhanced root turnover (Stocker et al., 1999), experimental evidence for such a phenomenon is elusive and some studies did not detect the expected “C overflow” from the roots to the soil (Arnone et al., 2000). The latter authors, reviewing the existing literature, report that studies showing a large increase in root biomass under elevated CO₂ are mainly growth chambers experiments (see for example Newton et al., 1994; Newton et al., 1996; Fitter et al., 1996; Fitter et al., 1997) whereas field studies generally report lower or no CO₂-induced stimulation (Leadley et al., 1999; Arnone et al., 2000). They hypothesise that growth conditions might create an artefact by artificially increasing the growing period in growth chambers and therefore conclude that in “real life” root growth might not be significantly stimulated under elevated CO₂.

Arnone et al. (2000) in their own study assessed root standing biomass, growth and mortality and showed clearly that in their system elevated CO₂ did not affect the C flow from roots to the soil. Nevertheless some of the other in situ studies they use as example to draw their conclusions only report effects of elevated CO₂ on root standing biomass (Körner et al., 1997; Sindhøj et al., 2000). Our own results clearly show that root standing biomass and growth can respond very differently to elevated CO₂ (Chapter 4) so I believe that, based on these examples, it cannot be concluded that elevated CO₂ does not affect root C flux to the soil. It is possible, as Arnone et al. (2000) state, that growth chambers conditions artificially enhance the root system response to elevated CO₂ but it is equally probable that root standing biomass measurements, when taken alone in in situ studies, give an incomplete picture of the biomass flux and underestimate fluxes from the roots to the soil. Consequently, in my view, the existing literature tends to validate the results presented in Chapter 4.

6.2.3. Exudation under elevated CO₂

Root exudation of easily decomposable C compounds is the third pathway though which elevated CO₂ potentially affects soil organic matter (Fig. 6.1). The
problem here is not to assess the changes in quality/quantity of plant litter returns but how the alteration of easily decomposable C by the roots may modify the fate of this litter by decreasing the mineralisation of complex soil organic matter (Lekkerkerk et al., 1990) or on the other hand, by stimulating its mineralisation through a priming effect as suggested by the results of Zak et al. (1993).

Root exudation in this study was not measured in situ on a variety of plant species but assessed on individual Lolium perenne plants under glasshouse conditions (Chapter 5); therefore extrapolation of our results should be cautious. Nevertheless some interesting patterns emerged from this experiment. Firstly, it appeared that exudation was increased by elevated CO2 since more 14C was recovered in the soil and from the microbial biomass under elevated CO2 and this increase was independent of root biomass. Given the increase in root production at elevated CO2 shown in Chapter 4, it suggests that an even larger increase in C exudation under elevated CO2 might be occurring in the field. The enhanced availability of readily decomposable C drove an increase in the soil microbial biomass; this would be expected since microbial biomass is usually C limited (Smith and Paul, 1990). In this experiment, the results indicate that the enhanced microbial biomass led to a priming effect through a tendency to greater decomposition of polymerised organic matter. In other words, even if the microbial population partially changed its feeding source by metabolising more exuded-C (as shown by the higher relative specific activity of the microbial biomass, Chapter 5), the extent of the priming effect led to a situation that was not detrimental for organic matter decomposition. In our study, the priming effect was independent of N fertilisation. This shows the possible limits of the conceptual model proposed by Hungate and Chapin (1995, in Cardon, 1996). In their scheme, if mineral nutrients are abundant in the soil, microorganisms will utilise readily decomposable C and will immobilise mineral N leading to a suppression of organic matter decomposition. If, instead nutrients are scarce, microorganisms will use the exuded C as a C source but break down more organic matter in order to obtain nutrients. This leads to the question of whether the fate of soil organic matter in grassland under elevated CO2 can be simply assessed based on C-N interactions or if a more complex model is required. This will be discussed in the following paragraphs.
6.2.4. Accumulation of soil organic matter under elevated CO$_2$

The results concerning the decomposability of the litter and the priming effect caused by enhanced exudation under elevated CO$_2$ should tend to create an equilibrium where soil organic matter does not accumulate, or even decreases. This raises the question: why is the increased flux of organic matter entering the soil, in particular in the form of dead roots, not totally processed by the microbial biomass? Results from Chapter 4 clearly indicate an increase in the coarser fraction of particulate organic matter under elevated CO$_2$ showing that the extra organic matter entering in the soil organic matter pool cannot be totally metabolised. In addition, this supplementary organic matter exhibits a lower C/N ratio suggesting together with long term results from the same site (Ross et al., 2003) that lower N availability is not the driving factor of this accumulation under elevated CO$_2$. Clearly, there are one or more limiting processes in the soil leading to this organic matter accumulation. In particular, if an increase in microbial biomass was observed under laboratory conditions at elevated CO$_2$ (Chapter 5) this effect was not observed under field conditions (Chapter 4, Ross et al., 2003) suggesting that the microbial population might be controlled by some processes we did not assess directly in our studies.

A possible answer to this question may be changes in soil fauna. A recent study based on the same experimental site (Yeates et al., 2003) showed a dramatic increase in nematode abundance under elevated CO$_2$. In that study, however, the relative abundance of the different trophic groups (omnivorous, root-feeding, bacterial-feeding and fungal-feeding nematodes) was not affected by elevated CO$_2$ making it relatively unlikely that a specific soil function was depressed under elevated CO$_2$. Moreover, even an augmentation in bacterial or fungal-feeding nematodes does not induce a suppression of microbial activity but may rather be associated with an increase in microbial activity (G. Yeates, personal communication) and the subsequent decomposition of soil organic matter. Therefore in the sward studied here, the extra nematode abundance is more likely to be the a sign of an enhanced overall soil activity that should be favourable to organic matter processing, though it should kept in mind that the interactions among the various functional groups of below-ground microbes and microfauna are complex and, so far, largely unpredictable (Wardle et al., 2001; Wardle, 2002).
Figure 6.2: The effect of the intensity of grazing (the average leaf area index at which the sward is sustained) on the major physiological components of herbage growth and utilisation (from Parsons et al., 1983).
A second possible answer to this accumulation of organic matter under elevated CO₂ would be to look further than possible C and N limitations. It is indeed likely that the availability of other macro-nutrients (P in particular) and possibly meso- and micro-nutrients can control ecosystem response to elevated CO₂ and more specifically microbial response to an increased organic matter supply. Studying the effects of elevated CO₂ on soil microbial biomass in a calcareous grassland, Niklaus (1998) concluded that microbial growth in their system under elevated CO₂ was limited by N but not by P availability. However, in this study, there was evidence that P might be a limiting factor, at least for aboveground plant growth. This leads to an interesting observation. In grasslands, such as this, containing a significant amount of legumes, the relative increase in N demand (compared to C) can be, at least partially counterbalanced by the increased proportion of legumes under elevated CO₂. Other nutrients however do not benefit from a similar mechanism and might become relatively scarce. This implies that future studies should widen their range of investigation by assessing the numerous controls that other nutrients can represent for ecosystem response to elevated CO₂. In addition there is probably also a strong need for a further integration of above and belowground processes. If, in relatively short-term experiments, including this one, ecosystem responses to elevated CO₂ are (apparently) mainly driven by plant and primary soil decomposers, it is probably hopeless to predict “real” ecosystem response to climate change without assessing indirect CO₂ effects on the multiple trophic levels of the soil food web.

6.3. An exploration of the possible impacts of grazing on N cycling in a pasture exposed to elevated CO₂

A second aspect of the results presented earlier that deserves further exploration is the implication for N cycling under elevated CO₂ of including the ruminants in the experimental system. In order to assess this, a simple conceptual model was developed to simulate the effects of an increasing herbage utilisation by ruminants on the amount, the form and fate of N returns in a pasture. This model has no quantitative objectives; its outcomes will only provide a basis for further discussion about possible interactions between grazing and elevated CO₂. In particular
Figure 6.3: Relative N fluxes from herbage to litter and animal utilisation in a grazed pasture (modified from Thornley, 1998)

Figure 6.4: Relative C fluxes from herbage to litter and animal utilisation in a grazed pasture (modified from Thornley, 1998)
I will compare the relative and absolute N returns that occur through litter decomposition and faeces/urine deposition as a consequence of grazing intensity.

6.3.1. Description of the model

Grazing intensity can affect N cycling in a pasture through two main mechanisms. Firstly, the amount of N that is recycled is a function of the average biomass that is sustained in the pasture. Secondly, the amount of this biomass that is removed by animal grazing determines the proportion of N returned to the soil through leaf litter or animal excreta. The effects of grazing intensity on these two processes are described in Fig. 6.2. More specifically this figure taken from Parsons et al. (1983) describes the effects of the amount of leaf material that is sustained in the long-term in a pasture, on gross photosynthesis and on the balance between respiration and root growth, gross shoot production and intake by the grazers. In swards maintained on average at high leaf area index (LAI) (low grazing intensity, left side of the graph) canopy gross photosynthesis is close to a maximum, and the maximum amount of plant biomass is produced. To maintain the sward in this state however, only a small proportion of the leaves produced can be harvested and therefore most of the leaves produced die and return to the soil as litter. In swards maintained at lower LAI (right side of the graph), a greater proportion of plant material is grazed and the litter flux is therefore lower. In spite of a reduced gross photosynthesis induced by the lower LAI, this management achieves a better balance between photosynthesis, gross tissue production, death and intake. At even higher intensities of defoliation (extreme right of the graph), all components of production and utilisation are reduced: the sward is overgrazed. This figure gives us clear relationships between grazing intensity, amount of tissue production and the relative proportion of this production that is grazed or lost through litter decomposition. For the model presented here Parsons et al.’ figure provides a relationship between two variables: herbage biomass production (BM) and the proportion of this biomass that is grazed by the animal, the utilisation (U), in relation to grazing intensity.

Figure 6.3 (modified from Thornley, 1998) describes schematically the conceptual model and presents the three main processes that generate major
Figure 6.5: Effect of an increasing herbage utilisation by the grazers on a) the average C/N ratio of the organic matter returns, b) the total N recycled and lost from the pasture and c) the proportion of cycling N lost from the pasture.
discrepancies between grazed and ungrazed systems. Firstly, the herbage utilisation determines the relative volume of plant material returning to the soil as litter or through the grazing animals. Secondly, due to N resorption (R) during leaf senescence, the N concentration of the plant material deposited as litter is lower than the N concentration of the grazed material. Thirdly, because N is only marginally retained by grazing animals, nearly all of the ingested N is excreted in faeces or urine, their relative proportion depending on the N partitioning (P). Figure 6.4 presents a simple C cycle in a grazed pasture to highlight the fact that grazing animals "decouple" C and N cycles (compared to organic matter returns through litter) because most of the ingested C is not returned to the soil. This has major implications for the overall C/N ratio of the organic matter returned to the pasture.

6.3.2. Effects of grazing intensity on N cycling in a pasture

The most important effect of an increased grazing intensity on N cycling is to alter the form of the N returns by consistently decreasing their average C/N ratio (Fig 6.5 a). Under minimal grazing, all the N returns occur through litter deposition and therefore exhibit a maximum C/N ratio value (i.e. the one of the litter). When the proportion of grazed material increases, the overall C/N ratio of the returns continuously decreases for two reasons: the animals feed on material with higher N concentration (compared to the litter) and while most of the ingested N is returned to the soil, a great part of the ingested C is lost through respiration and methane production. Importantly, the total amount of N recycled is also greatly affected by grazing intensity. In referring to amount of N recycled, I mean all the N that is taken from the live plant pool and that is returned to the soil by whatever pathway. When going from low to intermediate herbage utilisation rates, the total amount of recycled N increases. Indeed the lower herbage biomass (and thus the smaller N pool in the herbage) induced by the lower LAI at higher utilisation rates, is more than counterbalanced by the fact that the grazed herbage pool has a higher N concentration than the herbage returning to the soil as litter. Nevertheless at even higher utilisation rates the total amount of N recycled decreases because the herbage standing biomass and associated fluxes are increasingly reduced, so limiting N cycling. Obviously, the greater the herbage utilisation by animals, the greater the proportion of cycling N that
Figure 6.6: Impacts of various value of partitioning of ingested N between urine and faeces on a) total N losses from the pasture b) the proportion of recycled N lost from the pasture.
is potentially lost from the system, since animal returns, urine in particular, are highly susceptible to both volatilisation and leaching.

In summary, if we focus on the comparison between a well managed grazed grassland (with a herbage utilisation of about 0.3-0.4) and a “natural” grassland (with a very lenient defoliation management) which is the type of system used in most other CO₂ experiments (Owensby et al., 1999; Niklaus et al., 2001), it is obvious that there are numerous differences that may affect the response to elevated CO₂. The most important of these are that the total amount of N cycling is greater and the overall C/N ratio of the material in which N is returned is dramatically lower. The compounding of these two processes maximises the potential for N losses from the system.

6.3.3. Possible interactions with elevated CO₂

Clearly this simple model cannot take into account the full range of effects of elevated CO₂ on N cycling. Nevertheless this tool allows me to explore some possible effects of elevated CO₂ that would not occur in an ungrazed system. In particular, Chapter 2 presented an unexpected effect of elevated CO₂: a shift in N partitioning between faeces and urine. Though this shift will leave the total amount of N cycling in the system unchanged, it will nevertheless affect the amount of N that remains in the system. Figure 6.6 indeed shows that an increased proportion of ingested N returning to the soil as urine creates greater potential for N losses, both in term of relative and absolute amounts. This is an outcome of the relatively lower potential N losses from dung than from urine (Haynes and Williams, 1993). Not only do greater N losses through volatilisation and/or leaching have a negative environmental impact, they also “unlock” the grassland N cycle under elevated CO₂ and contribute to a lower N availability for plant growth.

A second interesting outcome of this model is its response to changes in N resorption rate during senescence. It is generally accepted that elevated CO₂ does not directly affect N resorption in senescing leaves (Norby et al., 2000) although a small decrease in resorption efficiency can occur in situations where the average proportion of leaf N resorbed during senescence is high (Arp et al., 1997). However, changes in
Figure 6.7: Impacts of various leaf N resorption rates on a) the total amount of N recycled in the pasture and b) the C/N ratio of the organic matter returns.
plant nutritional status or shifts in botanical composition can affect overall resorption efficiency since under low N availability plants tend to retain more N during senescence (Berendse and Aerts, 1987). Species-specific differences are also evident (Aerts, 1996). Data from Chapter 3, for example, tend to show that clover, a species that does not essentially rely on soil N availability, retains less N during senescence than grasses. Therefore an increase in the proportion of legumes under elevated CO₂ may lead to a lower resorption rate at the community scale. Fig. 6.7 shows that a lower resorption rate tends to offset the effect of increasing herbage utilisation on the C/N ratio of the returns. Indeed, with a low resorption rate the average N concentration in green herbage and litter are more similar. In relation to the total amount of N recycled, lower resorption rates increase the proportion of N that is recycled through litter and thus increase the total amount of N recycled. Nevertheless the shape of the curve defining the relationship between herbage utilisation and total N changes: at very low resorption rates total N is near maximum at a low utilisation rate and rapidly decreases driven by the negative effect of grazing pressure on herbage biomass.

Again, this conceptual model was not designed to be a predictive tool. Its only aim was to highlight the fact that N cycling (and other nutrients as well) under grazing is intrinsically different from N cycling in a cut grassland. Therefore this creates a real need for studies assessing the effects of elevated CO₂ specifically in grazed pastures. This thesis has provided a small contribution to this need but numerous key processes have not been studied. In particular the fact that animal N return is, unlike litter decomposition, a spatially heterogeneous process. Due to the deposition of faeces and urine, a pasture is a mosaic of patches ranging from very high to very low nutrient availability, creating potential for interactions not apparent when studying steady-state component processes. This possibility of complex interactions has major implications for herbage production, botanical composition and individual plant chemical composition. Because of its multiple influences on grassland processes, heterogeneity of nutrient availability is a characteristic that has a great potential to interact with elevated CO₂ (Newton et al., 2001). Therefore, it seems to me that one of the great challenges of future studies dealing with elevated CO₂ and grasslands will be to understand the role of spatial heterogeneity if we are to predict more accurately the effects of elevated atmospheric CO₂ on grasslands.
6.4. References


Stimulation de la croissance et du turnover racinaire d’espèces prairiales par l’augmentation de la [CO₂]

Allard, V. et Newton P.C.D.

Dans le contexte actuel de changement climatique et plus particulièrement de l’augmentation de la concentration en CO₂ atmosphérique ([CO₂]ₐ), une des questions centrales relatives à l’étude des écosystème terrestres sous [CO₂]ₐ enrichi, est leur capacité ou non à stocker dans les sols le flux de carbone supplémentaire créé par la stimulation de l’activité photosynthétique des couverts végétaux. En plus de leur importance économique majeure, les écosystèmes prairiaux ont la particularité d’associer dans leurs sols des stocks importants et un turnover rapide de matière organique ce qui en fait un système particulièrement sensible à un changement de quantité ou qualité de la matière organique entrant dans le système.

Afin de quantifier l’effet de l’élévation en [CO₂]ₐ sur ces flux de matière organique vers le sol, nous nous sommes concentrés sur l’étude de la croissance et du turnover racinaire d’espèces dominantes dans le système étudié, ce compartiment étant primordial quantitativement sous prairie pâturée mais aussi particulièrement sensible à l’élévation de la [CO₂]ₐ. Cette étude a été menée dans une prairie pâturée tempérée pourvue d’un système FACE (Free Air CO₂ Enrichment) à Palmerston North, Nouvelle-Zélande. Le dispositif consiste en 6 anneaux (Ø = 12 m), trois desquels étant enrichis en CO₂ (valeur cible 475 ppm), les trois autres laissés à concentration ambiante (≈ 370 ppm). La croissance racinaire mesurée par la méthode des « ingrowth cores » est stimulée de plus de 30% par l’élévation de la [CO₂]ₐ. De plus une estimation du temps de résidence des racines vivantes montre que celui-ci est diminué de près de 50% par le CO₂. De ce fait, le flux de matière organique quittant le compartiment « racines vivantes » et rejoignant la matière organique du sol est stimulé de l’ordre de 80% ce qui représente un flux supplémentaire de matière sèche d’environ 1.1 t par hectare et par an. L’élévation de la [CO₂]ₐ crée donc une forte augmentation des entrées de matière organique dans les sols prairiaux, ce qui peut potentiellement modifier fortement les cycles du C et de l’N dans le sol. Des mesures concernant la qualité et la dégradabilité des racines sont en cours et sont nécessaires pour conclure à une éventuelle capacité de ces sols à stocker cet excès de C.

Interactions between grazing by ruminants and elevated CO₂ on nutrient cycling in pastures.

V. Allard and P. C. D. Newton

Abstract
In a New Zealand temperate grazed pasture currently exposed to either ambient air or air enriched to 475 μl l⁻¹ CO₂ using a Free Air CO₂ Enrichment (FACE) technology, the pattern of nitrogen (N) return through sheep has been investigated. We show here that changes in the composition of the diet of ruminants at elevated CO₂ result in a different partitioning of N in the nutrient returned which could create a greater potential for N losses from grazed pasture in a high CO₂ world.

Keywords: elevated CO₂, N cycling, grazed pasture, faeces, urine

Introduction
It has often been hypothesised that the fertilisation effect of elevated CO₂ for plant communities may be buffered in the long term by a decrease in litter quality. It now seems that these changes are too small to consistently alter decomposition processes (Norby and Cotrufo, 1998). In grazed pastures about 40% of the N return to the system occurs through ruminant excreta so any CO₂-induced changes in faeces chemical composition and urine N concentration are likely to strongly affect N cycling at the pasture scale.

Materials and methods
This study was conducted during summer 2000 in a FACE experiment located in a temperate grazed pasture in Bulls, New Zealand. The facility consists of 6 experimental units (12 m diameter "rings"), three of them were enriched with CO₂ (475 μ l⁻¹) since October 1997 while the other three were kept at ambient CO₂ concentrations (see Edwards et al., 2001 for details). Each ring is fenced off in order to control the timing and duration of the grazing by sheep. Five sheep were allowed to graze for 24 hours successively in each ring, firstly in the three ambient, then in three elevated ones. Dung was sampled using dung bags on the last day of grazing in each treatment (day 3 and 6). Dung subsamples were placed in nylon bags and returned to the rings to determine in situ decomposition rates. A crossover design was used to test potential direct CO₂ effects on decomposition. Prior to and after the grazing, the herbage was dissected for botanical composition determination and was analysed for chemical composition by NIR. Chemical composition of the dung samples was determined by wet chemistry analysis.
Results and discussion
The botanical composition of the grazed herbage was significantly different between the two treatments (Figure 1). At the single species scale, elevated CO₂ significantly decreased the leaf N concentration (Figure 2). This buffered the increase in legume content and as a result the N concentration of the ingested feed was similar in the two treatments (2.67% N in average). The digestibility of the herbage from the enriched plots was higher (76%) than from the ambient (74%), which is consistent with the higher proportion of legumes. The nitrogen content of the dung produced when sheep were grazing in high CO₂ rings was significantly lower (3.1%) than from the low CO₂ rings (3.5%) which induced a significant decrease in dung decomposition rate (Figure 3) during the first month of decomposition. The proportion of dietary N present in urine was higher when sheep were fed on high CO₂ herbage (72% against 66%). The same pattern was found in a subsequent experiment, which showed significantly greater N concentration in urine (+104%) from sheep grazing elevated CO₂ rings.

Conclusion and perspectives
Under our experimental conditions, elevated CO₂ induced a modification of the dietary N partitioning between faeces and urine. Since the absolute N intake was unaltered by elevated CO₂ despite the botanical composition changes, this modification may be induced by complex interactions between herbage N digestibility (protein type) and carbon pools quality (Beever, 1993). This can potentially strongly affect N cycling at the pasture scale since a slower dung decomposition under elevated CO₂ would contribute to an increased N immobilisation in the soil organic
matter pool. In addition the increase in N concentration in the urine at elevated CO₂ might well result in greater N loss through volatilisation and leaching.

Figure 3: Decomposition (in % of the initial DW remaining) of faeces produced by sheep, which have been grazing in ambient (...) and enriched (——) submitted to either ambient (■) or enriched (○) atmosphere during decomposition.

References
Appendix 3: Poster presented at the 2003 “Journées d’Ecologie Fonctionnelle” meeting, Nancy, France. March 2003. (See next page)
Comparaison des effets court-terme et long-terme de l'élevation en CO₂ atmosphérique sur l'activité photosynthétique d'une graminée prairiale

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Objectifs

Le CO₂ étant le substrat primaire de la photosynthèse chez les végétaux, son augmentation dans l'atmosphère du fait de la combustion d'énergies fossiles favorise l'assimilation de carbone par les plantes. Cependant les ressources en autres nutriments, en particulier en azote restant limitées, certaines plantes réduisent leur capacité photosynthétique en allouant leur nutriments vers d'autres compartiments métaboliques. On parle d'acclimatation de la photosynthèse. Dans des écosystèmes naturels tels que des prairies, soumis à un enrichissement en CO₂ de longue durée, des rétro-actions complexes peuvent aussi se produire, si par exemple, la disponibilité en minéraux du sol est affectée par l'élévation en CO₂. Notre objectif est donc de comparer les effets long-terme du CO₂ (via le sol) et court-terme (agissant directement sur la plante) sur l'activité photosynthétique des plantes afin de pouvoir prédire plus finement le comportement de ces dernières sous des climats futurs.

Méthodologie

1) En Nouvelle-Zélande une prairie est soumise à un enrichissement en CO₂ depuis 5 ans. Du sol de parcelles ambiantes (S-) ou enrichies en CO₂ (S+) est prélevé.
2) Mise en culture du sol S+ ou S- avec une graminée (Lolium perenne) sous serre à atmosphère ambiante (A-) ou enrichie (A+) en CO₂. Deux niveaux de fertilisation azotée ont aussi été appliqués: 100 UN (N+) ou 5 UN (N-).
3) Après 3 mois de croissance, l'assimilation de CO₂ est mesurée pour une gamme de concentration en CO₂. Ces données sont ensuite ajustées au modèle photosynthétique de Farquhar qui fournit les variables \( V_{c_{max}} \) et \( J_{c_{max}} \).

Résultats

1) Les plantes ayant eu une croissance sous atmosphère enrichie en CO₂ expriment des paramètres photosynthétiques plus faibles. Notamment, l'assimilation à atmosphère ambiante est réduite de 20%. Il y a donc acclimatation de la photosynthèse.
2) Cependant, l'effet «sol» que l'on assimile à un effet long terme du CO₂ induit une augmentation de l'assimilation mesurée sous atmosphère ambiante. Cette augmentation est de même ampleur que celle induite par la fertilisation azotée. Les traitements S+ et N+ donnent donc en partie l'acclimatation induite par le traitement A+.
3) Tous traitements confondus, l'assimilation mesurée à atmosphère ambiante est corrélée fortement à la teneur en azote des feuilles. Nous faisons donc l'hypothèse que le traitement S+ favorise le maintien de la teneur en azote foliaire.

Conclusion

L'acclimatation négative de la phosynthesis que l'on observe sur les plantes exposées à une atmosphère enrichie en CO₂ est tempérée sur sol ayant été lui même exposé à un enrichissement en CO₂. Cet effet positif du sol enrichi semble être lié à une fourniture d'azote plus grande. Cette étude met donc l'accent sur la nécessité de travailler sur des systèmes couvert/sol si l'on veut prédire la réponse des écosystèmes terrestre au changement climatique.
Appendix 4: Poster presented by Gregor Yeates at British Ecological Society Annual Symposium "Biological Diversity and Function in Soils" March 2003, Lancaster University, England. (See next page)
Elevated carbon dioxide has marked effect on soil microfauna under grazed pasture on sand

Introduction

Increasing levels of atmospheric CO₂ affect plant tissue quality and its decomposition. To assess long-term implications we need to understand impacts of elevated CO₂ on soil processes, Free-air CO₂ enrichment (FACE) technology allows us to manipulate CO₂. A FACE array in a pasture permits alteration of total labile soil organic matter in a few years. Assessing the mechanisms underlying such changes is also critical in predicting responses in plant growth and below-ground carbon storage.

Methods

• >30-year-old pasture on Pukewatu black sand
• 6 rings, each 12 m in diameter
• 3 ambient controls (360 µl CO₂ l⁻¹)
• 3 enriched rings maintained at 475 µl CO₂ l⁻¹ during photoperiod since October 1997
• Baseline nematode samples collected from each ring in May 1997; ‘equilibrated’ samples collected May, August, November 2001, February 2002
• 5 samples per ring collected each time from 0-10 cm soil depth
• Soil nematodes extracted using the tray method
• Root growth measured in ingrowth cores

Results

• We found significantly more (1.45x) nematodes in the elevated CO₂ pasture rings than in the ambient control rings after enrichment (Table 1). Rotifers also increased (4.0Bx) but enchytraeids were depressed
• Eleven of 21 abundant nematode taxa had significantly higher populations in rings under elevated CO₂ (Table 2). Only the bacterial-feeding Cervidellus had a lower abundance m⁻² under elevated CO₂
• The greatest nematode response to elevated CO₂ was by the root-feeding Longidorus elongatus (Figure 1). Its mean population increased from 87 to 372 per 100 g soil (4.26x) (Table 2)
• The greatest response of bacterial-feeding nematodes was a 2.03x increase in Alomus. Alomus was the only nematode to show a significant, positive response in related growth chamber experiments. It may feed on dissolved organic matter
• The abundance of each of the six nematode feeding groups showed a significant, positive response to elevated CO₂ (Table 3). These responses differ with trophic level
• Root mass was 21% lower under elevated CO₂ but root growth rate 22-37% greater. Thus root turnover was greater (i.e. residence time was lowered) (Table 4)
• Other responses to elevated CO₂ found in these FACE rings include: uncharged microbial biofilms and N-accumulation
• increased earthworm abundance and biomass (see Box A)

Conclusions

• Increase of atmospheric CO₂ from ambient to 475 µl l⁻¹ was associated with significant changes in the soil fauna
• The 4.3% increase in the root-feeding nematode Longidorus was associated with increased root turnover under elevated CO₂. This suggests a significant increase in the partitioning of photosynthate below-ground
• The uncharged soil microbial biomass and doubling of populations of omnivorous and predacious nematodes suggest a multiplicative response in CO₂ and greater fluxes through soil biota
• Better understanding of plant-microbial-microfaunal interactions in soil is necessary. Until the processes are better understood, the impact of increasing atmospheric CO₂ levels on net plant productivity and carbon sequestration in soils cannot be evaluated

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