High potential, but low actual, glycine uptake of dominant plant species in three Australian land-use types with intermediate N availability

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Abstract The traditional view of the nitrogen (N) cycle has been challenged since the discovery that plants can compete with microbes for low molecular weight (LMW) organic N. Despite a number of studies that have shown LMW organic N uptake by plants, there remains a debate on the overall ecological relevance of LMW organic N uptake by plants across ecosystems with different N availabilities. We here report patterns of glycine N uptake by plants from three different Australian land-use types with intermediate N availability and low inherent glycine concentrations in the soil. Using 15N labeled tracers, we tested the potential of these plants to acquire glycine in ex-situ laboratory experiments and attempted to validate these results in the field by determining actual uptake of glycine by plants directly from the soil. We found in the ex-situ experiments that plants from all three land-use types were able to take up significant amounts of glycine. In contrast, glycine uptake directly from the soil was minimal in all three land-use types and 15N tracers were largely immobilized in the soil organic N pool. Our study confirms that the potential for LMW organic N uptake by plants is a widespread phenomenon. However, our in-situ experiments show that in the three land-use types tested here plants are inferior competitors for LMW organic N and rely on NH4+ as their main N source. In contrast to several previous studies in arctic, alpine and even temperate ecosystems, our study suggests that in ecosystems with intermediate N availability, mineral N is the plants' main N source, while LMW organic N is of less ecological relevance to plant N nutrition.

Keywords Amino acids · Glycine · Microbial competition · Mineral nutrition · Stable isotopes

Introduction

The nitrogen (N) cycle is an essential element of ecosystem biogeochemistry (Vitousek and Howarth 1991; Vitousek et al. 1997b). A profound understanding of the N cycle is therefore important to evaluate and predict the consequences of globally increasing N deposition for the structure and function of ecosystems (Vitousek et al. 1997a). Historically, microbial N
mineralization in the soil had been viewed as the most critical aspect of the N cycle. In this traditional concept only N that was mineralized by microbes in excess of their own demand and released back into the soil (i.e. net N mineralization) was available to plants. As a consequence, net N mineralization was considered the key process that controlled N availability for plants (Haynes and Goh 1978; Kinzel 1982; Runge 1970). This classic view of the N cycle has been challenged over the past years by a number of studies that have shown the potential of plants to circumnavigate the mineralization bottleneck in the N cycle by effectively taking up intact forms of low molecular weight (LMW) organic N, such as amino acids (Bardgett et al. 2003; Chapin et al. 1993; Näsholm et al. 1998; Schimel and Bennett 2004).

Originally, LMW organic N uptake by plants was tested under laboratory conditions. These experiments revealed that the potential for LMW organic N uptake is a widespread phenomenon of plant species from a broad range of different ecosystems (Falkengren-Gergerup et al. 2000; Finzi and Berthrong 2005; Kielland 1994; Raab et al. 1999; Schmidt and Stewart 1999; Warren and Adams 2007; Weigelt et al. 2005). Plants in laboratory experiments are, however, removed from their ecological context where they have to cope with microbial competition, abiotic immobilization of nutrients and where they are deprived of their mycorrhizal associations. A number of studies have therefore tested the uptake of LMW organic N by plants directly from the soil, either in pots or in-situ in intact ecosystems. To date we have evidence for LMW organic N uptake directly from the soil by plants from arctic (Clemmensen et al. 2008; Nordin et al. 2004), alpine (Miller et al. 2007), tundra (McKane et al. 2002; Schimel and Chapin 1996) and boreal ecosystems (Näsholm et al. 1998; Nordin et al. 2001; Persson et al. 2003) as well as from improved and unimproved temperate grasslands (Bardgett et al. 2003; Harrison et al. 2007; Streeter et al. 2000) and agricultural soils (Näsholm et al. 2000, 2001; Xu et al. 2008).

Despite a growing body of literature that reports LMW organic N utilization by plants, the ecological relevance of LMW organic N uptake by plants in different ecosystems is still controversial (Jones et al. 2005). In particular, strong microbial competition for mineral N, and especially for LMW organic N, seems to conflict with observations of plant LMW organic N uptake (Finzi and Berthrong 2005; Hodge et al. 2000; Jackson et al. 1989; Kaye and Hart 1997; Lipson and Monson 1998; Lipson et al. 1999; Schimel et al. 1989; Sorensen et al. 2008). Compared to plant roots, microbes have better substrate affinity, higher mobility in the soil and a higher surface to mass ratio and should therefore out compete plants for LMW organic N (Owen and Jones 2001). Schimel and Bennett (2004) therefore proposed that the main form of N taken up by plants varies across a gradient of increasing N availability and microbial competition and depends on the distribution of N-rich micro sites in the soil. For example, in extremely N poor ecosystems such as some boreal, arctic or alpine ecosystems as well as in acidic grasslands decomposition and mineralization rates are low so that plants and microbes have to compete for LMW organic N to meet their N demands. Given their competitive strength microbes would win the majority of the competition for amino acids while plants would capture only a small but constant proportion of LMW organic N. However, the slow growth and longevity of plants in these ecosystems combined with their potential to effectively retain N could allow a small but consistent uptake of LMW organic N to meet the overall N demand of these plants. In ecosystems with greater N availability, microbes would be less N limited and begin to mineralize organic N to NH$_4^+$. However, mineralization would be restricted to N-rich micro sites and LMW organic N and NH$_4^+$ would diffuse away from the mineralization micro sites into the soil where plants and microbes would actively compete for it. Under such conditions, plants would have access to some LMW organic N but their nutrition would mainly rely on NH$_4^+$. With increasing N availability, microbial demand would be increasingly satisfied by LMW organic N in the microsites, reducing the competition between plants and microbes for diffusing NH$_4^+$. Eventually denitrifiers would become established and plant N capture would increasingly rely on NH$_4^+$ and NO$_3^−$ as suggested in the classic view of the N cycle (Schimel and Bennett 2004).

The large number of studies that have reported LMW organic N uptake by plants in arctic, alpine, boreal ecosystems as well as in unimproved grasslands on acidic soils (see references cited above) give credible evidence for the low end of the N availability gradient along Schimel and Bennett’s model. Also, Nordin et al. (2001) and Bardgett et al. (2003) demonstrated that the main form of N taken up by plants does in fact vary across a gradient of increasing N availability as hypothesised. In their studies LMW organic N uptake
decreases at the expense of inorganic N uptake along a boreal forest productivity gradient and along the transition of an unimproved acidic soil to agriculturally improved grassland, respectively. Alternatively, studies made at the high N end of the N availability gradient have shown LMW organic N uptake by plants in heavily fertilized agricultural ecosystems, where competition between plants and microbes for soil N is limited (Näsholm et al. 2001; Xu et al. 2008). Given ample evidence from either end of this N availability gradient, it is remarkable that very little information for LMW organic N uptake by plants is available from ecosystems with intermediate N availability where net N mineralization occurs (in contrast to high latitude or alpine ecosystems) but where ecosystems are not N saturated (in contrast to agricultural ecosystems or locations with large inputs from atmospheric deposition). For a full understanding of the underlying basic ecological mechanisms, as well as for the overall ecological relevance of organic LMW N uptake for plants across a broad range of ecosystems, investigations in ecosystems with intermediate N availability are now critically important.

In the study presented here, we tested LMW organic N uptake of plants in relation to mineral N uptake in three different Australian land-use types with intermediate N availability: a Pinus radiata plantation, a Eucalyptus globulus plantation and a grassland pasture. Specifically, we used the amino acid glycine as an indicator of LMW organic N uptake and tested (1) the overall potential of the plants to take up glycine relative to mineral N in ex-situ root incubation experiments. Further, (2) we determined actual glycine uptake relative to mineral N uptake directly from the soil in the presence of abiotic immobilization and microbial competition. Finally, (3) we determined the recovery rates of glycine $^{15}$N tracers over time, as well as mineral $^{15}$N tracers, in different compartments of the ecosystems such as plant roots, mineral N pools and the immobile soil organic N pool.

Methods

Study area

The study area was located near Dereel (37° 49'S, 143° 45'E), 20 km south of Ballarat in the state of Victoria in south-eastern Australia. The climate in this area is temperate, with an annual rainfall of 700 mm and mean monthly maximum temperatures ranging from 11–25°C. Sample plots (ca 0.25 ha, $n=5$) were established in three adjacent land uses types that are common the south eastern part of Australia: a perennial grassland pasture, a P. radiata plantation established on pasture in 1994 and an E. globulus plantation established on pasture in 1999. The three land-use types were all located within a 50 m radius. The grassland was an improved pasture that was dominated by the grass species Agrostis capillaris L., Holcus lanatus L. and Anthoxanthum odoratum L. and contained a number of native Australian grassland species (Hypericum graminea G. Forst, Aceana echinata Nees, Austrodanthonia sp) and typical pastoral herbs (Hypochaeris radiata L., Taraxacum officinale Weber ex F.H.Wigg, Trifolium sp. and Oxalis sp.). The two plantations were both established after soil ripping, mounding and an initial fertiliser addition. By the time of the experiment both plantations had achieved a closed canopy with an approximate height of 12 m (pine) and 15 m (eucalyptus). The topography is gently undulating, and the soil type in the upper 10 cm is classified as a silt loam (USDA) with 11–16% clay and 10–17% silt.

Uptake of mineral and glycine $^{15}$N by roots

In February 2005, root uptake of $^{15}$NO$_3^-$, $^{15}$NH$_4^+$ and $^{15}$N-glycine by E. globulus (Eucalyptus plantation), P. radiata (Pine plantation) and grassland plants (not separated by species) was studied using three different approaches. We studied the $^{15}$N uptake of 1) excavated but intact roots in the field (ex-situ attached roots), 2) excavated and detached roots in the lab (ex-situ excised roots), and 3) intact and in-situ rhizosphere roots by injecting $^{15}$N labeled solutions into the soil (in-situ ecosystem N uptake). We specifically applied three different methods to test for different aspects of root $^{15}$NO$_3^-$, $^{15}$NH$_4^+$ and $^{15}$N-glycine uptake. With the two ex-situ methods we tested the plant’s potential uptake capabilities for the different N species when provided in solution at similar concentrations. In addition, we tested the plant’s actual N uptake patterns under more realistic in-situ conditions where plants have to cope with abiotic immobilization and microbial competition, as well as the different concentrations the three N species are available at. Therefore, the concentrations of the $^{15}$N species injected into the soil in the in situ
experiments were adjusted to only fractionally increase (~10%) the in-situ concentrations of NO$_3^-$, NH$_4^+$ and glycine measured in the soil at the time of the experiments (Murphy et al. 2003).

N uptake by excavated but intact roots

To study the $^{15}$N uptake of attached roots we carefully excavated root tips of *P. radiata* and *E. globulus* in the upper 5 cm of the soil in each of the five blocks in the two plantations, (one root tip for $^{15}$NO$_3^-$, $^{15}$NH$_4^+$ and $^{15}$N-glycine uptake, replicated five times in each of the two plantations). Excavation of individual roots in the grassland was not possible because plants developed a thick root layer and roots that were too fragile to be excavated while still attached to the plant. After washing the excavated roots with water, root tips were incubated in 10.0 mL of nutrient solution. The solution contained 60 atom% $^{15}$NO$_3^-$, 60 atom% $^{15}$NH$_4^+$, 99.9 atom% $^{15}$N-glycine. The roots were simultaneously offered all three N sources in one solution at a concentration of 100 $\mu$mol N L$^{-1}$ each, to provide comparable access to different N sources, whilst only $^{15}$N labeling one N source. In addition, N uptake solutions contained 10 mg L$^{-1}$ ampicillin to minimize microbial activity in the uptake solutions and 100 $\mu$mol L$^{-1}$ CaCl$_2$ for membrane stability. Ampicillin has been shown to effectively inhibit bacterial growth and N mineralization in similar experiments (Warren 2006; Warren and Adams 2007). We are therefore confident that the $^{15}$N recovered in plant roots in the $^{15}$N-glycine treatment originated from the uptake of intact glycine. After 2 h of incubation, the submerged root tip, and the first 10 mm of the unsubmerged root, were cut and washed in 50 mM KCl and then deionized water. Roots were then dried at 80°C for 48 h, ground to a fine powder and analyzed by IRMS.

N uptake by excavated and detached roots

The $^{15}$N uptake of excised roots was studied on the same day as $^{15}$N uptake of attached roots. We collected five soil cores (50 × 50 mm) in each of the three different land use types. The soil cores were kept on ice whilst transported back to the laboratory within 2 h of sample collection and roots were separated from the soil by sieving and washing in deionised water. From each core roots were then incubated in 10 mL of the same labeled nutrient uptake solution as described above. After 1 h of incubation the roots were washed with 50 mM KCl and deionised water to remove excess nutrient solution, dried at 80°C for 48 h, ground to a fine powder and analyzed by IRMS.

N uptake by intact and in-situ roots

The uptake of $^{15}$N by roots from their rhizosphere environments was studied in the same week as the other uptake approaches. Fifteen large (250 mm diameter) plastic soil collars were installed in each of the three land-use types. The litter layer around the circumference of the collar was cut, and the collar slightly inserted into the soil so that the litter-fermentation layer remained intact and the roots were not disturbed inside the collar. In order to create comparable soil moisture conditions within, but also across, the three different land-use types and to minimize disturbance from the injection of N solutions into the soil, the collars were slowly (over 1 h) irrigated with 9.0 L of deionized water 3 days before the experiment, so that the upper five cm of soil was saturated and could then drained back to field capacity. Three days later, two soil cores (50 mm diameter, 50 mm deep) were inserted into the soil inside each collar. One core was sampled immediately to determine soil water content (48 h at 105°C), inorganic N concentration (1:4, soil:1 M KCl, m/v), and the natural abundance $^{15}$N signature of total soil N and root samples ($t=0$). The other soil core remained in-situ, and was sampled after 168 h (i.e. after the in-situ N uptake experiment) and processed as before for soil water content and inorganic N concentration to calculate net ammonification and nitrification rates for the time of the experiment (for details see below).
diameter, 50 mm deep) were injected in each collar with a solution containing three N species: NO$_3^-$, NH$_4^+$, and glycine. There were three versions of this triple N-mix solution where only one of the three N species was $^{15}$N enriched. The solutions spiked with $^{15}$N-NO$_3^-$ and $^{15}$N-NH$_4^+$ were enriched to 60 atom % $^{15}$N, whereas the solution spiked with $^{15}$N-glycine was enriched to 99.9 atom % $^{15}$N. Within a triple-mix solution, the concentrations of N for three N sources were: 0.71 mM for NH$_4^+$, 0.071 mM for NO$_3^-$ and 0.071 mM for N-glycine. In contrast to the solutions used in the ex-situ experiments described above, these solutions did not contain ampicillin or CaCl$_2$ as we did not want to alter soil microbial activity, or need to ensure membrane stability. We injected 5 mL of solution at three depths (1, 2.5 and 4 cm) through five injection points into each soil column (1 mL per injection point). This resulted in the application of 50µg of NH$_4^+$, 5µg of NO$_3^-$ and 5µg glycine-N to the soil column, equivalent to 25.5, 2.5 and 2.5 mg N m$^{-2}$, respectively. N concentrations in the solution were specifically chosen so that the N added to the soil would only increase the existing soil N pools by about 10% and therefore not dramatically alter the natural concentrations of NH$_4^+$, NO$_3^-$ and glycine in the soil. The injected soil volumes were not disturbed by inserting a core so that root uptake and rhizosphere microbial function could continue. Instead, three small plastic markers were inserted at equi-distant points along the circumference of each soil volume to record their exact locations. In total, for every land-use type there were five collars with three $^{15}$N-NO$_3^-$ labeled soil columns, five collars with three $^{15}$N-NH$_4^+$ labeled soil columns and five collars with three $^{15}$N-glycine labeled soil columns. From each collar, we sampled these soil volumes in sequence, at 6 h, 24 h and 72 h after $^{15}$N solution injection. Samples were kept on ice and transported to the laboratory within 2 h. Roots were separated from soil, washed, dried and analyzed for $^{15}$N content as described above.

To calculate N uptake rates by these in-situ roots, we determined µmol $^{15}$N excess in the labelled roots compared to µmol $^{15}$N in the control roots taking the total amount of $^{15}$N tracer injected into account. In addition, we accounted for the dilution of the injected $^{15}$N tracers by the pools and fluxes of NO$_3^+$, NH$_4^+$ and glycine in the soil during the incubation. This method of determining N uptake rates allows calculating N uptake rates that are independent of the amount of tracer added to the ecosystem. For a detailed description of the calculations see Kahmen et al. (2006).

Soil N pools and fluxes

Soil N dynamics in the three land-use types were in detail assessed for the time that we investigated plant N uptake patterns. In particular, net nitrification and ammonification were measured from the difference in soil NO$_3^-$ and NH$_4^+$ concentrations in the soil cores sampled at the beginning of the N uptake experiment ($t=0$) and from soil cores left in-situ and sampled 168 h after the beginning of the experiment ($t=168$) following a method described by (Adams and Attiwill 1984). Soil cores were processed for soil water content and inorganic N extraction (1:4, soil:1 M KCl, m/v). Soil KCl extracts were shaken for 1 h at 250 rpm before being filtered (Whatman 42) and the filtrate analyzed for NO$_3^-$ and NH$_4^+$ concentration using a segmented continuous flow auto-analyser (Technicon™). Net nitrification and ammonification were calculated as:

$$N_{net} = \frac{N_{t1} - N_{t0}}{t}$$

where $N_{t1}$ is the NO$_3^-$-N concentration (mg kg$^{-1}$) at the end of the in-situ containment, $N_{t0}$ is the NO$_3^-$ concentration at the start of the in-situ containment and $t$ is the number of days.

To calculate rates of gross nitrification and ammonification and to determine the amount of $^{15}$N tracer remaining in the mineral N soil pool over time we used the $^{15}$N labelled soil columns sampled for root N uptake in the in-situ experiment at $t=24$ and $t=72$. After roots were removed from the soil columns, the soil was extracted with 100 mL 1M KCl as described above and then the solution filtered directly into 500 mL glass jars. A 5 mL sub-sample of the KCl filtrate was removed to determine NH$_4^+$-N and NO$_3^-$-N concentrations as described before. The remaining KCl filtrate (~80 mL) received an excess addition of MgO (~0.3 g) to raise solution pH above 10, so that NH$_4^+$-N was volatilized to NH$_3$. Immediately after adding MgO, the lid of the glass jar was sealed. Underneath of the lid (inside the jar) was a stainless-steel hook with an acidified (2.5 M, KHSO$_4$) filter paper disc attached. The volatilized NH$_3$ was captured on to the acidified disc through conversion back to NH$_4^+$ over 7 days of enclosed micro-diffusion (Brooks et al. 1989). After 7 days, the jar was shortly opened,
the filter paper removed and dried in a dessicator. Twenty-four hours later, excess Devarda’s alloy (~0.20 g) was added to reduce the remaining NO₃⁻ to NH₄⁺, which was subsequently converted to NH₃ and also volatilised and captured on a fresh acidified filter paper disc. The NH₄⁺-N and NO₃-N on separate, dried filter paper discs were analysed for ¹⁵N atom-% enrichment by IRMS.

Gross nitrification and ammonification were calculated using the change in NO₃⁻ and NH₄⁺ pool size and atom-% enrichment between soil cores sampled 24 and 72 h (Murphy et al. 2003) after the injection of ¹⁵NO₃⁻ or ¹⁵NH₄⁺ tracer based on ¹⁵N isotope pool dilution as:

\[ N_2^* = N_1^* \left(1 + \frac{\theta t}{N_1}\right)^{R/\theta} \]  

where \( N \) is the labelled nitrate pool (mg kg⁻¹), \( t \) is incubation time (days), \( R \) is the rate of nitrification, * indicates atom-% excess ¹⁵N, \( N_1 \) is the first time of sampling, \( N_2 \) the second and \( \theta \) is the rate at which the labelled nitrate pool changed (Barraclough 1991).

In addition to the mineral soil N pools and fluxes, we determined glycine concentrations in the soils of the three land-use types. The original attempt to determine soil glycine concentrations during the experiment failed, so we repeated this under comparable environmental conditions in February 2005. From each land-use type we collected 10 soil samples from 0–5 cm depth using a soil core (4.5 cm diameter, 0–5 cm depth). Glycine was extracted using 1 M KCl as described for mineral N above. Soil glycine concentrations were analyzed with a post-column ninhydrin-derivatization high-performance liquid chromatography-based amino acid analyzer (Model 6300; Beckman Instruments, Palo Alto, CA, USA) according to Aidar et al. (2003).

To place the soil N pools and fluxes measured during this experiment in the context of overall seasonal dynamics of mineral soil N pools in the three land-use types, we collected composite soil samples (i.e. five subsamples were combined to a single sample) from each of the three land-use types to investigate the seasonal variability in soil NO₃⁻ and NH₄⁺ pool size on a monthly basis between October 2004 and March 2005. The soil samples were sieved (2 mm) and subsamples taken for mineral N extraction (1:4, soil:1 M KCl, m/v) and gravimetric water content as described above. The samples collected in October and November 2004 were also analyzed for total soil C and N and soil pH. Samples were air-dried, sieved (2 mm) and ground (Retsch™, Haan, Germany) and analyzed for total C and N concentration by dry combustion in a CHN elemental analyzer (LECO™ CNH-2000, St. Joseph, MI, USA). Soil pH was measured in deionised water at a 1:5 soil:water ratio.

We determined soil bulk density in all three land-use types in order to relate the soil N pools and fluxes to the amount of tracer N that we injected into a given soil volume. Immediately after the collection of all in-situ root N uptake samples we collected three soil samples at 0–5 cm using a stainless steel soil core (Ø 70 mm, height 50 mm) adjacent to each collar. Samples were oven dried at 105°C for 48 h and then weighed to determine dry soil mass per unit volume (g cm⁻³).

Microbial immobilization and recovery of ¹⁵N in different ecosystem N pools

We determined microbial and abiotic immobilization of the ¹⁵N tracers injected to the soil. After the \( t=6 \), \( t=24 \) and \( t=72 \) h soil samples (25 g) had been extracted with 100 mL of 1 M KCl (see above), the extraction was repeated twice so that all trace of inorganic N had been removed from the soil residue (Recous et al. 1999). The triple-KCl extracted soil residue was then rinsed with 100 ml of deionised water, centrifuged and the solution discarded and dried at 60°C for 48 h. Sub-samples of the dried soil residue were fine-ground on the Retsch™ ball mill and analysed for ¹⁵N atom-% enrichment using the IRMS.

To determine the recovery rates of ¹⁵N tracers in plant roots, mineral N and soil organic N we calculated µmol ¹⁵N excess in labelled plant roots, mineral soil N and soil organic N as compared to µmol ¹⁵N in the unlabeled controls (\( t=0 \)) and expressed the recovery rates as percent of the ¹⁵N tracer originally injected into the soil.

Statistics

We used one-way ANOVAs with Turkey’s post-hoc tests to determine separately for the individual land-use types a) statistical differences for the uptake of a given N species across different ‘N-uptake methods’, and b) statistical differences among NO₃⁻, NH₄⁺ and glycine uptake for a given ‘N uptake method’. In both tests, we used N uptake rate of a given N species as dependent variable and either ‘N uptake method’ (a) or ‘N species’ (b) as factor.
Results

NO$_3^-$ and glycine concentrations were at least one order of magnitude smaller than NH$_4^+$ concentrations across the three land-use types at the time we assessed root N uptake patterns (Fig. 1). Similarly, gross and net ammonification were more than one order of magnitude higher than gross and net nitrification in all three land-use types (Table 1). Gross and net ammonification was highest in the pasture and lowest in the E. globulus plantation. Likewise, gross and net nitrification was extremely low in the E. globulus plantation and only slightly higher in the pasture and P. radiata plantation. These patterns are consistent with the seasonal trend that we observed for mineral soil N pools in the three land-use types. Although there was some seasonal variability over the growing season, NH$_4^+$ dominated the plant available mineral soil N pool throughout the year in all land uses (Fig. 2). Overall, this suggests that NH$_4^+$ is the dominant form of plant available N in these ecosystems.

Soil pH ranged from 4.33 in the P. radiata plantation to 4.52 in the E. globulus plantation and 4.89 in the pasture. Soil C was significantly higher in the P. radiata and E. globulus plantations (5.76% and 5.03%, respectively) as compared to the pasture (4.21%). In contrast, soil N was comparable across all three land-use types, ranging from 0.42% in the P. radiata plantation to 0.48% in the E. globulus plantation (Table 1).

When uptake of different N species was tested using excavated attached roots, P. radiata and E. globulus relied exclusively on NH$_4^+$ and glycine, and showed only marginal NO$_3^-$ uptake (Fig. 3a). Laboratory incubations of excavated and detached roots revealed that roots from the pasture took up all three N species with a preference for NH$_4^+$, while P. radiata and E. globulus again only took up NH$_4^+$ and glycine, not NO$_3^-$ (Fig. 3b). In both ex-situ N uptake experiments, glycine was a substantial N source for plant roots from all three land-use types. In contrast, when N uptake directly from the soil was tested in-situ with intact roots, NH$_4^+$ was the exclusive N source for plants from all three land-use types, while neither NO$_3^-$ nor glycine contributed to a plant’s N uptake in significant amounts in any of the three land-use types (Fig. 3c).

Root uptake rates of NO$_3^-$, NH$_4^+$ and glycine were in the same order of magnitude in all three N uptake experiments, independent of method (Fig. 3). However, we found marked differences with regard to the relative amount of different N species acquired by plant roots depending on the method used to measure N uptake (Fig. 4). When N uptake was measured ex-situ, using either attached or excised roots, glycine contributed significantly to overall N uptake, ranging from 63.2% for attached P. radiata roots to 20.8% for excised E. globulus roots. In contrast, when N uptake from the soil was measured for roots in-situ NH$_4^+$ dominated the N taken up, while glycine uptake declined to 4.7% for E. globulus, 2.7% for P. radiata and 1.7% for pasture roots of overall N uptake (Fig. 4). Each of the three methods used to measure N uptake by plant roots influenced the perceived plant N uptake pattern, a finding that was consistent across all three land-use types: organic N uptake declined at the expense of NH$_4^+$ uptake from ex-situ attached roots, to ex-situ excised roots and finally to in-situ direct N uptake from the soil (Fig. 4).

Total recovery of the $^{15}$N label injected into the soil of the three land-use types ranged from between 45.7% (P. radiata, $^{15}$NO$_3^-$ treatment) and 119.1% (pasture, $^{15}$NO$_3^-$ treatment) with mean recovery rates of 65.8%, 85.9% and 59.2% for the E. globulus, pasture and P. radiata land-use type, respectively. $^{15}$N tracers were recovered in all soil N pools except for the NO$_3^-$ pool, which was very low to non-detectable (Fig. 5). Of the recovered $^{15}$N label, the largest proportion was recovered in the insoluble organic N soil pool (Fig. 5). Substantial amounts of


\[ 15\text{N} \text{ tracer were, however, also recovered from the mineral N pool (i.e. the soil NH}_4^+ \text{ pool). The recovery rates in the mineral N pool in all three land-use types declined over time at the expense of the insoluble organic N pool, which increased over time. In contrast to the insoluble organic N pool and the mineral N pool, the proportion of label recovered in the roots was small, ranging from 1.2–3.0% in } E. \text{ globulus roots, 3.6–12.1% in pasture roots and 1.0–3.7% in } P. \text{ radiata roots (Fig. 5). For roots, we were not able to detect an increase or decrease in recovered } 15\text{N} \text{ label over time.} \]

**Discussion**

Ammonium and glycine were the dominant forms of root N uptake in both, the excavated attached roots and the excavated detached roots experiment (Fig. 4). The high levels of glycine uptake that we detected using ex-situ N uptake methods show the potential of plant species from all three land-use types to use LMW organic N as an N source. Our findings are consistent with a broad range of other studies that have found LMW organic N uptake from solutions (Falkengren-Grerup et al. 2000; Finzi and Berthrong 2005; Kielland 1994; Raab et al. 1999; Schmidt and Stewart 1999; Warren and Adams 2007). Our study therefore supports the suggestion that the potential for LMW organic N uptake by plant roots is a widespread functional trait of plants from different ecosystems around the globe.

The uptake rates for NO\(_3^-\) from the solutions were minimal for } E. \text{ globulus and } P. \text{ radiata in both ex-situ experiments compared to NH}_4^+ \text{ and glycine uptake. Only the pasture roots showed some NO}_3^- \text{ uptake (Figs. 3 and 4). Low levels of NO}_3^- \text{ uptake reflect the small NO}_3^- \text{ pools and fluxes that we measured in the soils of all three land-use types (Figs. 1 and 2, Table 1) and could therefore be the result of root adaptation to high NH}_4^+ \text{ and low NO}_3^- \text{ concentrations in these soils. Also, the presence of NH}_4^+ \text{ and amino acids has been shown to inhibit root uptake of NO}_3^- \text{ from solution, which could also explain the low levels of NO}_3^- \text{ uptake in our ex-situ experiments (Haynes and Goh 1978; Thornton and Robinson 2005).} \]

Both ex-situ methods that we used to determine the potential of plants for glycine uptake have several
disadvantages that can impact the N uptake rates of roots (Lucash et al. 2007). For excised roots it has been argued that 2 h after a root has been cut, soluble sugar concentrations and metabolic activity in that root sharply decline which affects the capacity for active uptake of compounds and may lead to an underesti-
mation of root N uptake capacity (Farrar 1985). Similarly, it has been estimated that measuring nutrient uptake on roots that remain attached to a tree could underestimate N uptake rates by 10% as some of the N is transported away from the root towards the trunk of the tree (Hawkins et al. 2005; Warren and Adams 2007). It is noteworthy that the general N uptake patterns we obtained from both ex-situ experiments are similar, given the uncertainties regarding both methods in determining N uptake from solution and despite the higher relative NH$_4^+$ uptake in excised roots as compared to attached roots. The magnitude of absolute NH$_4^+$ and glycine uptake, as well as the general patterns with respect to N preferences of the different species, is comparable in both ex-situ methods; both E. globulus and P. radiata show a clear overall preference for NH$_4^+$ and glycine uptake, but neither tree species takes up much NO$_3^-$ (Figs. 3 and 4).

Measuring the uptake of N from nutrient solutions by roots ex-situ from their soil environment provides important information on N uptake kinetics or the potential of plants to acquire LMW organic N. However, these ‘root ex-situ in solution’ studies reveal relatively little information on the ecological importance of LMW organic N uptake by plant roots (Jones et al. 2005; Näsholm et al. 1998; Owen and Jones 2001). We

Fig. 3 Uptake rates for NO$_3^-$, NH$_4^+$ and glycine for E. globulus, P. radiata and pasture plant roots determined with excavated but intact roots (a), excavated and detached roots (b) and intact rhizosphere roots by injecting $^{15}$N labeled solutions into the soil (c). The attached root method was not tested in the grassland (n.d.). We used one-way ANOVAs with Turkey’s post-hoc tests to determine separately for the individual land-use types a) statistical differences for the uptake of a given N species across different ‘N-uptake methods’ (capital letters), and b) statistical differences among NO$_3^-$, NH$_4^+$ and glycine uptake for a given ‘N uptake method’ (lower case letters). Error bars represent one standard deviation from the mean.

Fig. 4 Uptake rates of NO$_3^-$, NH$_4^+$ and glycine in relation to total N uptake in comparison of three different methods determining N uptake of E. globulus, P. radiata and grassland roots. A: excavated, attached roots, E: excavated and detached roots and ES: recovery of a $^{15}$N tracer from the soil. The attached root method was not tested in the grassland (n.d.)
therefore, determined in-situ N uptake patterns of intact plant roots in the three land-use types. Interestingly, glycine uptake in the in-situ experiment was extremely small, contributing between 2% and 5% to the overall N uptake of *E. globulus*, *P. radiata* or pasture grasses. In contrast, NH$_4^+$ was the dominant N source for plant roots contributing more than 93.5% to overall N uptake in all land-use types (Fig. 4). While our ex-situ experiments have clearly shown the potential for the plants to acquire LMW organic N in the form of glycine, the small uptake rates measured in-situ suggest that the actual ecological relevance of LMW organic N in the form of glycine may be limited in these three ecosystems—at least during the time of this experiment.

Small concentrations of glycine in the soil in combination with microbial competition and abiotic immobilization of LMW organic and mineral N could explain why plants that have a high potential for LMW organic N uptake show low in-situ uptake rates for glycine (Hodge et al. 2000; Jones et al. 2004; Kaye and Hart 1997). It has been shown for a cold temperate forests, for example, that more than 90% of $^{15}$N labeled glycine was recovered in either NH$_4^+$ or in the soil microbial biomass only 15 min after it was injected into the soil (Finzi and Berthrong 2005). Similarly, other studies have shown that microbes compete very effectively with plants for LMW organic N in temperate grassland ecosystems (Bardgett et al. 2003; Harrison et al. 2007) or subarctic tundra (Sorensen et al. 2008). In agreement with these previous studies, we recovered more than 70% of the $^{15}$N label in the insoluble soil organic N pool within 6 h of injection in all three land-use types and for all three labeled N species (Fig. 5). In contrast, we recovered only between 1.2 and 9.4% of $^{15}$N in plant roots 6 h after injection, suggesting that glycine, like mineral N, in these three ecosystems is rapidly immobilized by microbial uptake or abiotic processes, leaving only a small proportion available for root uptake. It is interesting to note, however, that the relative rates of immobilization were largely independent of N species as was the fraction of a given N species that was available for root uptake (Fig. 5). This suggests that not microbial preference for a given N species but the small concentrations of glycine and NO$_3^-$ that we found in the soil as compared to NH$_4^+$, explains why glycine and NO$_3^-$ contributed little to the overall N uptake of plants in the three ecosystems.
Our study provides evidence that N uptake patterns for mineral N and glycine largely reflect the availability of the different N species in the soil when tested with intact roots in-situ (Figs. 1 and 3). This suggests that in-situ N-uptake studies need to be carefully designed in order not to bias the observed N uptake patterns. In our experiment, we were particularly cautious not to alter the relative abundances of the different N species in the soil by injecting NH$_4^+$, NO$_3^-$ and glycine at concentrations that ranged roughly around 10% of the naturally occurring concentrations. In addition, we applied a novel method that allowed us to estimate the absolute uptake rates of different N species by accounting for the dilution of the tracer through net N mineralization fluxes as well as existing pools of the relevant N species (Kahmen et al. 2006; Kahmen et al. 2008). As such, the calculated N uptake rates are independent of the amount of tracer applied. To do so, we assessed concentrations of NH$_4^+$, NO$_3^-$ and glycine during the experiment as well as the net NH$_4^+$ and NO$_3^-$ fluxes. Since it was not possible to determine the net glycine fluxes in this experiment, we made the assumption that net glycine fluxes are in the same order of magnitude as net NH$_4^+$ fluxes. While this introduces uncertainty to our calculations of in-situ glycine N uptake, the net fluxes that we determined in this study were small compared to the respective pools (Fig. 1, Table 1). As a result, the error that our assumption introduces to our estimates of glycine uptake is unlikely to affect the overall patterns that we detected in this study. We acknowledge that methodological difficulties can introduce error to our estimates of N uptake, in particular determining the exact flux rates of NH$_4^+$, NO$_3^-$ and glycine (Näsholm et al. 2001). However, the method we used provides us with more realistic estimates of N uptake rates than the typically reported $^{15}$N recovery rates that are biased by the amount of tracer added to the ecosystem, and that are affected by the dilution of the tracer in the soil (Jones et al. 2005; Näsholm et al. 2001; Nordin et al. 2004).

In the study presented here we have used glycine as a proxy for root uptake of LMW organic N compounds. Glycine is, however, only one out of a large number of LMW organic N compounds that can be taken up by plants (Weigelt et al. 2005). Our study might therefore underestimate the overall contribution of LMW organic N compounds to plant N uptake in the three land-use types. However, the low abundance of glycine in the soil, the high net mineralization rates and the high rates of microbial immobilization observed provide little evidence that other LMW organic N compounds would be substantially more abundant in these soils and contribute differently to overall plant N uptake in the investigated ecosystems.

Uptake of intact LMW organic N directly from the soil is often studied using $^{13}$C and $^{15}$N labeled compounds (Näsholm et al. 1998). In our study, we used glycine that was only $^{15}$N labeled that does not allow to distinguish if the $^{15}$N has been taken up as intact organic compound or has been mineralized prior to uptake. However, this does not confound the findings of our study since organic N is of limited relevance for overall plant N uptake in all three investigated ecosystems. Further, given the large unlabeled C pool in the roots of plants compared with the small amount of $^{13}$C that would have been taken up in this study from $^{13}$C and $^{15}$N labeled glycine, we question if significant amounts of the $^{13}$C tracer could have been detected at all in this study (von Felten et al. 2008).

At first glance the data we present seems to contradict previous studies that have shown LMW organic N contributes to the overall N uptake of plants (McKane et al. 2002; Näsholm et al. 1998; Näsholm et al. 2001; Nordin et al. 2004; Persson et al. 2003; Xu et al. 2008). However, the glycine uptake patterns we report for these three Australian ecosystems fit well into Schimel and Bennett’s (2004) general conceptual model of LMW organic N uptake across ecosystems with varying N availability and microbial competition. While previous studies have reported LMW organic N uptake by plants either from arctic, alpine or boreal ecosystems where N mineralization rates and levels of plant available N are low or from N saturated agricultural ecosystems, where microbial competition for N is limited, the three land-use types we investigated in this study showed intermediate levels of N availability at the time of this experiment: ammonification and NH$_4^+$ pools in the soil were high compared to nitrification and NO$_3^-$ and glycine pools (Table 1, Fig. 2). Yet microbial competition and abiotic immobilization for LMW organic N as well as mineral N was still high as indicated by the high immobilization rates after in-situ application of tracers in all land-uses (Fig. 5). In accordance with Schimel and Bennett (2004) our study therefore suggests that under these conditions of intermediate N availability LMW organic N is of limited ecological relevance to the overall N nutrition of the dominant plant species.
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