TRANSFORMATIONS OF NITRATE ¹⁵N UNDER DIFFERENT FOREST HARVEST RESIDUE REGIMES IN A HOOP PINE PLANTATION IN AUSTRALIA

G. X. Pu*, P. G. Saffigna & Z. H. Xu

Cooperative Research Centre for Sustainable Production Forestry, Australia Australian School of Environmental Studies, Faculty of Environmental Sciences, Griffith University, Nathan, Queensland 4111, Australia

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PU, G. X., SAFFIGNA, P. G. & XU, Z. H. 2005. Transformations of nitrate ¹⁵N under different forest harvest residue regimes in a hoop pine plantation in Australia. A study was conducted to quantify the effects of harvest residue management on denitrification, leaching and immobilization of ¹⁵N-labelled nitrate applied at 20 kg N ha-1 to 1-year-old hoop pine (Araucaria cunninghamii) in subtropical Australia. The experiment was undertaken in PVC microplots of 235 mm diameter and 300 mm long, driven into the soil (Lithosol) to a depth of 250 mm. Three replications were undertaken for each of the seven treatments: control without any residue, ground and unground foliage at 20 Mg dry matter (DM) ha-1, ground foliage at 40 Mg DM ha-1, ground and unground branches at 40 Mg DM ha-1, and ground branches at 80 Mg DM ha⁻¹. In 15 days after simulated daily rainfall of 100, 50 and 25 mm respectively in the first three days, 6-26% of applied ¹⁵N was lost via denitrification, 14-35% was immobilized and 32-53% was leached. The treatment incorporating foliage materials lost more ¹⁵N (21–26%) via denitrification than other treatments. Measurement of ¹⁵N gases (15N₉+15N₉O) showed higher 15N gas emission on day 1, followed by low gas emissions thereafter. This study showed that significant amounts of mineral N could be lost through leaching and denitrification during plantation establishment.

Key words: Denitrification – leaching – immobilization – ${\rm ^{15}N_2}$ – ${\rm ^{15}N_2O}$ – water-soluble carbon

PU, G. X., SAFFIGNA, P. G. & XU, Z. H. 2005. Perubahan nitrat ¹⁵N di bawah rejim sisa tuaian hutan yang berlainan di dalam ladang pain di Australia. Kajian ini dijalankan untuk menilai kesan pengurusan sisa tuaian terhadap denitrifikasi, larut lesap dan ketakmobilan nitrat berlabel 15N (20 kg N ha-1) pada pokok pain (Araucaria cunninghamii) berumur satu tahun yang tumbuh di kawasan subtropika Australia. Ujian dijalankan dalam mikroplot PVC berukuran 235 mm diameter dan 300 mm panjang. Mikroplot ini dipacu ke dalam tanah (Litosol) sehingga kedalaman 250 mm. Tiga ulangan dijalankan untuk kesemua tujuh perlakuan: kawalan tanpa sisa, dedaun yang dikisar dibubuh pada kadar 20 Mg jirim kering (DM) ha-1, dedaun yang tidak dikisar dibubuh pada kadar 20 Mg DM ha-1, dedaun di tanah dibubuh pada kadar 40 Mg DM ha⁻¹, dahan yang dikisar dibubuh pada kadar 40 Mg DM ha⁻¹, dahan yang tidak dikisar dibubuh pada kadar 40 Mg DM ha-1 dan dahan yang dikisar dibubuh pada kadar 80 Mg DM ha⁻¹. Ujikaji ini didedah kepada hujan tiruan harian sebanyak 100 mm, 50 mm dan 25 mm untuk tiga hari berturut-turut. Lima belas hari selepas itu didapati 6%–26% daripada ¹⁵N yang dirawat hilang melalui denitrifikasi, 14%–35% ¹⁵N menjadi tak mobil dan 32%–53% ¹⁵N terlarut lesap. Perlakuan yang menerima dedaun kehilangan lebih

*Present address: Sustainable Land Management, Natural Resource Sciences, Department of Natural Resources and Mines, 80 Meiers Rd Block B, Indooroopilly QLD 4068, Australia. E-mail: grantpu88@yahoo.com.au ^{15}N (21%–26%) melalui denitrifikasi berbanding perlakuan lain. Kajian gas ^{15}N ($^{15}N_2 + ^{15}N_2O$) menunjukkan pelepasan gas ^{15}N yang lebih tinggi pada hari pertama kajian, diikuti oleh pelepasan gas yang rendah pada hari-hari seterusnya. Kajian ini menunjukkan bahawa jumlah bahan galian N yang bererti boleh hilang secara larut lesap dan denitrifikasi semasa penubuhan ladang.

Introduction

Both land management and climate can have a significant impact on the dynamics of terrestrial nitrogen (N) cycling processes such as denitrification, leaching and immobilization (Firestone 1982, Robertson & Tiedje 1988, Nadelhoffer *et al.* 1999). Considering the large number of denitrification studies in agricultural ecosystems (Firestone 1982, Bouwman 1998), few studies have been made in forest ecosystems, particularly in the tropics and subtropics (Carnol & Ineson 1999, Hall & Matson 1999, Priha & Smolander 1999).

Almost all of the new hoop pine (Araucaria cunninghamii) plantations in south-east Queensland, Australia are established on second-rotation sites. In subtropical Australia, hoop pine plantations have a much higher N requirement than plantations of Eucalyptus and Pinus species (Bubb et al. 1999, Prasolova et al. 2000). Therefore, minimizing N losses during harvesting and site preparation is likely to be an important management practice for the long-term sustainability of these plantations (Matson et al. 1987, Bubb et al. 1998a). Besides residue removal or burning, N losses from leaching or denitrification can occur in the inter-rotation period as a result of higher nitrification rates due to site disturbance caused by clear fell and site preparation (Robertson & Tiedje 1988, Hall & Matson 1999). Holt and Spain (1986) reported that soil organic C, total N, available P and exchangeable Ca, Mg and K in a mature hoop pine plantation were substantially lower than those in the adjoining rain forest area in tropical north Queensland, Australia due to harvesting removal and site preparation. Although all the new hoop pine plantations are now established on second-rotation sites, the effects of site management practices such as harvest residue management regimes on nutrient cycling and soil fertility status are poorly understood (Bubb et al. 1999). There is limited information on the impacts of harvest residues on N cycling processes such as denitrification, leaching and N immobilization of hoop pine plantations. Nitrogen losses via denitrification have been considered relatively unimportant in most forest ecosystems (Attiwill & Leeper 1987), but Palm et al. (1993) have reported that denitrification in tropical forest ecosystems, either disturbed or undisturbed, is higher than in temperate forest ecosystems. Several studies have shown a significant increase in N losses via denitrification in disturbed forest sites compared with undisturbed sites (Hulm & Killham 1988, Myrold 1988, Dutch & Ineson 1990). Verchot et al. (1997) have mentioned that nitrate loss in some forests is almost exclusively through denitrification.

Studies of Bubb *et al.* (1998a) and Prasolova *et al.* (2000) have shown that there is a significant amount of mineral N (80–120 kg N ha⁻¹) present during hoop pine plantation establishment. Of this amount, about 20–40 kg N ha⁻¹ exist as nitrate-N due to active nitrification. The objective of this study was, therefore, to quantify the effects of harvest residue management regimes on denitrification, leaching and immobilization of ¹⁵N-labelled nitrate applied at 20 kg N ha⁻¹ to a 1-year-old, second rotation hoop pine plantation in south-east Queensland, Australia.

Materials and methods

Site and soil properties

The experimental site was located in a hoop pine plantation within the State Forest of Amamoor (26 ° 16' S, 152 ° 37' E), south-east Queensland, Australia, where a second-rotation plantation had been established for one year. The soil was classified as Lithosol according to the FAO Soil Classification System. The contents of sand, silt and clay were 63, 29 and 8% respectively for the 0–100 mm soil and 57, 33 and 10% for the 100–200 cm soil. For the 0–100 mm soil, pH (1:5 H_2O) was 6.4, total N 0.21%, organic C 3.2%, total K 1308 mg kg⁻¹, electrical conductivity 0.13 dS m⁻¹. The corresponding values were 6.2, 0.19%, 2.0%, 1259 mg kg⁻¹ and 0.08 dS m-1 respectively for the 100–250 mm soil. The average temperature was 25 °C in summer and 14 °C in winter. The average annual rainfall was about 1200 mm (Bubb *et al.* 1998a) and more than half of rainfall occurred in summer (November–February) featured with heavy thunderstorms. During the period of this experiment (mid-summer 1997), an average daily evaporation rate of 7 mm was recorded and the maximum and minimum average temperatures were 32 and 23 °C respectively.

Experimental treatments and procedures

The experiment consisted of seven treatments and replicated three times using a randomized complete block design. They were (1) control, without any residues incorporated, (2) unground foliage materials (chopped to < 35 mm in length) applied at 20 Mg dry matter (DM) ha⁻¹, equivalent to the average rate of foliage materials remaining on the ground after harvesting of first-rotation hoop pine plantations in south-east Queensland (3) ground foliage materials (to pass a 2 mm sieve) at 20 Mg DM ha⁻¹, (4) ground foliage materials at 40 Mg DM ha⁻¹, (5) unground branch materials (chopped to a size of any dimensions < 35 mm) at 40 Mg DM ha⁻¹ equivalent to the rate of branch materials remaining on the ground after harvesting, (6) ground branch materials at 40 Mg DM ha⁻¹ and (7) ground branch materials (to pass a 2 mm sieve) at 80 Mg DM ha⁻¹. All residues were incorporated into the top 100 mm soil within the microplots. Total C and N were 410 and 10 g kg⁻¹ (C/N ratio 41:1) respectively for foliage materials, and 478 and 4.3 g kg⁻¹ (C/N ratio 111:1) for branch materials.

Soil was collected at two layers (0–100 and 100–250 mm) from 21 random positions using PVC tubes (235 mm diameter \times 300 mm long). Soil samples were bulked and homogenized to give one composite sample for each depth respectively. Soil from 100–250 mm was repacked into each PVC core positioned at the original sampling places. Soil from 0–100 mm was mixed with oven-dried (60 °C) hoop pine residues according to treatments as described above. Soil plus residue was then repacked into each PVC tube. The procedure ensured that the top 250 mm

soil in the PVC tube was mixed and replaced to simulate soil disturbance during the site preparation at the establishment of the hoop pine plantation.

Bromide (Br) was used as a companion tracer to monitor the potential path of nitrate movement in the soil profile (Smith & Davis 1974, Saffigna *et al.* 1977, Kessavalou *et al.* 1996, Turpin *et al.* 1999). The difference between the recovered Br and ¹⁵N was assumed to be ¹⁵N loss through denitrification. The percentage difference between applied and recovered Br was regarded as the equivalent leaching losses of applied nitrate N.

Adequate water was added to the soil inside the PVC tubes until the soil reached field capacity (~42% w/w) in the top 250 mm soil. Water was also applied to the soil surrounding the PVC tubes at the same rate as inside the PVC tubes. The ¹⁵N-labelled KNO₃ (98% ¹⁵N excess) at 20 kg N ha⁻¹ and bromide at 100 kg Br ha⁻¹ as solutions were evenly applied to the soil surface within the PVC tubes. Application of 20 kg N ha⁻¹ as ¹⁵N-labelled nitrate would represent the lower end of 20–40 kg NO₃-N ha⁻¹ present in the hoop pine soil during plantation establishment and would not be expected to interfere significantly with the soil mineral pools (about 100 kg N ha⁻¹) and fluxes (Bubb *et al.* 1998a). More water was added to simulate a rainfall event of 100 mm and the topsoil was saturated, which is common in summer in this area.

Evolved ¹⁵N gases were collected daily by covering the microplots (Pu *et al.* 1999) for three hours each day, assuming the ¹⁵N gas emission rate in the three-hour period was the average rate on that day. The gas was collected into a 10 ml venoject vial via a sampling port on the cover, after mixing all the air within the headspace of the microplot using a 10 ml syringe (Pu *et al.* 1999). The recovery of ¹⁵N in gas form was regarded as the result of direct denitrification. After collecting gas samples on day 2 and day 3, more water, equivalent to 50 mm and 25 mm of rainfall respectively, was added to the soil surface. The amounts of added water were to simulate the common summer rainfall events of more than 100 mm in the experimental area. A natural rainfall event of 24 mm occurred on day 6. At the completion of the 15-day gas collection period, the soil inside the PVC microplot was removed and separated into the 0–100 and 100–250 mm layers. The PVC microplot was then driven into the soil two times to collect soil samples of 250–500 and 500–750 mm increments below the enclosed area. All soil samples were oven dried at 45 °C and finely ground to a powder (< 0.1 mm) for chemical analysis.

Chemical analysis

Water-soluble C analysis in soil

A modified Walkley-Black method (Batonda & Waring 1984) was used to measure water-soluble C. Twenty grams of moist soil collected at the end of the field study were shaken in 80 ml distilled water for one hour. The extracts were filtered through No. 42 Whatman filter paper and the filtrates were refiltered through 0.2 μ m Sartorius cellulose filter paper to remove micro-organisms. A total of 15 ml of concentrated sulphuric acid were added to the mixture of 5 ml filtrate and 5 ml of

0.01 M K_2Cr_2O7 . The solution was then titrated using 0.03 M FeSO₄ with 2% o-phenanthroline solution as an indicator.

¹⁵N (
$$N_2O+N_2$$
) gas analysis

Gas samples collected were transferred to an arc vessel (Strong *et al.* 1991) and arced for a total of 250 s. The cooled arc vessel was connected to the inlet system of the mass spectrometer and the system evacuated to give a vacuum of less than 4 Pa. The arc vessel stopcock was then opened to allow the gas sample to expand through two cold traps immersed in liquid N. The gas sample was then introduced into the mass spectrometer for ¹⁵N abundance analysis. Recovery of ¹⁵N in the gas sample was calculated using an equation described by Buresh and Austin (1988).

Soil total N and ¹⁵N analysis

A variable amount of soil subsamples (depending on the soil total N content) was weighed into a tin capsule and introduced into a combined C/N analyser–mass spectrometer (Roborep CN (7001) / Tracermass System (9001) by an automatic sampler. ¹⁵N-labelled (NH₄)₂SO₄ solution, prepared in the laboratory, and a series of (¹⁵NH₄)₂SO₄ standard solutions of 0.38, 0.50, 1.00 and 2.00% ¹⁵N excess, supplied by the International Atomic Energy Agency, were used for calibration of the mass spectrometer.

Soil mineral N and 15N analysis

Soil mineral N was extracted with 2 M KCl in a ratio of 1:4 (soil:solution). Mineral N was determined on a subsample of the extract by a flow injection analyser, using a method recommended by the Lachat Instruments (QuickChem Method 10-107-06-4-D). For ¹⁵N analysis, a second subsample of the extract was acidified to pH 3 with 0.1 M H₂SO₄ and dried in an oven at 70–80 °C. The dried extract was analysed for ¹⁵N enrichment using a mass spectrometer.

Soil Br analysis

A method described by Saffigna *et al.* (1977) was used to measure Br in soil. Soil was extracted using 0.0025 M K_2SO_4 solution in a ratio of 1:4 (soil:solution), and extract filtered using Whatman 42 filter paper. About 0.6 ml total ionic adjustment buffer solution (5 M NaNO₃) were added to 30 ml of extract. Bromide was measured using a Br electrode with a reference electrode. The electrode was calibrated using a series of standard solutions between 0 and 10 mg Br l⁻¹.

Other soil chemical analysis

Soil pH, organic C and total K were determined as reported by Xu et al. (1993).

Statistical analyses

The differences between treatment means were tested for statistical significance using the least significant difference procedure in Microsoft Excel (Windows 95).

Results and discussion

Water-soluble C

No significant difference in water-soluble C was found between treatments in the soil at 250-500 and 500-750 mm depths (Figure 1). However, there was a significant difference (p < 0.05) in water-soluble C between treatments for the top 250 mm soil, especially at the 0–100 mm depth. The insignificant difference in levels of water-soluble C in the soil of 250-500 and 500-750 mm depths for all treatments suggested that while Br and nitrate could be leached down to deeper layers of the soil profile by rainfall, the water-soluble C released from the incorporated residues did not move below the top 250 mm soil. This may be because the decomposition and release of water-soluble C were slower processes, compared with the leaching of Br and nitrate in the soil profile (Figures 2 and 3), or maybe water-soluble C was oxidized at the soil surface before it could be leached to deeper soil layers.



Figure 1 Water-soluble C in the top 750 mm soil profiles under different residue management regimes 15 days after watering. Control: no residues incorporated; 1×FUG: unground foliage at 20 Mg DM ha⁻¹; 1×FG: ground foliage at 20 Mg DM ha⁻¹; 2×FG: ground foliage at 40 Mg DM ha⁻¹; 1×BUG: unground branches at 40 Mg DM ha⁻¹; 1×BG: ground branches at 40 Mg DM ha⁻¹; 2×BG: ground branches at 80 Mg DM ha⁻¹.



Figure 2 Recovery of applied bromide (Br) in the top 750 mm soil profiles under different residue management regimes 15 days after watering. Control: no residues incorporated; 1×FUG: unground foliage at 20 Mg DM ha⁻¹; 1×FG: ground foliage at 20 Mg DM ha⁻¹; 2×FG: ground foliage at 40 Mg DM ha⁻¹; 1×BUG: unground branches at 40 Mg DM ha⁻¹; 1×BG: ground branches at 40 Mg DM ha⁻¹; 2×BG: ground branches at 40 Mg DM ha⁻¹; all the residues were incorporated into the top 100 mm soil inside the microplots before ¹⁵N-labelled nitrate was added.

Therefore, when heavy rainfall occurred, both Br and nitrate could be leached down to sub-soil layers in the first instance, together with some of the water-soluble C already existing in the soil before the soil was watered. The newly released water-soluble C, which may be stimulated by watering (Pu *et al.* 1999), is more likely to remain in the surface soil and available for the processes of denitrification (if the surface soil is wet enough) and immobilization.

Water-soluble C in the soil in microplots with unground foliage materials at 20 Mg DM ha⁻¹ incorporated into the top 100 mm soil was much higher (43 mg C kg⁻¹) than that of unground branch materials (25 mg C kg⁻¹) at 40 Mg DM ha⁻¹ even though the quantity of incorporated branch materials was twice that of the incorporated foliage treatment (Figure 1). This could be attributed to decomposition of foliage materials being faster than that of branch materials (Bubb *et al.* 1998b). Grinding residues increased water-soluble C in soil in microplots treated with branch materials, but this was not the case for the microplots treated with foliage.

¹⁵N gas emissions

All microplots emitted ¹⁵N labelled gases (${}^{15}N_2 + {}^{15}N_2O$) from day 1 after the soil was watered and 0.2–4.5% of applied ¹⁵N was lost on the first day (Figure 4).



Figure 3 Distribution of mineral ¹⁵N (MN) and immobilized ¹⁵N (IN) in the top 750 mm soil profiles under different residue management regimes 15 days after watering. Control: no residues incorporated; 1×FUG: unground foliage at 20 Mg DM ha⁻¹; 1×FG: ground foliage at 20 Mg DM ha⁻¹; 2×FG: ground foliage at 40 Mg DM ha⁻¹; 1×BUG: unground branches at 40 Mg DM ha⁻¹; 1×BG: ground branches at 40 Mg DM ha⁻¹; 2×FG: ground branches at 40 Mg DM ha⁻¹; 2×FG: ground branches at 40 Mg DM ha⁻¹.

Microplots with ground foliage materials at 40 Mg DM ha⁻¹ experienced the highest ¹⁵N gas emission (4.5% of applied ¹⁵N) on day 1 while control microplots had the lowest ¹⁵N gas emission (< 0.3%). The ¹⁵N gas emission rates were then low for all treatments over the 15 days. This was different from the ¹⁵N gas emission patterns reported by Avalakki *et al.* (1995a, 1995b) and Pu *et al.* (1999) for cereal soils of poor drainage in subtropical Australia. The ¹⁵N gas emission rates in those experiments generally had a three-phase pattern: an initial phase of low ¹⁵N gas losses



Figure 4 Accumulated daily ¹⁵N gases (N₂+N₂O) emitted from microplots under different residue management regimes in the first 15 days after watering. Daily ¹⁵N-gas emission rate was calculated by multiplying a factor of 8 to the 3-hour value. Control: no residues incorporated; 1×FUG: unground foliage at 20 Mg DM ha⁻¹; 1×FG: ground foliage at 20 Mg DM ha⁻¹; 2×FG: ground foliage at 40 Mg DM ha⁻¹; 1×BUG: unground branches at 40 Mg DM ha⁻¹; 1×BUG: ground branches at 40 Mg DM ha⁻¹; 1×BG: ground branches at 40 Mg DM ha⁻¹; 1×BG: ground branches at 80 Mg DM ha⁻¹; all the residues were incorporated into the top 100 mm soil inside the microplots before ¹⁵N-labelled nitrate was added. The standard errors of total ¹⁵N gas emissions on days 1 and 15 are presented as vertical line bars.

and a declining phase. The low rate of ¹⁵N gas emissions found in this study after day 1 could be attributed to the lack of saturated conditions in the soil profiles due to the good drainage of the soil under the experimental conditions. However, numerous reports (Firestone 1982, Robertson & Tiedje 1988) have shown that anaerobic microsites are common under non-saturated conditions in the aggregates of fine textured, well structured soils and under these conditions nitrification proceeds in the aerobic zone and appreciable denitrification occurs at the microsites.

The low and quite steady level of ¹⁵N gas emission after day 1 from this experiment suggested that N loss via denitrification under field conditions occurred in soil microsites in which anaerobic conditions were maintained even over continued drying condition from day 6 to day 15. Microplots with ground foliage materials at 40 Mg DM ha⁻¹ showed the highest ¹⁵N gas emission rate between zero and day 1, compared with those receiving other residue treatments. However, subsequent rates were similar. This initial higher rate could be attributed to the fact that the large

quantity of foliage residues could immediately release available C for denitrifiers and also create anaerobic microsites in the soil by (1) accelerating microbial activity and depleting local oxygen supply as well as (2) retaining soil moisture. The presence of ample C substrate can cause rapid microbial O_2 consumption and therefore its depletion in soil microsites, thereby indirectly enhancing denitrification activity (Firestone 1982). Little difference in ¹⁵N gas emission was found between control microplots and microplots treated with unground branch materials. This indicated that hoop pine branch materials might contain relatively low level of available C. The little available C would be rapidly used in moist soil by denitrifiers. When the residues were ground, more available C could be released quickly and available to denitrifiers.

N transformations

Recoveries of the applied ¹⁵N and Br ranged from 33 to 59% and from 47 to 69% respectively in the top 750 mm of soil (Table 1). The difference between the recovered Br and ¹⁵N or denitrification loss ranged from 6 to 26%. The lower recovery of ¹⁵N compared with Br was assumed as the ¹⁵N loss via denitrification. However, this might be an underestimation as some of the ¹⁵N might have already been denitrified before moving to deeper soil layers. The immobilized ¹⁵N ranged from 16 to 34%, leaving 13 to 24% of the applied ¹⁵N-NO₃ remaining in the top 750 mm of soil (Figure 3). The recoveries of both applied ¹⁵N and Br from control microplots were significantly (p < 0.05) lower than those of the other treatments. It appears that the incorporation of residues into the top 100 mm of soil inside the respective microplots played an important role in reducing leaching and increasing the recoveries of the applied Br and ¹⁵N in the soil profiles.

Surprisingly, denitrification loss was lowest (6%) for microplots with ground branch materials incorporated at 80 Mg DM ha⁻¹ (Table 1). This result contradicted

Residue management regime	¹⁵ N recovery	Br recovery	Denitrification loss
Control without any residue added	33.4 (10.0)	46.9 (5.1)	13.6 (5.8)
Unground foliage materials at 20 Mg DM ha ⁻¹	48.6 (4.1)	69.4 (4.4)	20.8 (4.4)
Ground foliage materials at 20 Mg DM ha ⁻¹	38.7 (2.2)	65.0 (7.7)	26.2 (6.8)
Ground foliage materials at 40 Mg DM ha ⁻¹	48.6 (3.1)	69.3 (7.1)	20.7 (10.3)
Unground branch materials at 40 Mg DM ha ⁻¹	48.2 (13.7)	63.6 (12.3)	15.4 (8.0)
Ground branch materials at 40 Mg DM ha ⁻¹	41.7 (4.8)	54.2 (12.6)	12.5 (10.0)
Ground branch materials at 80 Mg DM ha ⁻¹	58.5 (3.5)	64.2 (4.8)	5.8 (6.9)
LSD (p < 0.05)	11.2	12.0	10.1

Table 1Recoveries of added ¹⁵N and Br in the top 750 mm soil profiles 15 days after watering the soils and
estimated ¹⁵N losses via denitrification regimes in a 1-year-old hoop pine plantation

the finding that these microplots had the second highest denitrification loss, measured by ¹⁵N gases evolved (Figure 4). Denitrification losses for all microplots treated with foliage materials were significantly higher than those of the control, suggesting that the added residues could promote denitrification loss. These results were consistent with many other studies (Xu *et al.* 1992, Avalakki *et al.* 1995a, Pu *et al.* 1999, 2001).

Incorporating branch materials into the soil did not show any increase in denitrification loss although there was generally a significant increase in watersoluble C (Figure 1). This suggests that while the foliage materials with lower C:N ratio (41:1) quickly released available C for soil denitrifiers, branch materials with high C:N ratio (111:1) were slower in releasing available C and might contribute little to the early stage of denitrification activity. This agrees with the findings that narrow C:N ratio encourages denitrification (Aulakh *et al.* 1991, 2000, Laverman *et al.* 2001). Grounded materials created much more surface areas in contact with soil and hence released more soluble extractable compound as indicated by the water-soluble carbon levels of the different regimes and hence increased denitrification activity.

About 31 to 53% of the applied Br could not be recovered from the 750 mm soil profiles. More than half of the recovered Br was in the 250–500 mm soil depth and 5–10% of the applied Br recovered in the 500–750 mm soil depth (Figure 2). Distribution of Br in the soil profiles suggests that the unrecovered Br could have been lost due to either leaching of Br below the 750 mm soil profiles or the lateral movement of Br outside the microplot area below the 250 mm soil profiles, which were not sampled. It was estimated that about two thirds of the uncovered Br was due to lateral movement using data of the recovered Br from another field experiment (Pu *et al.* 2001). Xu *et al.* (1992, 1993) reported that significant amounts of fertilizer ¹⁵N could move outside the microplots of 250 mm soil in two to three months after ¹⁵N application at 40 kg N ha⁻¹ in a semi-arid environment of tropical Australia. Nevertheless, their studies have indicated that the downward movement of fertilizer ¹⁵N in the soil profiles is much greater than the lateral movement below the 250 mm soil.

While useful information could be obtained using Br as a tracer to investigate the leaching and denitrification loss of applied ¹⁵N, it should be noted that denitrification loss calculated by the difference between the recovered Br and ¹⁵N-labelled nitrate could be underestimated. The leaching loss, determined by the difference between the applied and recovered Br, could be an overestimation. The possible underestimation of denitrification loss and overestimation of leaching loss may be attributed to the differences in the transformation behaviour between the applied ¹⁵N-labelled nitrate and Br in the soil. Significant amounts of the applied ¹⁵N-labelled nitrate could be immobilized and this part of applied ¹⁵N would not be subjected to further leaching loss or denitrification. This is not the case for applied Br, which could be leached below the soil surface whenever there is vertical downward water movement in the soil profiles. Furthermore, the lateral movement of ¹⁵N out of the microplots would underestimate denitrification and overestimate leaching loss. Therefore, we suggest that a circular plot of 1000 mm diameter should

be established surrounding the 235 mm diameter microplot and the same rates of Br and ¹⁵N-unlabelled (¹⁵N natural abundance with about 99.6% ¹⁴N) nitrate could be applied to the surrounding area as those applied to the area inside the microplot when a field microplot experiment would need to be conducted under the similar experimental conditions. This suggests improvement in the microplot installation has been confirmed in a denitrification study of Pu *et al.* (2001).

¹⁵N recoveries as mineral N ranged from 13 to 24% and ¹⁵N immobilized varied from 16 to 34 % (Figure 3). Total recovered mineral ¹⁵N in the 0–100, 100–250 and 500–750 mm soil depths was less than in the 250–500 mm depth for all treatments, indicating that there was lateral movement of applied 15N. There was no significant difference in mineral ¹⁵N recovery between treatments, except for the ground branch materials applied at 80 Mg DM ha⁻¹. Low recovery of the applied ¹⁵N (Figure 3) and high level of water-soluble C (Figure 1) found in the top 250 mm soil suggested that the surface 250 mm soil was an active zone for denitrification and immobilization.

There was much more immobilized ¹⁵N in the 0–100 and 100–250 mm soil depths. This suggested that while denitrification rate was low from day 2 as indicated by directly measuring ¹⁵N gases, the process of immobilization was probably able to continue during the period of this study since immobilization would not require anaerobic conditions in the soil. The study by Pu *et al.* (1999) showed that ¹⁵N immobilization is a slower process than denitrification after water logging poorly drained soils. However, in fully aerobic soils, ¹⁵N immobilization is a stronger competitor for the available soil nitrate than denitrification. All microplots treated with the hoop pine residues showed a higher rate of immobilized ¹⁵N, compared with those of the control. This indicated that available C released from the residues (incorporated into the soil) after watering also promoted ¹⁵N immobilization. The denitrification ¹⁵N losses measured by the gas emission method were generally lower than those by the mass balance method, similar to the findings by Pu *et al.* (2001).

Conclusions

This study showed that after simulated heavy rainfall, substantial soil mineral N may be lost from the ecosystem through leaching (31-53%) and denitrification (6-26%). Incorporating foliage materials in the top 100 mm soil generally increased denitrification loss and promoted ¹⁵N immobilization in the surface soil. Incorporating branch materials did not show any significant increase in the ¹⁵N loss via denitrification, although ¹⁵N immobilization was greater than that in control microplots. Water-soluble C released through the residue decomposition, which could be stimulated by watering, would be most likely to remain in the surface soil due to the slow residue decomposition process compared with the leaching process of Br and ¹⁵N. The ¹⁵N gas emissions indicated that for the well-drained soil, high rates of denitrification (~4% of applied ¹⁵N/day) could only occur over a short period, but low rates of denitrification (~0.1% of applied ¹⁵N/day) could continue for a longer period.

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