Soil biogenic emissions of nitric oxide from a semi-arid savanna in South Africa

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Abstract

Soils of arid and semi-arid ecosystems are important biogenic sources of atmospheric nitric oxide (NO), however, there is still a shortage of measurements from these systems. Here we present the results of a laboratory study of the biogenic emission of NO from four different landscape positions of the Kruger National Park (KNP), a large conservation area in a semi-arid region of South Africa. Results show that the highest net potential NO fluxes come from the low lying (footslope) landscape regions, which have the largest nitrogen stocks and highest rates of nitrogen input into the soil. Net potential NO fluxes from midslope and crest regions were considerably lower. The maximum release of NO occurred at fairly low soil moisture contents of 10%–20% water filled pore space. Using soil moisture and temperature data obtained in situ at the Kruger National Park flux tower site, net potential NO fluxes obtained in the laboratory were converted to field fluxes for each of the four landscape positions for the period 2003 to 2005. The highest field NO flux is from footslope positions, during each of these years and emissions ranged from 1.5–8.5 kg ha\(^{-1}\) yr\(^{-1}\) (in terms of mass of nitrogen). Remote sensing and Geographic Information Systems techniques were used to upscale field NO fluxes on a regional basis indicating that the highest emissions occurred from the midslope positions, due to their large geographical extent in the considered research area. Emissions for the KNP Skukuza land type (56 000 ha) ranged from 20\(\times\)10\(^3\) kg in 2004 to 34\(\times\)10\(^3\) kg in 2003. The importance of landscape characteristics in the determination of regional biogenic NO soil emissions is emphasized.

1 Introduction

Emissions of nitric oxide (NO) and its conversion to nitrogen dioxide (NO\(_2\)) (collectively referred to as NO\(_x\)=NO+NO\(_2\)) are important in regulating chemical processes in the atmosphere (Crutzen, 1995; Crutzen and Lelieveld, 2001; Levine et al., 1997). Nitrogen oxides are key catalysts in the chemical processes that generate or destroy
ozone. The ambient NO$_x$ mixing ratio is a threshold which determines whether ozone is generated or destroyed (Chameides et al., 1992; Crutzen and Lelieveld, 2001; Meixner and Yang, 2006). The ozone forming or destroying reactions of nitric oxide are also involved in chemical processes that result in the production or consumption of the hydroxyl radical (OH), which is the chemical species most responsible for cleaning the atmosphere (Monks, 2005). Nitrogen oxides are removed from the atmosphere in a series of photochemical reactions that produce nitric acid (Monks, 2005). Nitric acid is an important component of acid deposition and acts as a source of N deposited from the atmosphere, in both the wet and dry form (Feig et al., 2007; Logan, 1983; Remde et al., 1993).

In 1860 it was estimated that the global NO$_x$ emission was 13.1 Tg a$^{-1}$ (all values mentioned are in terms of mass of nitrogen), of which 10.5 Tg a$^{-1}$ came from natural sources. By the early 1990s it was estimated that the emissions of NO$_x$ gases totalled 45.9 Tg a$^{-1}$ (Galloway et al., 2004). The recent estimate of the Intergovernmental Panel on Climate Change is 42–47 Tg a$^{-1}$ (Denman et al., 2007). The most important sources of NO$_x$ is fossil fuel combustion in power stations and vehicles (45%–67% of the total) (Denman et al., 2007; Kasibhatla et al., 1993) followed by lightning (5%–16%) and biomass burning (13%–29%) with the biogenic emission from soil (either natural or under agriculture) accounting for between 10% and 40% of the total (Davidson and Kingerlee, 1997; Denman et al., 2007). Nitrification has long been considered the main source for NO from the soil (Conrad, 1996; Coyne, 1999; Davidson et al., 1993; Garrido et al., 2002; Remde et al., 1989; Russow et al., 2000; Russow et al., 2008; Stehfest and Bouwman, 2006; Tabachow et al., 2001). The proportion of nitrified N emitted as NO has been shown to range between 0 and 2.5% of the total N involved in nitrification (Garrido et al., 2002). While it is known that soil processes are responsible for a sizable proportion of the global NO$_x$ budget, there is considerable uncertainty about exactly how much. Davidson and Kingerlee (1997) provided a global inventory of NO emissions from soils, based on field measurements world-wide. Their estimate of the global NO soil source strength is 21 Tg a$^{-1}$ (with an error margin of 4 to 10 Tg a$^{-1}$),
while the 4th IPCC estimate is 8.9 Tg a⁻¹ (Denman et al., 2007) up from the 3rd IPCC estimate of 5.6 Tg a⁻¹ (IPCC, 2001). The emission of NO from soil is affected by both biotic and abiotic factors; in addition, some ecosystems are better studied than others and therefore the quality of estimates differ between regions. While temperate regions and agricultural systems have been fairly well studied, a recent review by Meixner and Yang (2006) identified only 13 studies in natural drylands (annual precipitation below 400 mm). This is unfortunate since arid and semi-arid regions make up 40% of the earth’s surface area (Veron et al., 2006) and are known to be an important potential source of biogenic NO emissions (Davidson et al., 1993).

The microbial activity controlling biogenic NO emissions and consumption from the soil is controlled by a number of factors including soil moisture, soil temperature and soil nutrient status (Conrad, 1994; Ludwig et al., 2001; Meixner et al., 1997). These in turn are largely controlled by geographic factors, for example the climatic conditions, the position in the landscape, the land use (Davidson, 1991b) and biological factors such as the vegetation.

In drier ecosystems, soil water seems to be the most important factor regulating emissions of NO. When the soil moisture is too low to maintain microbial activity there are very low levels of NO emitted (Garrido et al., 2002; Meixner et al., 1997) and when soil moisture level are too high to maintain aerobic conditions, the emission of NO is negligible (Skopp et al., 1990). The optimal emission of NO seems to occur at low soil moisture levels, but where microbial activity can still take place. In a previous field study in the Kruger National Park (KNP), NO emissions followed the predicted soil moisture curve where they were lowest at high (>54%) and at low (<8.7%) values of Water Filled Pore Space (WFPS) (Parsons et al., 1996). Even when desert soils are wetted, emissions of N containing gasses can occur with the maximum emission of NO occurring at fairly low soil moisture contents (10% WFPS) (Hartley and Schlesinger, 2000).

A secondary controller of biogenic NO emissions is the soil nutrient status; many studies have found a relationship between the emissions of NO and either the concen-
trations of ammonia or nitrate (Erickson et al., 2002, 2001; Hartley and Schlesinger, 2000; Hutchinson et al., 1993; Ludwig et al., 2001; Meixner et al., 1997; Parsons et al., 1996) or the N cycling rate (Erickson et al., 2002; Erickson et al., 2001; Hartley and Schlesinger, 2000; Parsons et al., 1996). Therefore, natural or anthropogenic actions that result in the modification of the inputs of nutrients or the rates of nutrient turnover are likely to have an effect on the NO production rates.

Catenal development is an important landscape process in many of the savanna regions of southern Africa, where the distribution of soil physical and chemical properties and vegetation community occurrence is determined by topographic position. Landscape processes, such as catenal development provide a natural example of a process that can control the distribution of moisture and nutrients across a landscape. Therefore the aim of this study was to determine the emission of NO across differing landscape positions in a semi-arid savanna ecosystem in southern Africa.

2 Methods and materials

2.1 Site description

This research occurred at the Kruger National Park (KNP) Flux Tower site (25°01.184’S; 31°29.813’E; 365 m above sea level) which is situated in a semi-arid savanna in the north eastern part of South Africa (Fig. 1a), 13 km WSW of the tourist camp and administrative centre of Skukuza (for further details see Scholes et al. (2001)). It is a well characterised site that has been used in various other studies. Exchanges of CO₂, energy and H₂O have been measured continuously since April 2000 using the eddy covariance technique (Harley et al., 2003; Scholes et al., 2001). Meteorological parameters have been measured at the main tower, while soil moisture and temperature has been measured at various depths in the two main surrounding ecosystem types since 2000. The flux tower is positioned exactly on the ecotone between a ridge-top broad-leaved Combretum savanna on sandy soil, dom-
inated by *Combretum apiculatum* and *Sclerocarya birrea* and a midslope fine leaved *Acacia* savanna on clayey soil, dominated by *Acacia nilotica*, *A. nigrescence* and *S. birrea* (Scholes et al., 2001) (as shown in Fig. 1b). This characteristic catenal pattern is repeated throughout the wider region (Venter et al., 2003). The fine-leaved savannas have higher concentrations of soil nitrogen than the broad-leaved savannas (Table 1) (Scholes et al., 2003; Woghiren, 2002). The climate at the site is semi-arid subtropical with rainfall averaging 550 mm annually. Summers are hot and wet while winters are warm and dry, for more detailed climatic information see Scholes et al. (2001) and Venter et al. (2003).

The soil physical and chemical properties are presented in Table 1 and include values reported by Woghiren (2002) and measurements conducted during this study. The soils are sandy loams in the Crest and midslope positions and sandy clay loams in the Footslope positions as determined through a sedimentation technique (Day, 1969). The soil pH was determined using the method of Anderson and Ingram (1993) and ranges from 6.1 to 6.5. The total C and N was measured at the Microanalytical laboratory at the University of Mainz using a Vario MICRO Cube universal microanalyser set up to measure total C, H, N and S content of our soil samples.

### 2.2 Sampling and instrumental set up

Soils were sampled in April 2005, using stainless steel soil cores with a length of 5 cm and radius of 2.5 cm, from four landscape positions (see Fig. 1b); the Crest, 20 m upslope of the Ecotone referred to as “Upslope”, 20 m downslope of the Ecotone, referred to as “Downslope” and the Footslope. Soil samples were comprised of 10 sub-samples taken from the top 5 cm in each of the landscape positions, the ten samples were taken in a band parallel to the Ecotone at approximately 1 m intervals. The bulk samples were sieved through 2 mm mesh, air dried and then stored at 5°C until analysis. When soil is rewetted after a long period of inactivity there is often a large and rapid release of NO from the soil, either as the result of displacement of NO rich air from the soil or from the rapid mineralisation of readily available nitrogen sources. This pulsing effect
(a) is of short duration and is not repeated or the magnitude is greatly reduced on sub-
sequent rewetting (Kirkman et al., 2001; Otter et al., 1999; Scholes et al., 1997), and
(b) may add to the total NO flux, but it is thought to be fairly insignificant, less than
6% of the annual NO flux (Scholes et al., 1997). Therefore, two days before begin-
ning the laboratory analysis, an aliquot from the soil samples was soaked with deionised
water and allowed to drain freely at room temperature (22°C) to avoid the effect of puls-
ing after the initial wetting of soil after a long period of inactivity (Scholes et al., 1997;
Snyder and Tartowski, 2006). The basic methodology for the laboratory measurement
of NO flux from soil has been previously described (Aranibar et al., 2004; Kirkman et
al., 2001; Meixner and Yang, 2006; Otter et al., 1999; Van Dijk et al., 2002) and is
only briefly described here. A known mass of sieved, wetted soil (approximately 100 g
dry weight) was placed in one of five Plexiglas chambers (volume 9.7 × 10^{-4} m^3) in a
thermo-controlled cabinet (four chambers were used for soil samples while one was
kept empty as a reference). Pressurised air that had passed through a purification
system (four traps consisting of glass wool, activated charcoal, silica gel, and molec-
ular sieve to provide a dry and “NO free” air stream) was supplied to each chamber
at a flow rate of 4.2 × 10^{-5} m^3 s^{-1} (2.5 L min^{-1}), controlled by five mass flow controllers.
The air in each chamber was well mixed using a Teflon coated fan (Micronel®, USA)
in each chamber, flow rate 8.7 × 10^{-4} m^3 s^{-1} (52 L min^{-1}). The outlet of each cham-
ber was connected via (a) a reverse nafion tube (Perma Pure® MD-125-127), and
(b) a switching valve to a NO Chemiluminescence analyser (Eco Physics Switzerland,
model CLD780TR) and a H_2O/CO_2 analyser (Rosemont Analytical, USA, Binos IR gas
analyser). Occasionally, nitric oxide standard gas (200 ppm) was diluted into the air
purification system via a mass flow controller (Mass-Flo® 200 sccm Range, MKS in-
struments, USA); this allowed (a) calibration of the NO Chemiluminescence analyser
and (b) control of the chamber headspace NO concentration when determining NO up-
take in the soil (see below). The soil moisture content was determined by tracking the
loss of water vapour throughout the measurement period and relating it to the gravimet-
ric soil moisture content at the start and end of the measurement period. The purpose
of the reverse Nafion driers is (a) to keep the humidity of the chambers’ headspace air
high, and hence (b) to slow the dehydration of the soil, allowing the microbes in the soil
time to equilibrate to changes in the soil moisture content.

2.3 Measurements of the net, NO release rate

The net release of NO ($J_{NO}$, in ng kg$^{-1}$ s$^{-1}$) is calculated from the difference between
the NO mixing ratio at the outlet of the reference cuvette and the outlet of each of the
incubation cuvettes (since the air in the cuvette is well mixed, the air at the outlet is
assumed to have the same composition as the air in the headspace) according to:

$$J_{NO} = \frac{Q}{M_{soil}} (m_{NO, out} - m_{NO, ref}) \times \left(\frac{M_N}{V_m} \times 10^{-3}\right)$$

Where $Q$ is the flow rate through the cuvette ($4.2 \times 10^{-5}$ m$^3$ s$^{-1}$ or 2.5 L min$^{-1}$), $M_{soil}$ is
the dry mass of the soil (kg), $M_N$ is the molar mass of N (14.0076 kg kmol$^{-1}$), $V_m$ is
the molar volume (24.465 m$^3$ kmol$^{-1}$ at 25°C 1013.25 hPa) and $m_{NO, ref}$ and $m_{NO, out}$ are
the mixing ratios of NO in ppb, at the outlets of the control and incubation cuvettes
respectively (the factor $M_N/V_m \times 10^{-3}$ is needed to convert NO mixing ratio (ppb) to NO
concentration (ng m$^{-3}$)).

The release of NO from the soil is the result of the microbial production and con-
sumption of NO in the soil, processes that occur simultaneously (Conrad, 1994, 1996;
Conrad and Smith, 1995). As a result, the NO release rate ($J_{NO}$) is always a net release
rate. However if the NO consumption is greater than production in the soil sample then
$J_{NO}$ becomes negative. This will only occur if the in-coming (= reference) NO mixing
ratio is greater than the headspace NO mixing ratio in the soil containing chamber.

It has already been shown experimentally that there is a linear relationship between
the NO release rate ($J_{NO}$) and the rate of NO production ($P$) and consumption ($k$)
(Ludwig et al., 2001; Remde et al., 1989) so that the measured release rates can be
described according to:

\[ J_{NO} = P - k \times m_{NO,\text{out}} \times \left( \frac{M_N}{V_m} \times 10^{-3} \right) \]  \hspace{1cm} (2)

This equation implies that the NO production is independent of the NO mixing ratio in the headspace \((m_{NO,\text{out}})\), while the NO consumption is dependent on the NO mixing ratio in the headspace, and can be approached as a first order decay process. To determine the values of \(P\) and \(k\), Eq. (2) was used with the measured release rates \((J_{NO})\) from two sets of incubation measurements, \(m_{NO,\text{ref}}=0\) ppb and \(m_{NO,\text{ref}}=58\) ppb. This allowed us to calculate \(P\) (ng kg\(^{-1}\) s\(^{-1}\)) and \(k\) (m\(^3\) kg\(^{-2}\) s\(^{-2}\)), where \(k\) can be determined from the slope of Eq. (2),

\[ k = \frac{\Delta J_{NO}}{\Delta [\text{NO}]} = \frac{J_{m,\text{high}} - J_{m,\text{low}}}{m_{\text{high}} - m_{\text{low}}} \times \left( \frac{V_m}{M_N} \times 10^{3} \right) \]  \hspace{1cm} (3)

Where \(m_{\text{low}}\) is the actual NO mixing ratio (ppb) in the head space of the cuvette under fumigation with NO free air and \(m_{\text{high}}\) is the actual NO mixing ratio in the cuvette headspace under fumigation with 58 ppb NO. In this study, we will present values of the NO release rate \((J_{NO})\) as a function of soil moisture, in terms of the soil WFPS.

Water filled pore space is a useful concept because it indicates the amount of water in the soil that is available for microbial activity and also the amount of air in the soil and therefore the soil oxygen status. The WFPS is calculated (a) from the amount of water lost from the enclosed cuvettes through evaporation during the incubation process and (b) through determining the gravimetric water content \((\Theta)\) of the rewetted sample at the start of the incubation. The WFPS is calculated according to Eq. (4):

\[ \text{WFPS} = \Theta \times \frac{BD}{1 - \frac{BD}{PD}} \]  \hspace{1cm} (4)

Where BD is the soil bulk density in (kg m\(^{-3}\)) measured at the site of sampling, by driving a stainless steel core of known volume driven into the soil and removing a soil
sample and drying the soil at 105°C for 48 h, and PD is the particle density of the average soil mineral (quartz) with a value of $2.65 \times 10^3$ kg m$^{-3}$ according to (Parton et al., 2001).

2.4 Detection limit and estimation of the release rate error

The detection limit of our laboratory technique was determined in a study by Gelfand et al. (2008)$^1$, but is briefly described here. Inert glass beads and autoclaved Israeli desert soils were used to measure the “blank” net release of NO. It was found that the “blank” net release of NO from the glass beads was at a rate of 0.02 ng kg$^{-1}$ s$^{-1}$, with a random deviation of 0.02 ng kg$^{-1}$ s$^{-1}$ irrespective of the moisture content, therefore, we consider an experimentally derived detection limit for $J_{NO}$ of 0.08 ng kg$^{-1}$ s$^{-1}$ this results from the mean release rate of autoclaved soils plus 3 standard deviations (corresponding to a confidence interval of 99.7%). Similarly, the detection limit of the autoclaved soils is 0.11 ng kg$^{-1}$ s$^{-1}$, therefore the more conservative estimate from the autoclaved soils is being used as our detection limit.

To quantify the precision of $J_{NO}$ measurements, the NO net release rate was determined experimentally through the simultaneous measurement of four replicates across the full range of soil moisture. The mean standard deviation on the NO net release rate was found to be 0.03 ng kg$^{-1}$ s$^{-1}$ irrespective of WFPS, this is lower than the experimentally derived detection limit of $J_{NO}$ and we therefore consider $\pm 0.05$ ng kg$^{-1}$ s$^{-1}$ as a conservative estimate of the experimentally derived precision of $J_{NO}$.

2.5 Calculation of the net potential NO flux

The laboratory derived net release of NO ($J_{NO}$, in ng kg$^{-1}$ s$^{-1}$) from the soil was converted to a net potential NO flux ($F_{lab}$, in ng m$^{-2}$ s$^{-1}$) using a simple diffusion based algorithm (Eq. 5), originally developed by Galbally and Johansson (1989), modified by

van Dijk et al. (2002). The net potential laboratory NO flux, as a function of WFPS and $T_{\text{soil}}$, is calculated according to:

$$F_{\text{lab}}(T_{\text{soil}}, \text{WFPS}) = \sqrt{BD \times k(T_{\text{soil}}, \text{WFPS}) \times D(\text{WFPS})} \times \left( \frac{P(T_{\text{soil}}, \text{WFPS})}{k(T_{\text{soil}}, \text{WFPS})} - [\text{NO}]_{\text{Headspace}} \right)$$  \hspace{1cm} (5)

where $D(\text{WFPS})$, in m$^2$ s$^{-1}$, is the WFPS dependent diffusion coefficient of NO through the soil which is calculated after Moldrup et al. (2000), from WFPS and the gas diffusion constant for free air (m$^2$ s$^{-1}$) equal to $1.9 \times 10^{-5}$ m$^{-2}$ s$^{-1}$ (Gut et al., 1998), and where $[\text{NO}]_{\text{Headspace}}$ is the NO concentration (in $10^{-12}$ kg m$^{-3}$) in the headspace of the cuvette. The diffusion coefficient is dependent of the soil moisture content and the soil bulk density and therefore is calculated for each soil sample and each soil moisture interval.

For a given soil temperature, an algorithm has been developed (Meixner and Yang, 2006) to fit our net potential NO fluxes as a function of the WFPS (Eq. 6). This algorithm describes the net potential NO flux as a power increase until optimal soil moisture followed by an exponential decrease:

$$F_{\text{lab}}(T_{\text{soil}} = \text{const.}, \text{WFPS}) = a \text{WFPS}^b \exp(-c \text{WFPS})$$  \hspace{1cm} (6)

Where parameters $a$, $b$ and $c$ are related to observed values by:

$$a = \frac{F_{\text{lab}}(\text{WFPS}_{\text{opt}})}{\text{WFPS}_{\text{opt}}^b \exp(-b)}$$ \hspace{1cm} (7)

$$b = \frac{\ln \left[ \frac{F_{\text{lab}}(\text{WFPS}_{\text{opt}})}{F_{\text{lab}}(\text{WFPS}_{\text{upp}})} \right]}{\ln \left( \frac{\text{WFPS}_{\text{opt}}}{\text{WFPS}_{\text{upp}}} \right) + \frac{\text{WFPS}_{\text{upp}}}{\text{WFPS}_{\text{opt}}} - 1}$$ \hspace{1cm} (8)

$$c = \frac{-b}{\text{WFPS}_{\text{opt}}}$$ \hspace{1cm} (9)
Where, for a given $T_{\text{soil}}$, $WFPS_{\text{opt}}$ is the soil moisture where the maximum laboratory NO flux is observed, $F_{\text{lab}}(WFPS_{\text{opt}})$ is the maximum laboratory NO flux at the optimal soil moisture content, and $WFPS_{\text{upp}}$ is the soil moisture content where $F_{\text{lab}}$ approximately equals zero ($F_{\text{lab}}(WFPS_{\text{upp}}) = 1/100 F_{\text{lab}}(WFPS_{\text{opt}})$) for $WFPS > WFPS_{\text{opt}}$.

The temperature dependence of the laboratory NO flux was determined by measuring the net potential laboratory NO flux at two soil temperatures, 25°C and at 35°C. The temperature dependence usually shows an exponential increase and can be expressed as the increase of $F_{\text{lab}}$ for a 10°C increase in soil temperature, otherwise known as a $Q_{10}$ function (Eq. 10) (Lloyd and Taylor, 1994).

$$Q_{10}(WFPS) = \frac{F_{\text{lab}}(T_{\text{soil}}=35^\circ \text{C}, WFPS)}{F_{\text{lab}}(T_{\text{soil}}=25^\circ \text{C}, WFPS)}$$  \hspace{1cm} (10)

The $Q_{10}$ function can then be included into Eq. (6), as a “temperature amplification factor” of the reference NO flux ($T_{\text{ref}}=25^\circ \text{C}$), so that the laboratory NO flux can be estimated as a function of both soil temperature and soil moisture (Eq. 11):

$$F_{\text{lab}}(T_{\text{soil}}, WFPS) = a_{T_{\text{ref}}}$WFPS$b_{T_{\text{ref}}} \exp \left(c_{T_{\text{ref}}} \times WFPS \right) \times \exp \left[\frac{\ln Q_{10}(WFPS)}{10} \times (T_{\text{soil}} - T_{\text{ref}})\right]$$ \hspace{1cm} (11)

2.6 Compensation point mixing ratio

The compensation point mixing ratio ($m_{\text{NO,comp}}$) is an important concept for the bi-directional exchange of NO (see Conrad, 1994). Since it determines what the ambient mixing ratio of NO in the atmosphere has to be before a net NO uptake into the soil can occur. The compensation point mixing ratio is calculated by resolving Eq. (5) for the NO concentration where $F_{\text{lab}}(WFPS, T_{\text{soil}})=0$:

$$m_{\text{NO,comp}}(T_{\text{soil}}, WFPS) = \frac{P(T_{\text{soil}}, WFPS)}{k(T_{\text{soil}}, WFPS)} \times \left(\frac{V_m}{M_N} \times 10^3\right)$$ \hspace{1cm} (12)
2.7 Up-scaling to the landscape positions

Once we have derived the net potential NO flux ($F_{\text{lab}}$) from our laboratory measurements as a function of both, soil moisture and temperature, the flux of NO from the four landscape positions ($F_{\text{NO, up-scaled}}$) at the KNP Tower site could be estimated by suitable up-scaling procedures using field data of soil temperature and moisture. For that we used the half hourly soil moisture and temperature data obtained from measurement stations within the fine-leaved savanna and the broad-leaved savanna, corresponding with the Crest and Footslope landscape positions and the soil BD and D values measured at each of the landscape positions. The $F_{\text{NO, up-scaled}}$ from the two midslope positions was approached by the soil moisture and temperature data from the closest measurement site (the Crest for the Upslope position and the Footslope for the Downslope position). These soil moisture and temperature data were used in Eq. (11) to calculate the up-scaled flux on a half hourly basis. These half-hourly flux estimates were used to calculate monthly averages of $F_{\text{NO, up-scaled}}$.

2.8 Up-scaling to the landscape scale

The Kruger National Park (KNP) Geographical Information System (GIS) data base divides the Skukuza land system (56 000 ha) into 3 landscape positions; the Crest comprising 12% of the surface area (6720 ha), the midslope comprising 80% of the surface area (44 800 ha), and the Footslope, comprising 8% of the surface area (4480 ha). Since the midslope landscape position encompasses both, the Downslope and Upslope sites of our sampling scheme, the midslope area was divided equally between the Downslope and Upslope sites, giving each 40% of the land type surface area (22 400 ha). The monthly average NO fluxes for each of our designated landscape positions (Sect. 2.7) were apportioned according to the proportion that each of the landscape positions comprised of the total Skukuza land type (Venter et al., 2003) to estimate a total emission for the Skukuza land type for the years 2003–2005. This technique assumes that the soil BD, the soil temperature and soil moisture are the same.
across the whole region, for this reason the up-scaling attempt that we have made is limited to the Skukuza land type which is where the flux tower is situated, and where we can assume with reasonable confidence that on the whole similar conditions occur.

To assess the potential distribution of NO emissions from the Skukuza land type the landscape positions were associated with differing slope classes (Venter et al., 2003) as shown in Table 1 (bottom line). Shuttle Radar Tomography Mission (SRTM) digital elevation data (supplied by the Consortium for Spatial Information, of the Consultative Group for International Agricultural Research (CGIAR-CSI)) for the classification of the slope classes was used and incorporated into a geographical information system (GIS) database to give the distribution of the landscape classes for the Skukuza land type (Fig. 6). The annual point NO fluxes associated with the individual landscape positions were then assigned to the corresponding slope classes to provide a distribution of the annual NO emission from the various landscape positions of the Skukuza land type.

3 Results

3.1 General behaviour of the net potential NO fluxes ($F_{lab}$)

For the reference soil temperature of 25°C, an overview of the maximum derived net potential fluxes ($F_{lab}$) and the $Q_{10}$ values are given in Table 2. At 25°C the maximum $F_{lab}$ ranged from 1.3 ng m$^{-2}$ s$^{-1}$ in the Downslope soil to 2.4 ng m$^{-2}$ s$^{-1}$ in the Footslope. At 35°C the laboratory NO flux ranged from 3.1 ng m$^{-2}$ s$^{-1}$ in the Downslope soil to 3.5 ng m$^{-2}$ s$^{-1}$ in the Upslope soil. With the full temperature dependence included, the net potential NO fluxes were calculated for the four landscape positions as a function of soil moisture and temperature and are shown in Figure 2 a-d. Differences in the graphs reflect the differing emission potential as a function of soil WFPS and soil temperature for the individual soils. The highest $F_{lab}$ occurs in the Footslope, where at 40°C the $F_{lab}$ is 3.2 ng m$^{-2}$ s$^{-1}$, while the lowest $F_{lab}$ (2.5 ng m$^{-2}$ s$^{-1}$) occurred in the Downslope at 40°C.
The maximum $F_{\text{lab}}$ for all four landscape positions was calculated to lie between 10% and 20% WFPS, (Table 2). The optimum WFPS differed according to the temperature of incubation. At 25°C, it ranged from 11% for the Upslope position to 17% in the Footslope. The optimum WFPS was always greater under the higher incubation temperatures and occurred at 18% WFPS (Footslope, Downslope, Upslope) at 35°C. Maximum $F_{\text{lab}}$ differed according to the incubation temperature and increased in the Footslope, and the two midslope positions. The measurement of $F_{\text{lab}}$ at 35°C from the Crest soil was unsuccessful, since the aliquot of the corresponding soil sample got wet during the storage period between the measurements at 25°C and 35°C. Therefore, the $Q_{10}$ value for the Upslope site was used for the Crest site to estimate the temperature dependence during subsequent up-scaling calculations.

The maximum NO consumption constant ($k$), calculated from Eq. (3) as a function of $T_{\text{soil}}$ and WFPS, occurs at approximately the same WFPS as the maximum of $F_{\text{lab}}$ (see Fig. 3), namely between 10%–25% WFPS. The highest NO consumption occurs in the Footslope soils ($2.7 \times 10^{-4}$ m$^3$ kg$^{-1}$ s$^{-1}$) while the consumption of the midslope and Crest soils appears to be similar and less than $1 \times 10^{-4}$ m$^3$ kg$^{-1}$ s$^{-1}$. In all the soils the $k$ values were higher at the lower incubation temperatures.

The compensation point mixing ratio ($m_{\text{NO,comp}}$) was calculated according to Eq. (12) from laboratory measurements derived data of $P$ and $k$ (each as function of $T_{\text{soil}}$ and WFPS). If ambient NO equals $m_{\text{NO,comp}}$, then there is no net uptake or release of NO from the soil, since the production of NO equals the NO consumption in the soil. Figure 4 shows that $m_{\text{NO,comp}}$ at 25°C in each of the soils varies according to the soil moisture. The highest $m_{\text{NO,comp}}$ occurs in the Upslope soils where a $m_{\text{NO,comp}}$ of approximately 65–70 ppb is reached at 35% WFPS. The Footslope soils have the next highest $m_{\text{NO,comp}}$, with a peak of 55–60 ppb at 40% WFPS. The Crest and Downslope have a maximum $m_{\text{NO,comp}}$ between 45 and 50 ppb at 20% WFPS for the Downslope and 27% WFPS for the Crest.
3.2  \( T_{\text{soil}} \), WFPS, and \( F_{\text{NO, up-scaled}} \) at different landscape positions

Monthly mean soil temperature and volumetric soil moisture measured at 0.05 m depth at the Skukuza flux tower site are shown in Fig. 5 for the period 2003–2005.

The average monthly soil temperature ranged from 16.5°C to 36.5°C in winter and summer respectively (Fig. 5b and c). There was little difference between the soils in the Acacia savanna (Footslope position) and the soils in the Combretum savanna (Crest position); although in the winter months the Combretum soils tend to be slightly cooler. As there was only one measuring site in each vegetation type it is not known whether this is due to small-scale variability or whether it reflects a true difference.

In the Footslope soils there was always a higher soil WFPS than in the Crest soils (Fig. 5c), although they generally tracked each other quite closely during the dry periods. However in the high rainfall periods (Jan–Mar 2004, Dec 2004–May 2005 and Nov–Dec 2005, see Fig. 5d), the WFPS in the lower lying Footslope soils was much higher than that of the Crest soils. This is due to the drainage effects of the slope which resulted in water accumulating in the lower slope positions and the higher clay content of the Footslope would increase the water holding capacity of the Footslope soil.

Net up-scaled mean monthly NO fluxes \( (F_{\text{NO, up-scaled}}) \) at the four landscape positions were estimated from \( T_{\text{soil}} \) and WFPS and are shown in Fig. 5a for the period 2003–2005. In these soils, \( F_{\text{up-scaled}} \) ranges from less than the minimum detectable flux up to 3.5 ng m\(^{-2}\) s\(^{-1}\) and differs between the landscape positions (Fig. 5a). The \( F_{\text{up-scaled}} \) for the Downslope was generally lowest and never exceeded 3 ng m\(^{-2}\) s\(^{-1}\). The \( F_{\text{up-scaled}} \) is higher for the Footslope area than for the Crest and the Upslope soils except for a few periods during months of heavy rainfall (Fig. 5d). During these periods, the highest \( F_{\text{up-scaled}} \) came from the soils above the flux tower. The Crest and the Upslope soils generally followed a similar pattern, although the \( F_{\text{up-scaled}} \) from the Upslope soils tended to be slightly higher.

Of the four landscape positions the Footslope soils produced the highest NO emissions per unit area over the 2003–2005 period, between 0.5 and 0.8 kg ha\(^{-1}\) yr\(^{-1}\).
3.3 Estimates of NO emissions for the Skukuza land type area

When the area based annual NO fluxes were up-scaled to the entire Skukuza land type (Fig. 8), the total NO emission was $33 \times 10^3$ kg in 2003, $20 \times 10^3$ kg in 2004, and $25 \times 10^3$ kg in 2005 (all in terms of mass of nitrogen). When the total summed emission for the Skukuza land type is divided into the constitutive land landscape positions of the original Kruger National Park (KNP) Geographical Information System (GIS) data base, the midslope positions (= Upslope + Downslope) accounted for the majority of the emissions, due to their greater geographical extent.

4 Discussion

4.1 Net potential NO fluxes

In Fig. 9, we compare previously reported laboratory and field measurements of NO emissions from arid and semi-arid ecosystems with our net potential NO fluxes for the different landscape positions along a catenal sequence in the southern African Savanna of KNP. There have been two previous studies in the KNP, both of which were based on the use of field chambers (Levine et al., 1996; Parsons et al., 1996). In both studies, which occurred on long term fire return frequency experiments approximately 30 km away from the Skukuza flux tower, the measured NO field flux was between $0.34$ ng m$^{-2}$ s$^{-1}$ and $32$ ng m$^{-2}$ s$^{-1}$ at a range of soil moistures from 8.7% to 54% WFPS (Levine et al., 1996; Parsons et al., 1996). The higher values reported in the Levine et al. (1996) paper (see Fig. 9) can be attributed to measurements occurring under artificially irrigated conditions, which resulted in an increase in the emission of NO from the soil.

Three laboratory based measurements, similar to the method used in this study, have been made in southern Africa, these include studies in Nylsvley Savanna (Otter
et al., 1999), the Kalahari transect (Aranibar et al., 2004) and in Zimbabwe (Kirkman et al., 2001). In the Nylosvley Savanna in South Africa the NO flux ranged from 0.9–8 ng m\(^{-2}\) s\(^{-1}\) (Otter et al., 1999). For the study at Marondera (Zimbabwe) the NO flux ranged from 0.1–3.7 ng m\(^{-1}\) s\(^{-1}\) at soil moisture levels of between 9 and 15% WFPS (Kirkman et al., 2001). In a rainfall gradient transect measurement through the Kalahari it was found that the NO flux ranged from 0 to over 120 ng m\(^{-2}\) s\(^{-1}\) (Aranibar et al., 2004) although it should be noted that these values are a much higher than from the other similar sites (see Fig. 9).

In other natural arid and semi-arid regions (mean annual precipitation <700 mm) the measured median NO fluxes range from 0.07 ng m\(^{-2}\) s\(^{-1}\) to 5.3 ng m\(^{-2}\) s\(^{-1}\) (with reported values of up to 83 ng m\(^{-2}\) s\(^{-2}\) occurred during a pulsing event, such as when the soil is wetted or fertilized) (Davidson et al., 1993; Hartley and Schlesinger, 2000; Holst et al., 2007; Martin and Asner, 2005; Martin et al., 1998; Smart et al., 1999), the Aranibar et al. (2004) study showed a median value of 21 ng m\(^{-2}\) s\(^{-1}\) and is far out of the range of other reported values (Fig. 9). Therefore in comparison with the other published studies in arid and semi-arid ecosystems the net potential NO fluxes (\(F_{lab}\)) from this study are within the range reported for both field and laboratory measurements of NO. The NO fluxes from these arid and semi-arid ecosystems tend to be fairly low in comparison with some of the values reported from temperate and tropical forests where fluxes of 22 ng m\(^{-2}\) s\(^{-1}\) (Pilegaard et al., 2006) and 58 ng m\(^{-2}\) s\(^{-1}\) (Butterbach-Bahl et al., 2004) were reported in European coniferous and Australian tropical forests respectively.

4.2 Effect of landscape and vegetation

The biogenic emission of NO is controlled by the soil moisture, soil temperature and the soil nutrient status; it is through the combined effects of altering these modifiers of NO emission that landscape factors and vegetation can influence the production of NO from soils. Spatial and topographical factors influence the soil physical properties or the aboveground biomass; the soil physical properties and the biomass control the supply
and cycling on N in the soil (Akiyama and Tsuruta, 2002; Hartley and Schlesinger, 2000; Kirkman et al., 2001; Martin and Asner, 2005; Martin et al., 2003; Martin et al., 1998; Meixner et al., 1997). In the KNP the net N mineralization has been shown to change across soil textural sequences, and is highest in the low lying landscape positions, analogous to the Footslope position (Bechtold and Naiman, 2006; Scholes et al., 2003). Vegetation has been shown to have an effect on the emission of NO (Aranibar et al., 2004; Davidson, 1991a; Hartley and Schlesinger, 2000; Kirkman et al., 2001; Martin and Asner, 2005; Martin et al., 2003, 1998; Meixner et al., 1997; Ormeci et al., 1999; Otter et al., 1999; Pilegaard et al., 2006; Scholes et al., 1997; Serca et al., 1998; Van der A et al., 2008). This influence can be caused by the effect of:

1. Changes in the microclimate under the plant canopy, resulting in spatial heterogeneity in the soil through interacting biological and physical mechanisms;

2. Changes in the concentrations of soil nutrients; N, P and K have been shown to be enhanced under tree canopies, resulting from the relocation of absorbed nutrients and changes in the N mineralisation rates (Rossi and Villagra, 2003). The increase in nutrient contents under vegetation is either caused by the vegetation trapping or accumulating nutrients (Ludwig and Tongway, 1995; Ludwig et al., 1999), or through biological N fixation (Geesing et al., 2000; Schulze et al., 1991; Vitousek et al., 2002).

At the Skukuza flux tower site, the differences in the vegetation types along the catenal sequence have caused a significant difference in the amount of biological N fixation between the vegetation types. In the Combretum savanna (Crest position) the amount of N fixed is estimated at 4.8 kg ha\(^{-1}\) yr\(^{-1}\), while in the Acacia savanna (Footslope position) it is thought to amount to 21 kg ha\(^{-1}\) yr\(^{-1}\) (Scholes et al., 2003). Although the vegetation type has a marked influence on the biological N fixation, the deposition of N from the atmosphere in the form of both wet and dry deposition is constant across the vegetation types and amounts to 21.6 kg ha\(^{-1}\) yr\(^{-1}\) (Scholes et al., 2003).
These differences in the N cycling rate, the total amount of N in the system, and the amount of N entering the system by deposition from the atmosphere and biological nitrogen fixation, most likely account for the changes in the magnitude of the observed net potential NO flux. In this study there were clear differences in the NO production potential for the soils from the differing landscape positions. The highest potential NO flux occurred in the Footslope soils, which have been shown to have the highest total N content and N fixation rate. The lowest N emissions occurred in the less nutrient rich landscape positions (crest position Table 1) (Scholes et al., 2003; Woghiren, 2002). Loss of N from the ecosystem through the production of NO is in the range of 0.2–0.8 kg ha\(^{-1}\) yr\(^{-1}\) (Fig. 7) and therefore the loss of N in the form of NO is between 0.5% and 2.6% of the N that enters the ecosystem through biological N fixation and wet and dry deposition from the atmosphere.

4.3 Effect of soil moisture and temperature

The optimal soil moisture for the emission of NO from the four landscapes in the KNP was found to be between 10% and 20% WFPS. The optimal WFPS is the quantity of soil moisture, where the diffusion of NO through the soil is greatest, since diffusion is not limited by excessive soil moisture, while the bacterial metabolism, and therefore production of NO, is not limited by lack of soil moisture (Skopp et al., 1990). The optimum WFPS reported here correspond with the range reported in other studies in southern Africa; reported values lie between, 10% and 23% WFPS for a South African Savanna (Otter et al., 1999), a rainfall gradient in the Kalahari (Aranibar et al., 2004) and in Zimbabwe (Kirkman et al., 2001). Our study therefore confirms that the optimal NO flux occurs in a low and quite narrow range of WFPS in Southern African Savanna ecosystems (10–25%).

The landscape position affects the soil WFPS through changes in drainage and in the water holding capacity of the soil as influenced by the soil texture. Generally over the course of the year the \( F_{up-scaled} \) is higher for the Footslope than for the Crest or the Upslope soils since the soil WFPS is closer to the optimum. However during periods
of high rainfall, the highest \( F_{\text{up-scaled}} \) came from the soils above the flux tower which drained better, thereby maintaining a low soil moisture state more suitable for the production of NO. The Crest and the Upslope soils generally followed a similar pattern, although the \( F_{\text{up-scaled}} \) from the Upslope soils tended to be slightly higher (see Fig. 5d).

Many previous studies have reported an exponential increase in soil NO emission with increasing temperature, where the \( Q_{10} \) value is approximately 2 (Kirkman et al., 2001; Levine et al., 1996; Meixner and Yang, 2006; Van Dijk et al., 2002), which is similar to the \( Q_{10} \) values reported here (1.55–2.17). However, values of up to 4.6 have reported from the Kalahari (Aranibar et al., 2004) and from shortgrass steppe ecosystems (Martin et al., 1998). Under conditions such as very high temperatures (exceeding 40°C) there may even be a negative relationship the NO emission and the soil temperature (Passianoto et al., 2004), this indicates that an optimum soil temperature may exist, although due to the selection of incubation temperatures used in this study the position of the optimal temperature could not be determined.

The interplay between the influence of temperature and soil moisture content results in seasonal and annual patterns of NO emissions, in all the soils the lowest NO fluxes occurred during the southern hemisphere winter months when the soil temperatures were low and increased as the soil temperatures increased in spring (September, October) and remained high during the summer, unless the soil became too wet to allow optimal NO production.

The highest \( F_{\text{up-scaled}} \) occurred in 2003 and the lowest in 2004. The year 2003 was a relatively warm dry year which kept the soil WFPS low and in the range of the optimal temperature for NO production however, 2004 was a wet year and the soil was wetter than the optimum for extended periods (see Fig. 5d).

4.4 NO consumption rate and compensation mixing ratio

It is known that soils can both produce and consume NO, however there have been only a very few studies that have examined the uptake of NO in the soil, and none that have investigated the NO consumption rate constant (\( k \)) or the compensation point.
mixing ratio ($m_{\text{NO,comp}}$) across the full range of soil water contents.

The $k$ value found in this study is within the (very wide) range of values reported in the literature. The peak $k$ value was found to be at the same WFPS as the peak of NO production, between 10–20% WFPS and ranged between $5 \times 10^{-5}$ m$^{-3}$ s$^{-1}$ kg$^{-1}$ at 25°C (Fig. 3). At 35°C it was considerably lower for all the soil samples, less than $7 \times 10^{-5}$ m$^{-3}$ s$^{-1}$ kg$^{-1}$. The highest $k$ values were found in the Footslope soils, followed by the Crest soils. However, it appears to be a significant difference between the 2 mid-slope positions (Upslope and Downslope). To the best of our knowledge, there have only been four previous reports of $k$ from drylands; these are from Nylsvley savanna in South Africa (Otter et al., 1999), from the Kalahari transect in Botswana (Aranibar et al., 2004), from a study in Zimbabwe (Kirkman et al., 2001), and concerning an Egyptian soil (Saad and Conrad, 1993). The $k$ values found in our KNP soils were at least twice as high as those from the Nylsvley savanna and the Zimbabwe study where $k$ values of $1.3–3.2 \times 10^{-5}$ m$^{-3}$ s$^{-1}$ kg$^{-1}$ are reported. In comparison, however, the $k$ values reported from the Kalahari were up to two orders of magnitude higher than from the soils used in this study and ranged from $34–500 \times 10^{-5}$ m$^{-3}$ s$^{-1}$ kg$^{-1}$ (Aranibar et al., 2004). The $k$ values reported in the Egyptian soils ranged from $7.2 \times 10^{-5}$ m$^{-3}$ s$^{-1}$ kg$^{-1}$ at 7°C to $79 \times 10^{-5}$ m$^{-3}$ s$^{-1}$ kg$^{-1}$ at 25 and 30°C and 40% water holding capacity (Saad and Conrad, 1993). The $k$ values reported here fit right between the ranges of values previously reported. This, however, is the first time that the $k$ values have been reported across a full soil moisture scale; therefore, the cause of the wide range in reported $k$ values may be that previous measurements were made at different points along the soil moisture range. Soil temperature also has an effect on the $k$ values, although the Nylsvley, Kalahari transect and Zimbabwe studies all used an incubation temperature of 25°C.

The compensation point mixing ratio, $m_{\text{NO,comp}}$, has only been investigated in three studies in southern Africa:

1. $m_{\text{NO,comp}}$ of 152–157 ppb have been reported from Nylsvley (Otter et al., 1999).
2. Along a precipitation gradient in the Kalahari, the $M_{\text{NO,comp}}$ was reported to change with mean annual precipitation from 39 ppb at Tshane (365 mm Mean Annual Precipitation (MAP)) to 873 ppb for Mongu (880 mm MAP) (Aranibar et al., 2004),

3. Kirkman et al. (2001) reported that $M_{\text{NO,comp}}$ ranged from 5–11 ppb in the dry season in Zimbabwe and from 47–85 ppb during the wet season.

Our study, however, is the first to examine the effect of soil moisture on the $m_{\text{NO,comp}}$ covering as extensive a range of soil moisture contents. Within the WFPS range of the NO emission peak, the calculated $m_{\text{NO,comp}}$ in our study are between 40–70 ppb. Towards higher soil moisture conditions the compensation point mixing ratio decreases gradually to approx. 10 ppb (at 65% WFPS, see Fig. 4). The results of our study are within the range of values reported for Zimbabwe and Tshane (Kalahari), but much lower than those from Nylsvley and Mongu (Kalahari).

Knowing the WFPS dependent NO compensation point mixing ratios for the different Skukuza land types (as well as the average WFPS itself), the question could be tackled, whether the soils in KNP might ever act as a NO sink. The measured mean monthly WFPS at the Crest position ranges between 10–30% (Fig. 5c). NO uptake would only occur (c.f. Eqs. 5 and 12), if the ambient NO mixing ratio would be higher than $m_{\text{NO,comp}}$ (45–50 ppb, see Fig. 4). For the Footslope soils the recorded WFPS in the field occasionally reached levels of 60% (see Fig. 5c). Under these rather wet conditions the ambient concentration of NO, required before uptake could occur, would be much lower, only approximately 20 ppb. However, even an ambient NO mixing ratio of 20 ppb would be exceptionally high for such a remote location. Unfortunately, there are no measured data of ambient NO mixing ratio from Skukuza nor from the entire KNP region available. However, for comparable southern African sites, like Marondera in Zimbabwe, NO mixing ratios of 0.15–0.3 ppb have been reported (Meixner et al., 1997); aircraft measurements in the atmospheric boundary layer over northern Namibia revealed NO mixing ratios well below 0.5 ppb, even under conditions of high
biogenic soil emissions (approx. 30 ng m\(^{-2}\) s\(^{-1}\), see Harris et al. (1996)). Using the Max Planck Institute for Chemistry Atmospheric Chemistry Model MESSy (“Modular Earth Sub-model System”, Jockel et al., 2005), it was found that the modelled surface NO mixing ratio never exceeded 5 ppb, and that the mean NO mixing ratio was 0.08 ppb for the period January 2003–October 2005. It is therefore highly unlikely that the ambient NO mixing ratio has exceeded the \(m_{\text{NO,comp}}\) during 2003–2005. Considering the remoteness of the KNP region and the well established status as a National Park, it can be expected, that the KNP region is continuously acting as a biogenic NO source and not as a sink (like the majority of the southern African savanna ecosystems).

5 Conclusions

This laboratory study investigated the biogenic emission of NO from soils taken from four differing landscape positions in a semi-arid savanna under conservation land use (Skukuza, Kruger National Park, South Africa). The up-scaled emissions of NO did not exceed 3.5 ng m\(^{-2}\) s\(^{-2}\) for any of the landscape positions, even under optimal soil moisture conditions and high soil temperature; this is at the lower end of the range of previously published studies from comparable ecosystems. The optimum emission of NO occurred under fairly low soil water contents, between 10–20% WFPS. The highest NO flux came from the nutrient rich fine textured soils in the Footslope position, while the Downslope, Upslope and Crest positions had lower rates of NO emission. When the laboratory derived, net potential NO fluxes were up-scaled to the landscape level, the highest estimated NO fluxes were calculated to come from the sum of Downslope and Upslope landscape positions, because they make up the majority of the Skukuza landscape surface area. The biogenic emissions of NO from this ecosystem constitute a minor N loss mechanism and account for less than 2.6% of the N entering the system in the form of wet/dry deposition and biological N fixation. Although soils have generally both, the potential to emit and to take up atmospheric NO, the ambient atmospheric NO mixing ratios in this remote part of Africa are highly unlikely to reach and exceed 2818.
that level (i.e. the NO compensation point mixing ratio), where up-take of NO will take place. The laboratory based estimation of soil moisture and soil temperature dependent net potential NO fluxes from soil and the up-scaling procedure (using recorded soil moisture and temperature data from the field) is shown to be an effective method to quantify biogenic NO emissions on large temporal (annual) and spatial (regional) scales.

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Scholes, R. J., Gureja, N. G., Giannecchinni, M., Dovie, D., Wilson, B., Davidson, N., McLough-


Table 1. Physical and chemical properties of Skukuza KNP land type 971 soils (all units are in terms of mass of N and mass of dry soil; numbers in brackets indicate the standard deviation of the measurements; values taken from Woghiren (2002) are marked by a star (*)).

<table>
<thead>
<tr>
<th>Units</th>
<th>Footslope</th>
<th>Downslope</th>
<th>Upslope</th>
<th>Crest</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk Density</td>
<td>kg m⁻³</td>
<td>1.50×10³</td>
<td>1.40×10³</td>
<td>1.56×10³</td>
</tr>
<tr>
<td>Sand content</td>
<td>%</td>
<td>66.8</td>
<td>74.9</td>
<td>72.8</td>
</tr>
<tr>
<td>Silt content</td>
<td>%</td>
<td>12.1</td>
<td>10.0</td>
<td>10.1</td>
</tr>
<tr>
<td>Clay content</td>
<td>%</td>
<td>21.1</td>
<td>15.1</td>
<td>17.2</td>
</tr>
<tr>
<td>Texture</td>
<td></td>
<td>Sandy Clay Loam</td>
<td>Sandy Loam</td>
<td>Sandy Loam</td>
</tr>
<tr>
<td>pH</td>
<td>6.25±0.06</td>
<td>6.08±0.13</td>
<td>6.36±0.06</td>
<td>6.48±0.09</td>
</tr>
<tr>
<td>Total N*</td>
<td>kg ha⁻¹</td>
<td>4635</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Ammonium*</td>
<td>µg g⁻¹</td>
<td>0.15–2.41</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Nitrate*</td>
<td>µg g⁻¹</td>
<td>0.21–2.04</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Mean N</td>
<td>%</td>
<td>0.08% (0.00)</td>
<td>0.07% (0.01)</td>
<td>0.05% (0.00)</td>
</tr>
<tr>
<td>Mean C</td>
<td>%</td>
<td>0.88 (0.01)</td>
<td>0.83 (0.02)</td>
<td>0.59 (0.04)</td>
</tr>
<tr>
<td>Slope Class</td>
<td>%</td>
<td>0–2%</td>
<td>2–6%</td>
<td>6–15%</td>
</tr>
</tbody>
</table>
**Table 2.** Optimum soil moisture, maximum NO flux and $Q_{10}$ value of Skukuza KNP land type soils (mass units are in terms of mass of N); (*) measurement failed ($Q_{10}=2.17$ is used for the Crest soil).

<table>
<thead>
<tr>
<th></th>
<th>Footslope</th>
<th>Downslope</th>
<th>Upslope</th>
<th>Crest</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optimum WFPS at 25°C</strong></td>
<td>17%</td>
<td>14%</td>
<td>11%</td>
<td>12%</td>
</tr>
<tr>
<td><strong>Maximum net potential NO flux at 25°C</strong> (ng m$^{-2}$ s$^{-1}$)</td>
<td>2.4</td>
<td>1.3</td>
<td>1.7</td>
<td>1.5</td>
</tr>
<tr>
<td><strong>Optimum WFPS at 35°C</strong></td>
<td>18%</td>
<td>17%</td>
<td>18%</td>
<td>18%</td>
</tr>
<tr>
<td><strong>Maximum net potential NO flux at 35°C</strong> (ng m$^{-2}$ s$^{-1}$)</td>
<td>3.5</td>
<td>3.0</td>
<td>3.5</td>
<td>No Data</td>
</tr>
<tr>
<td><strong>$Q_{10}$ Value</strong></td>
<td>1.55</td>
<td>2.11</td>
<td>2.17</td>
<td>(*)</td>
</tr>
</tbody>
</table>
Fig. 1. (a) Map of the southern section of the Kruger National Park indicating the position of the Skukuza land type and the flux tower site, the position of the Kruger National Park in South Africa is shown in the insert. (b). A schematic diagram (not to scale) showing the position of the Skukuza flux tower in relation to the main ecosystem types, the smaller towers at the top and bottom of the slope are where the soil moisture and soil temperature measurements were taken.
Fig. 2. Net potential NO flux ($F_{lab}$) from the four landscape positions, Footslope (a), Downslope (b), Upslope (c), and Crest (d) as a function of soil moisture (WFPS) and soil temperature (mass of NO is expressed in terms of mass of nitrogen).
Fig. 3. NO consumption rate ($k$) as a function of WFPS for the 4 landscape positions at 25°C and at 35°C, solid lines represent the $k$ values at 25°C, while the dashed lines represent $k$ at 35°C.
Fig. 4. Compensation point mixing ratio ($m_{\text{NO,comp}}$) at $T_{\text{soil}}=25^\circ\text{C}$ as a function of WFPS for soils of the four landscape positions in the Kruger National Park.
Fig. 5. (a) Monthly averages of the NO flux ($F_{\text{NO,up-scaled}}$) from soils of the four landscape positions calculated for the period January 2003–October 2005 (error bars represent the daily variability expressed as standard error of the daily averages; mass of NO expressed in terms of mass of nitrogen)). (b) Recorded soil temperature for the period 2003-2005 (c). Recorded soil moisture (WFPS) for the period 2003–2005 Average monthly soil moisture contents and temperatures calculated from half hourly recordings in the fine-leaved Acacia savanna (Footslope), and the broad-leaved Combretum (Crest). Error bars indicate the monthly variability of the measurements (expressed as standard error) (d). Rainfall for the period 2003–2005.
Fig. 6. Map of the Skukuza land type showing the distribution of the different landscape positions.
Fig. 7. Annual means of the up-scaled NO fluxes (FNO,up-scaled) from soils of the 4 landscape positions for the period 2003–2005; error bars represent the standard deviation of the means (mass, of NO is expressed in terms of mass of nitrogen).
Fig. 8. Total area weighted NO emissions (expressed in terms of mass of nitrogen) from each of the four landscape positions in the Skukuza land type for the period 2003 to 2005; errorbars represent the standard deviation of the means.
Fig. 9. Comparison of the up-scaled NO fluxes with data reported in the literature for comparable arid and semi arid ecosystems (all expressed in terms of mass of nitrogen). Boxes in the figure indicate the inter-quartile range between the 25 and 75 percentile and the median value is marked by a vertical bar. The (horizontal) error bars represent the 10 and 90 percentile. The open circle represents the arithmetic mean of the reported data. Where the cited papers provide a mean and range of data only, values were statistically recreated to match these ranges. The reported ecosystem NO flux data is cited from:

- Semi-arid grassland (Inner Mongolia, China) (Holst et al., 2007);
- Chihuahuan Desert (New Mexico, USA) (Hartley and Schlesinger, 2000);
- Shortgrass Steppe (Colorado, USA) (Martin et al., 1998);
- Sagebush Steppe (Utah, USA) (Smart et al., 1999);
- Temperate Savanna (Texas, USA) (Martin et al., 2003);
- Dry Tropical Forest (Mexico) (Davidson et al., 1993);
- Savanna (Zimbabwe) (Kirkman et al., 2001);
- Kalahari Savanna (southern Africa) (Aranibar et al., 2004);
- Savanna (RSA)* (Parsons et al., 1996);
- Savanna (RSA) (Levine et al., 1996)