TRANC – a novel fast-response converter to measure total reactive atmospheric nitrogen

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Abstract. The input and loss of plant available nitrogen (reactive nitrogen: $N_r$) from/to the atmosphere can be an important factor for the productivity of ecosystems and thus for its carbon and greenhouse gas exchange. We present a novel converter for reactive nitrogen (TRANC: Total Reactive Atmospheric Nitrogen Converter), which offers the opportunity to quantify the sum of all airborne reactive nitrogen compounds ($\sum N_r$) in high time resolution. The basic concept of the TRANC is the full conversion of all $N_r$ to nitrogen monoxide (NO) within two reaction steps. Initially, reduced $N_r$ compounds are being oxidised, and oxidised $N_r$ compounds are thermally converted to lower oxidation states. Particulate $N_r$ is being sublimated and oxidised or reduced afterwards. In a second step, remaining higher nitrogen oxides or those generated in the first step are catalytically converted to NO with carbon monoxide used as reduction gas. The converter is combined with a fast response chemiluminescence detector (CLD) for NO analysis and its performance was tested for the most relevant gaseous and particulate $N_r$ species under both laboratory and field conditions. Recovery rates during laboratory tests for NH$_3$ and NO$_2$ were found to be 95 and 99 %, respectively, and 97 % when the two gases were combined. In-field longterm stability over an 11-month period was approved by a value of 91 % for NO$_2$. Effective conversion was also found for ammonium and nitrate containing particles. The recovery rate of total ambient $N_r$ was tested against the sum of individual measurements of NH$_3$, HNO$_3$, HONO, NH$_4^+$, NO$_3^-$, and NO$_x$ using a combination of different well-established devices. The results show that the TRANC-CLD system precisely captures fluctuations in $\sum N_r$ concentrations and also matches the sum of all individual $N_r$ compounds measured by the different single techniques. The TRANC features a specific design with very short distance between the sample air inlet and the place where the thermal and catalytic conversions to NO occur. This assures a short residence time of the sample air inside the instrument, and minimises wall sorption problems of water soluble compounds. The fast response time (e-folding times of 0.30 to 0.35 s were found during concentration step changes) and high accuracy in capturing the dominant $N_r$ species enables the converter to be used in an eddy covariance setup. Although a source attribution of specific $N_r$ compounds is not possible, the TRANC is a new reliable tool for permanent measurements of the net $\sum N_r$ flux between ecosystem and atmosphere at a relatively low maintenance and reasonable cost level allowing for diurnal, seasonal and annual investigations.

1 Introduction

1.1 Reactive nitrogen

Nitrogen is an essential nutrient for all living organisms. Beside dinitrogen (N$_2$), being practically inert and constituting 78 % of the earth’s atmosphere (Seinfeld and Pandis, 2006), the important nitrogen-containing trace species are nitric oxide (NO, also nitrogen monoxide), nitrogen dioxide (NO$_2$), nitric acid (HNO$_3$), ammonia (NH$_3$), and nitrous oxide (N$_2$O) (Sutton et al., 2011). The sum of NO and NO$_2$, the former emitted by both natural and anthropogenic sources, the latter formed in the atmosphere by oxidation of NO and
emitted in small quantities from combustion processes along with NO, is usually designated as NOx. Reactive odd nitrogen, denoted NOy, is defined as the sum of NOx and all products of the atmospheric oxidation of NOx. These include among others HNO3, nitrous acid (HONO), and peroxyacetyl nitrate (PAN). HNO3 is rapidly deposited on surfaces and in water droplets and – in the presence of NH3 – can form ammonium nitrate (NH4NO3) aerosols. Significant sources of NH3 are animal waste, ammonification of humus followed by emission from soils, losses of NH3-based fertilizers, and industrial emissions. In this paper, we define reactive nitrogen (N\textsubscript{r}) as all nitrogen-containing trace species except for N\textsubscript{2} and N\textsubscript{2}O with the latter being inert in the troposphere (Seinfeld and Pandis, 2006). The oxidation states of the main N\textsubscript{r} compounds range from +5 for HNO\textsubscript{3} to −3 for NH3.

The input of N\textsubscript{r} into an ecosystem through atmospheric dry deposition is an important factor for its productivity and thus for its exchange of carbon dioxide (CO\textsubscript{2}) and other greenhouse gases. The dry deposition of N\textsubscript{r} typically contributes between one third and two thirds to the total atmospheric N deposition (Simpson et al., 2006). The environmental effects of excess atmospheric N\textsubscript{r} deposition to ecosystems may include soil acidification, eutrophication of water bodies, nutrient imbalances, leaching of base cation and nitrate, loss of biodiversity, direct toxicity to plants, increased N\textsubscript{2}O emissions, and inhibition of soil methane (CH\textsubscript{4}) oxidation (Galloway et al., 2003; Erismann et al., 2007; Flechard et al., 2011).

1.2 Exchange measurements of reactive nitrogen compounds

Unlike wet deposition, which is widely monitored in regional networks of wet-only or bulk precipitation collectors, measurements of overall dry (turbulent) N\textsubscript{r} exchange fluxes have largely remained experimental and were limited to selected research sites and to measurement campaigns of typically a few days to a few months due to technical complexity and to the large equipment and operational costs involved (Flechard et al., 2011) except for multi-year NO\textsubscript{y} eddy flux measurements at Harvard forest presented in Munger et al. (1996, 1998) and in Horii et al. (2006). This is due to the fact that dry N\textsubscript{r} exchange constitutes a variety of compounds with different exchange patterns and that in general, concentration and flux measurements of N\textsubscript{r} species are challenging from a metrological point of view (e.g. von Bobrutzki et al., 2010; Wolff et al., 2010a). Well-established measurement techniques (e.g. using chemiluminescence detectors CLD, molybdenum oxide (MoO) or a gold (Au) converter operating at temperatures between 275 and 390°C, whereas Au converters need a reducing agent like CO or H\textsubscript{2}. Early work by Sigsby et al. (1973) and Winer et al. (1974) describe a chemiluminescent method for analysis of nitrogen containing compounds in mobile source emissions (NO, NO\textsubscript{2}, NH\textsubscript{3}) and the response of chemiluminescent NO-NO\textsubscript{2} analyzers to other nitrogen-containing compounds while stressing the importance of correcting for interferences, e.g. under high PAN levels, respectively. Using fast-response NO detection by CLD, some approaches of long-term NO\textsubscript{y} measurements using EC have been conducted. For example, Munger et al. (1996), report on five years of NO\textsubscript{y} eddy flux data measured at Harvard forest, where NO\textsubscript{y} was determined following reduction to NO by H\textsubscript{2} on a Au catalyst.

EC-based measurements of N\textsubscript{r} compounds other than NO\textsubscript{y} are extremely limited. Day et al. (2002) and Farmer et al. (2006) present the application of a thermal dissociation laser-induced fluorescence (TD-LIF) instrument suitable for the measurement of NO\textsubscript{2}, HNO\textsubscript{3}, alkyl (\textsubscript{\Sigma}AN) and peroxy...
nitrates ($\sum PN$). In this approach, each class of compounds thermally dissociates to NO$_2$ and an accompanying radical (RO$_2$, RO, OH) in separately thermostatted ovens at temperatures of 180, 330, and 550 °C for $\sum PN$, $\sum AN$, and HNO$_3$, respectively. Sintermann et al. (2011) use a system for fast NH$_3$ measurements with chemical ionisation mass spectrometry (CIMS) based on a commercial Proton Transfer Reaction-Mass Spectrometer (PTR-MS) within an EC setup over two agricultural fields in Oensingen, Switzerland, whereas the EC fluxes of NH$_3$ measured by Fanumari et al. (2004) during a two-month field campaign were conducted by the use of a TDLAS system.

In the absence of fast-response instruments suitable for EC measurements, the aerodynamic gradient method is frequently applied for N$_r$ compounds such as NH$_3$, HNO$_3$, HONO, and aerosol particles. It infers the flux from concentration measurements at several heights and measured turbulence characteristics. Concentration measurements are frequently performed by denuder or filter-pack sampling in combination with on- or offline analysis by ion chromatography and/or flow injection analysis, e.g., for NH$_3$ in the AMANDA system (Wyers et al., 1993), or for HNO$_3$ and aerosol particles in the GRAEGOR system (Thomas et al., 2009; Twigg et al., 2011). However, this method has proved to be relatively labour and cost extensive, often shows limited time resolution and brings along method-linked limitations. Only very few long term measurements are published (Flechard and Fowler, 1998; Flechard et al., 2010).

Measurements of total N$_r$ have been reported by McCalley and Sparks (2009) using selective thermal and chemical decomposition converters to reduce or oxidise all N$_r$ trace gases to NO. However, their setup was applied for relatively slow enclosure measurements over short experimental periods to determine the soil-atmosphere exchange of NO, NO$_x$, and NH$_3$ differentially by switching between different converter configurations. No information is given on the influence of the enclosures and sampling tubes in the field configuration on the performance of the system.

Although the knowledge of total N$_r$ exchange is of major importance for the compilation of matter flux balances on ecosystem level, none of the above-mentioned methods is capable of determining total N$_r$ in high time resolution within longterm monitoring setups. The reasons for this limitation are (a) the wide range of N$_r$ containing compounds, their different molecular and exchange characteristics and thus different sampling and measuring approaches, (b) the interactions between different N$_r$ compounds due to gas phase reactions (e.g. Meixner, 1994) and gas-aerosol particle interactions (e.g. Wolff et al., 2010b), and therefore the necessity to measure several parameters simultaneously, and (c) the complexity and labour-intensity of flux measurements of the individual compounds.

### 1.3 Motivation and objectives of present study

In this study, we present a novel converter named TRANC (Total Reactive Atmospheric Nitrogen Converter), which, in combination with a fast-response analyser (CLD), offers the opportunity to quantify the sum of all airborne N$_r$ compounds (denoted here as $\sum N_r$) in high time resolution. The design of the TRANC is supposed to fulfil the requirements needed for an operation within an EC setup. These requirements are an instantaneous oxidation and reduction of all N$_r$ compounds in the sample air to NO with the conversion occurring close to the air inlet, thereby reducing errors due to chemical reactions, particularly of NH$_3$, in the sampling tube. To our knowledge, this is the first approach that is on the one hand not limited to a single or a few N$_r$ compounds, but provides the quantification of the sum of all N$_r$ compounds, and that is on the other hand fast enough to be used for EC measurements if combined with a fast-response NO analyser, thus presenting a new methodology for permanent exchange measurements of $\sum N_r$ at a relatively low maintenance level. The purpose of this paper is the presentation of the converter, its characteristics and results of performance tests. Validation and long-term application of the converter for EC flux measurements is presented in companion papers by Ammann et al. (2012) and Brümmer et al. (2012b).

### 2 Experimental setup

#### 2.1 Converter principle and design

The basic concept of the TRANC is the full conversion of all N$_r$ compounds in the sample air to NO in two reaction steps. Initially, reduced N$_r$ compounds are being oxidised, whereas oxidised N$_r$ compounds are thermally converted to compounds of lower oxidation states. Particulate N$_r$ is being sublimated and oxidised or reduced afterwards. The conversions in the first reaction step, i.e. the thermal conversions, occur at temperatures $\geq$870 °C. In a second step, i.e. the catalytic conversion, remaining higher nitrogen oxides in the sample air or those originated in the first reaction step are converted on a Au surface ($\geq$300 °C) to NO with carbon monoxide (CO) used as reduction gas and by installing a platinum (Pt) gauze ensuring a complete catalytic conversion of NH$_3$ to NO. The main conversions resulting under these conditions are as follows:

\[
\begin{align*}
\text{NH}_4\text{NO}_3 & \xrightarrow{\Delta T} \text{NH}_3 \uparrow + \text{HNO}_3 \uparrow & \text{(R1)} \\
2 \text{HNO}_3 & \xrightarrow{\Delta T} 2 \text{NO}_2 + \text{H}_2\text{O} + 1/2 \text{O}_2 & \text{(R2)} \\
\text{HONO} & \xrightarrow{\Delta T} \text{NO} + \text{OH} & \text{(R3)} \\
4 \text{NH}_3 + 5 \text{O}_2 & \xrightarrow{\Delta T, \text{Pt}} 4 \text{NO} + 6 \text{H}_2\text{O} & \text{(R4)} \\
\text{NO}_2 + \text{CO} & \xrightarrow{\Delta T, \text{cat}} \text{NO} + \text{CO}_2 & \text{(R5)}
\end{align*}
\]
Finally, the NO concentration in the gas sample leaving the converter towards a CLD represents the initial $\sum N_r$ concentration that entered the converter through the sample air inlet. Regarding decomposition temperatures of the most abundant $N_r$ compounds being significantly lower than 870 and 300 °C during the thermal and catalytic reaction steps, respectively, we assume a full conversion of all $N_r$ in the sample air to NO. The conversion efficiency of single $N_r$ compounds was tested in several laboratory and field experiments and is described in Sects. 3.1 and 3.2.

A schematic overview of the TRANC is presented in Fig. 1. Core piece of the converter is a high-temperature resistant metal tube (length $l = 780 \text{ mm}$, outer diameter OD = 6.35 mm, inner diameter ID = 4.57 mm) consisting of an iron-nickel-chrome (FeNiCr) alloy. The tube is encapsulated in a double-walled, weather-sealed aluminium housing ($l = 705 \text{ mm}$, OD = 120 mm of outer box). The inner box ($l = 650 \text{ mm}$, OD = 90 mm) is filled with high-temperature resistant concrete granulate assuring effective heat insulation. Outside the outer housing, the FeNiCr tube is connected to a Pt gauze followed by a tee junction serving as gas inlet for CO and a stainless steel loop leading the sample air back to the inside of the TRANC. Here, the Au tube ($l = 490 \text{ mm}$, OD = 6 mm, ID = 4 mm), i.e. the location where the catalytic conversion occurs, is mounted. Another loop at the back end of the Au tube leads the stainless steel pipe outside the converter, where a 4 µm filter and a critical orifice assure the desired pressure drop (see Sect. 2.2) before the sample air passes through lightproof tubing and enters the CLD for NO analysis. Residence time in the converter at a flow rate of 5.1 min$^{-1}$ is $\sim 0.73$ s.

An operating temperature of $> 800$ °C is needed to fully oxidise the reduced $N$ compounds (McCalley and Sparks, 2009, Supplement). Such high temperatures also convert oxidised $N$ compounds like HNO$_3$ to lower oxidation states (Reaction 2; see e.g. Day et al., 2002). These conditions for the thermal conversion are implemented by the resistance-heated FeNiCr tube. A current of 80 A at a voltage of 5 V is fed to the tube over brass blocks at the two end fittings. Provision of electricity is effected by a switch-mode power supply with the temperature of the tube being regulated by a universal modulator and controlled by a thermocouple (see Fig. 1). The Au tube is indirectly warmed up to approximately 300 °C by the waste heat of the FeNiCr tube. The tube lead-throughs of both the inner and outer housing are electrically insulated by ceramic material except for two small additional inlets that are connected over flexible tubes with two pumps assuring permanent ventilation and removal of hot air between the inner and the outer housing at a flux rate of 5 l min$^{-1}$ each (not shown in Fig. 1).

Figure 2 shows the temperature distribution inside the FeNiCr and the Au tube during different experimental runs. The target temperature of the heating modulator was set to 895 °C to ensure temperatures above the desired values of 870 and 300 °C over a significant part of the FeNiCr and the Au tube, respectively. The direct heating of the FeNiCr tube ensures temperatures above 100 °C from the first millimetre of the inlet, although the target temperature is only reached in the heat-insulated part of the converter. The generally hot inner surfaces are supposed to minimise wall interaction effects for the sampled $N_r$ compounds.

### 2.2 Combination of TRANC with fast-response NO analysis

The TRANC is the key part of a comprehensive measuring system consisting of a CLD connected to a dry vacuum scroll pump (BOC Edwards XDS10, Sussex, UK) and a calibration unit (see Sect. 3.1) with a multi-gas calibrator and specific reference gases. In our setup, we used a commercial CLD (780 TR, ECO PHYSICS, Dürnten, Switzerland) for NO analysis. The measurement principle of this instrument is based on gas phase titration, i.e. the reaction of NO with O$_3$. It generates electronically excited NO$_5$ molecules, which rapidly decay to their ground state in a reduced pressure regime by emitting photons. The total light intensity in the reaction chamber, detected by a photomultiplier tube (PMT), is proportional to the NO mixing ratio (Fontijn et al., 1970; Rummel et al., 2002). To minimise electrical noise, the
temperature of the PMT is held constant at $-18 \, ^\circ C$. Flushing with oxygen or dry air avoids condensation. An ozone generator supplies the main chamber of the CLD with $O_3$-enriched air, where it reacts with NO to NO$_2$ and O$_2$.

To reduce the probability of energy loss of NO$_2$ molecules by collision with others, the gas phase titration in the main chamber occurs under low pressure, i.e. $<20\, hPa$. This is effected by the critical orifice located at the back end of the stainless steel tube that is leaving the TRANC towards the CLD (Fig. 1) and restricts the flow to $2.71\, \text{min}^{-1}$. In our setup, the instrument was running in a continuous mode, integrating the photons over 0.05 s. In the standard operation procedure, the CLD 780TR uses so-called pre-chamber measurements in order to correct for interferences which might be caused by e.g. hydrocarbons in the sample air that react with O$_3$ in a similar way like NO and are therefore detected as NO in the main reaction chamber. For fast response measurements this option is disabled. Frequent calibration aims at minimising related errors. Furthermore we estimate that the relative contribution of possible interferences is small in comparison to the $\sum N_r$ concentration. As O$_3$ is completely destroyed in the TRANC (thermal decomposition and reduction by CO), chemical NO loss by gas phase reaction ($NO + O_3 \rightarrow NO_2 + O_2$) could also be neglected. Even if a small amount of O$_3$ would pass the TRANC, the influence on concentration changes are expected to be lower than the detection limit of the CLD according to Rummel et al. (2002). A schematic overview of the TRANC-CLD system in an EC-setup is given in Fig. 3.

### 3 Results of performance tests

#### 3.1 Calibration and conversion of individual $N_r$ compounds

During field application, the TRANC-CLD unit was calibrated by regularly feeding calibration gas with different NO and NO$_2$ concentrations through the system (Fig. 3). In our setup, we used a multi-gas calibrator (S6100, Environics Inc., Tolland, USA) connected to two different reference gas cylinders containing 10 ppm NO and 10 ppm NO$_2$ each as well as a pure air generator (PAG 003, ECO PHYSICS, Dürnten, Switzerland). The calibrator mixed the desired gas concentration by dilution of the cylinder standard with zero air. Potential loss of NO in the TRANC converter, particularly with respect to the usage of a Pt-gauze catalyst (see Kliner et al., 1997) could be excluded as no significant reduction of NO concentration was found when adding NO calibration gas upstream and downstream of the TRANC.

We used gas concentrations of 6, 10, 20, and 50 ppb NO as well as pure air to calibrate the TRANC-CLD system. Additionally 50 ppb NO$_2$ and 50 ppb NO mixed with O$_3$ ($<50\, \text{ppb}$) was given into the system to determine the conversion efficiency of NO$_2$. During a typical calibration procedure, the calibrator started mixing the target concentration 3.5 h prior to the actual calibration run when the reference gas was fed into the system. At this time, the calibration valve near the TRANC inlet (Fig. 3) was still shut and the calibration gas was vented directly from the calibrator unit. After activating the TRANC valve, the system was calibrated for 30 min with one specific gas concentration. As the reference gas was fed into the tubes with a higher flow rate (approximately $51\, \text{min}^{-1}$) than that established by the vacuum pump and the critical orifice ($2.71\, \text{min}^{-1}$), the excess gas was leaving the system through the sample air inlet at the front side of the TRANC. After 30 min the TRANC valve was switched off to set the system back to the measurement mode for ambient air. To assure a well conditioned calibration gas tube and therefore a stable calibration gas concentration at the sample air inlet of the TRANC, only data of the 10-min period from 15 to 25 min after the initial switch were taken to calculate the NO concentrations. If necessary, spikes were removed and the 10-min window was shifted to the most stable interval during the respective half-hour calibration period. Over the field measurement period (see Fig. 4a), not a single calibration run was observed, where the 30-min interval had to be extended and data were not discarded due to other system instabilities.

For optimal data coverage during long-term measurement campaigns, the system was configured to calibrate for 30 min with only one gas concentration once a night, in this case from 03:30 to 04:00 a.m LT. The data obtained in these eight consecutive nights were used for the calibration regression of
Fig. 3. Schematic overview of the TRANC-CLD system (blue) in an eddy-covariance setup. Elements of the calibration unit are shown in red. Bold lines indicate tubing. The photo shows the field-setup at the Oensingen site.

this time period. With a new calibration interval starting on
day nine, the regression was updated with the data recorded
from day 9 to 16, accordingly. These steps with the respec-
tive regression update were repeated for all following 8-day
periods.

A low positive NO signal slightly different from zero with
a mean offset of 0.055 ($\pm$0.028) (Fig. 4b) was found while
calibrating with pure air. Beside the interference with hy-
drocarbons in the sample air (see Sect. 2.2) or impurities in
the added CO (Hegglin et al., 2006), this could have been
induced by the pure air generator through a small contami-
nation with any other $N_r$ compound. The overall calibration
performance of a multi-month observation period is given in
Fig. 4 (panels a and b). These data were taken from a field
campaign conducted at an agricultural site in Gebesee, Ger-
many, which will be described in more detail in Brümmer
et al. (2012b). $R^2$ values from the correlation between the
given calibration gas concentration and the corresponding
CLD signals of single 8-day periods were ranging between
0.81 and 0.99 with $R^2 = 0.98$ for the averaged fit ($n = 30$
for each gas concentration). In most cases, intercepts were found
to be slightly positive (Fig. 4b).

Over the same 11-month observation period, we found a
mean conversion efficiency for NO$_2$ of 91 % (Fig. 5) with
a standard deviation of $\pm$9 %. It has to be considered that

Fig. 4. Left panel: long-term stability of mean signal response of
the CLD coupled to the TRANC after calibrating with different
concentrations. Black squares: 0 ppb NO; red diamonds: 6 ppb NO;
yellow triangles: 10 ppb NO; green triangles: 20 ppb NO; blue trian-
gles: 50 ppb NO; blue open squares: 50 ppb NO$_2$; blue open circles:
50 ppb NO + O$_3$. DOY 0 represents 15 December 2007. Right panel:
averaged signal response of left panel data plotted against nominal
concentration. Data were taken from an 11-month field campaign
conducted at an agricultural site in Gebesee, Germany. Each data
point represents the mean of those calibration runs shown in the left
panel. Error bars indicate the standard error of the mean. Filled sym-
bols are NO calibrations, open square and open triangle are NO$_2$
and NO + O$_3$ calibration runs, respectively.
cylinder calibration standards for NO$_2$ are generally less accurate than for NO, because of possible non-ideal mixing in the gas cylinder over longer periods. The conversion rate for in-situ produced NO$_2$, resulting from 50 ppb NO mixed with O$_3$, was found to be 105%. We observed no correlation between any of the conversion efficiencies and meteorological parameters like air temperature, relative humidity, wind speed and direction, or precipitation events as an effect of possibly poor heat insulation.

### 3.1.2 Recovery rates of NH$_3$ and mixed sample gas (NO$_2$ and NH$_3$)

Since the preparation of reliable and accurate calibration gas mixtures for reactive nitrogenous gases is generally difficult, we concentrated the conversion efficiency tests for the TRANC on NO$_2$ and NH$_3$, the most abundant oxidized and reduced N$_r$ compounds, respectively, in central Europe (see e.g. Hesterberg et al., 1996; Flechard et al., 2011). The conversion of NH$_3$, NO$_2$ and a mixture of both gases was tested by parallel sampling of calibration gases by the TRANC-CLD system and specific NH$_3$ and NO$_2$ analysers. The NH$_3$ analyser (Picarro G1103, Sunnyvale, CA, USA) was calibrated prior to the measurements using a permeation oven manufactured by LN Industries (Geneva, Switzerland) equipped with a NH$_3$ permeation tube (VICI, Metronics Inc., Poulsbo, WA, USA). The permeation system was calibrated by directing the gas mixture through an impinger containing acidic solution (H$_2$SO$_4$, 0.01 m$^{-1}$) and subsequent analysis by ion chromatography. The NO$_2$ analyser (Thermo Environmental chemiluminescence NO-NO$_2$-NO$_x$ analyser, 42C Trace Level, called TEI hereafter), a chemiluminescence detector, in principle operating identical to the CLD described earlier but containing an additional channel for NO$_2$ measurements making use of a NO$_2$ specific photolytic converter (blue light converter BLC, Air Quality Design Inc., Wheatridge, USA; see e.g. Pollack et al., 2010) was calibrated with a NO standard parallel to the CLD. Both instruments thus agreed in NO concentrations. The conversion efficiency of the build-in NO$_2$ converter of the TEI was determined during an instrument service just after the conversion tests.

Conversion tests for NH$_3$ were performed by comparing the TRANC-CLD signal to NH$_3$ concentrations as determined by the NH$_3$ analyser. Both instruments sampled at the overflow of the same calibration tube which was flushed with the calibration gas for 48 h to ensure a stable NH$_3$ concentration. The TRANC-CLD system found 112.6 ± 0.7 ppb N$_r$, which corresponds to a conversion efficiency of 95% compared to NH$_3$ concentrations of 118.8 ± 0.6 ppb. Using the same setup for NO$_2$, the conversion efficiency was observed to be larger than 99% with 45.3 ± 0.4 ppb detected by the TRANC-CLD system in comparison to 45.5 ± 0.3 ppb NO$_2$. Finally, a mixture of NH$_3$ and NO$_2$ calibration gas was fed to the analysers (see Fig. 5). To ensure excellent conditioning of the calibration tube, it was flushed with NH$_3$ calibration gas for 3 days, adding the NO$_2$ calibration gas after 2 days. The calibration gas mixture showed a NO$_2$ concentration of 88.8 ± 0.4 ppb and an NH$_3$ concentration of 69.7 ± 0.4 ppb (mean values ± standard deviation after 1 and 3 days of tube conditioning). The TRANC-CLD system found 154.0 ± 0.9 ppb corresponding to a recovery rate of 97%. The uncertainty of the TRANC conversion efficiency not only depends on the precision of the individual measurements (indicated by the standard deviations) but also on the absolute accuracy of the independent measurement systems. The latter is limited by the absolute uncertainty of the calibration gas source and dilution leading to total absolute (2σ) errors in the order of 10% (see error bars in Fig. 5).
3.1.3 Aerosol particle conversion tests

Aerosol particle conversion efficiency of the TRANC was tested in cooperation with the Institute for Tropospheric Research (IfT), Leipzig, Germany. Aerosol particles were generated using a collision-type atomizer (TSI, St. Paul, USA) with a 0.3 mm nozzle from aqueous solutions of either NaNO₃, NH₄NO₃, or (NH₄)₂SO₄ (0.5 g l⁻¹, 1 g l⁻¹, 0.5 g l⁻¹, respectively), the three most common nitrogen containing aerosol compounds (e.g. Wexler and Seinfeld, 1991; Nemitz et al., 2009). The aerosol particles were dried and flushed into a stainless steel chamber of 1 m³, which was kept at ambient pressure. From this chamber, samples were taken in parallel by the TRANC-CLD system and a twin differential mobility particle sizer (TDMPS, Birmili et al., 1999), consisting of a differential mobility analyser combined with a condensational particle counter (UCPC3025 and CPC3010, TSI, St. Paul, USA). The recorded particle size and number distribution were used in combination with the known salt solid state densities to derive salt mixing ratios and consequently particle-bound Nr mixing ratios in the chamber air. The latter were compared to the \( \sum N_r \) measured by the TRANC-CLD system (Fig. 6). The results indicate mean apparent conversion efficiencies of 78 %, 142 % and 91 %, respectively. While the results show reasonably quantitative conversion of NaNO₃ and (NH₄)₂SO₄, NH₄NO₃ seems to be either overestimated by the TRANC-CLD or underestimated by the TDMPS. In contrast to the other two aerosol species, NH₄NO₃ is semi-volatile under ambient conditions, forming a thermodynamic equilibrium with gaseous NH₃ and HNO₃ (Stelson et al., 1979; Mozurkewich, 1993; Seinfeld and Pandis, 2006). For normal chamber air conditions (temperature of about 20 °C and relative humidity of about 50 %), the gaseous NH₃ and HNO₃ fraction may be significant. While the TRANC-CLD system would detect both, the gas and the aerosol phase, the evaporation of gaseous NH₃ and HNO₃ diminish the particle mass, measured by the TDMPS. Further differences between TDMPS and TRANC-CLD system may be caused by the use of ambient air to flush the chamber and to transport the generated aerosol. This led to a relatively high and variable Nr background (20 to 39 ppb) in the TRANC-CLD measurements, for which only an approximate correction was possible.

3.2 Field test with ambient N_r composition

In summer 2006, an intensive field campaign within the NitroEurope project allowed an in-field comparison between the concentrations of a variety of N_r compounds and the \( \sum N_r \) concentration measured by the TRANC-CLD system. The study took place in Central Switzerland, close to the village of Oensingen at an intensively managed grassland site. A detailed site description can be found in Ammann et al. (2009).

During this study NO and NO₂ were measured by a TEI chemiluminescence analyser in combination with a BLC

Fig. 8. Comparison between ΣN_r concentrations detected by the TRANC-CLD system and as the sum of measured individual species (acc. to Table 1) at the Oensingen field site during August 2006. Only those cases were included, for which measurements of all species (30 min means) were available. The solid line indicates the calculated linear regression (with corresponding equation), and the dashed line indicates the 1:1 line.

Photolytic converter (see Sect. 3.1.2), and gaseous NH_3, HNO_3, and HONO as well as particulate NH_4^+ and NO_3^- were measured using the GGradient of Aerosol and Gases Online Registrator (GRAEGOR, Thomas et al., 2009) – a wet chemical instrument, sampling with a wet annular denuder – steam jet aerosol collector combination, coupled to online analysis by ion-chromatography (HNO_3, HONO, NO_3^-) and flow injection analysis (NH_3, NH_4^+). Furthermore, NH_3 was measured additionally by a wet chemical AirRRmonia instrument (Erisman et al., 2001; Spirig et al., 2010; Flechard et al., 2010) (Table 1). Time series of NH_3 measured by the AirRRmonia system, gap-filled HNO_3, HONO, NH_4^+, NO_3^- data of the GRAEGOR, and NO, NO_2 data of the TEI are shown in Fig. 7. The exemplary time series reveal distinct fluctuations of the different N_r compounds as well as the large dominance of NH_3 and NO_2, accounting for more than 75 % of the site’s ambient ΣN_r concentration. The observed variability is related to the weather conditions with generally low concentrations in the rainy periods and also to the diurnal cycle with e.g. peak NO concentrations in the morning hours (due to the traffic source, limited vertical mixing and low ozone concentrations). The TRANC-CLD results compare very well to the sum of the individual measurements, capturing its fluctuations independently of the contribution of the different N_r species. A quantitative comparison between the two approaches for the ΣN_r concentration is shown in Fig. 8. The data points are generally close to the 1:1 line indicating an overall good agreement with slightly higher values by TRANC-CLD method (linear regression slope of 1.07 ± 0.013). The small systematic deviation can either result from missing N_r compounds in the individual measurements (organic nitrogenous compounds like e.g. PAN, MPAN, acetonitrile etc.) or combined calibration errors for the various analyser systems. Yet, there is no indication of a significant lack of conversion efficiency in the TRANC system under the varying field conditions.

3.3 Time response

An important feature for a converter designed for EC measurements is the fast detection of rapidly changing gas concentrations. Thus, we checked the TRANC-CLD system for time response characteristics during the calibration periods (Sect. 3.1.1) to make sure that even fast variations are clearly determined by the CLD without major attenuation effects. Figure 9a shows the time response of the NO signal in ppb recorded at a rate of 20 Hz after switching the TRANC valve from the calibration mode back to ambient air, which in that case was about 5 ppb. The course of the curve can be best approximated by exponential decay functions in the form of c(t) = Δc · e^{-t/τ} + c_{final} with a response (e-folding) time of τ = 0.30 s. The full step change was completed within about 1 s after switching the valve. Figure 9b and c show the time response of the NO signal recorded after switching the TRANC valve from the NO_2 and the NO plus O_3 calibration mode, respectively, back to ambient air. Here, τ values of 0.30 s were found as well. These numbers demonstrate the fast conversion and time response of the TRANC-CLD system.
Table 1. Overview of the instruments used for concentration measurements of nitrogenous compounds during the field inter-comparison in Oensingen, August 2006.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Oxidation level</th>
<th>Instruments</th>
<th>Detection principle</th>
<th>Observed concentration range</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO</td>
<td>+2</td>
<td>ThermoElectron 42C</td>
<td>Chemiluminescence</td>
<td>0–28 ppb</td>
</tr>
<tr>
<td>NO\textsubscript{2}</td>
<td>+4</td>
<td>ThermoElectron 42C + blue light converter</td>
<td>Photolytic conversion + chemiluminescence</td>
<td>1–22 ppb</td>
</tr>
<tr>
<td>NH\textsubscript{3}</td>
<td>−3</td>
<td>AirRmonia</td>
<td>Membrane scrubber + conductivity cell</td>
<td>1–19 ppb</td>
</tr>
<tr>
<td>pNH\textsubscript{4}</td>
<td>−3</td>
<td>GRAEGOR</td>
<td>Mist chamber + flow injection/conductivity</td>
<td>0–1.9 ppb</td>
</tr>
<tr>
<td>HNO\textsubscript{3}</td>
<td>+5</td>
<td>GRAEGOR</td>
<td>Rotating denuder + ion chromatography</td>
<td>0–0.8 ppb</td>
</tr>
<tr>
<td>pNO\textsubscript{3}</td>
<td>+5</td>
<td>GRAEGOR</td>
<td>Rotating denuder + ion chromatography</td>
<td>0–1.7 ppb</td>
</tr>
<tr>
<td>HONO</td>
<td>+3</td>
<td>GRAEGOR</td>
<td>Rotating denuder + ion chromatography</td>
<td>0–0.8 ppb</td>
</tr>
<tr>
<td>$\sum N_r$</td>
<td>−3 to +5</td>
<td>TRANC-CLD</td>
<td>Thermal and catalytic conversion + chemiluminescence</td>
<td>0–69 ppb</td>
</tr>
</tbody>
</table>

system. In another experiment (data not shown), two different NO concentrations were prepared and alternately fed into the system, one by the multi-gas calibrator and the other directly from the gas cylinder. In this test, we found e-folding times of 0.35 s. Given the fact, that the average residence time of air in the CLD reaction chamber ($V = 0.71$ l at a pressure of 20 mbar and a flowrate of 3 sl min\(^{-1}\)) is 0.28 s, the reaction time is largely limited by the flushing of the CLD reaction chamber with probably some additional longitudinal mixing due to the flow velocity profile within the tube.

4 Discussion and conclusions

High-frequency measurements of $N_r$ species have been rarely conducted in land-atmosphere exchange research and were usually limited to selected sites and short field campaigns. Neither a robust measurement technique for the long-term monitoring of $\sum N_r$ nor a technique that is fast enough to be applicable for EC measurements has been engineered and field-tested up to now. With the development of the TRANC, we present a unique methodology for both quantifying $\sum N_r$ and a device being robust and fast enough to be usable in an EC setup when combined with a fast response NO detector (CLD).

The specific design of the TRANC which features fast and strong heating of the sample gas and quick conversion of $N_r$ compounds to NO allows the detection of polar, water soluble and sticky substances like NH\textsubscript{3}, avoiding e.g. water surface effects (Kita et al., 2006; Sintermann et al., 2011). As the NO resulting from the conversion is relatively inert (in the absence of NO\textsubscript{2} and O\textsubscript{3}) it permits the use of long tubing between the TRANC and the analyser system and thus an installation of the TRANC on masts and towers for micrometeorological measurements. Other arising problems when using long tubing such as the correct determination of the lag time (to maximise the covariance), i.e. the time the air sample needs from entering the converter until it reaches the analyser, or possible high-frequency damping will be addressed in Ammann et al. (2012).

Testing recovery rates for single $N_r$ compounds during laboratory tests, we found values of 95 and 99 % for NH\textsubscript{3} and NO\textsubscript{2}, respectively, and 97 % when the two gases were combined. In-field long-term stability was approved by repeated conversion tests for NO\textsubscript{2} with an average value of 91 %. These conversion efficiency results were not significantly different from 100 %. For particulate mineral N, the laboratory conversion tests showed a less clear picture with more uncertain conversion efficiencies. However, while too many unknowns limit the possibility to explain all differences in the measurements, the overall qualitative picture (cf. Fig. 6) indicates an effective conversion of the aerosol particle N to NO in the TRANC.

The conversion efficiencies of the TRANC are comparable to values observed in other studies. McCalley and Sparks (2009), for example, report on 86 % for NH\textsubscript{3} measured with their converter system and 100 % for NO\textsubscript{2}. Horii et al. (2004) compared concentrations of NO\textsubscript{2} obtained by TDLAS versus photolysis-chemiluminescence instruments and found a slope of 1.1 ± 0.2 with a $R^2$ of 0.91. The TD-LIF instrument presented by Farmer et al. (2006) was tested by...
Thornton et al. (2003) and Rosen et al. (2004) with a conversion efficiency of >95% for NO2 and >94% for \( \sum PN \) using photolysis-chemiluminescence and gas chromatographs equipped with electron capture detectors, respectively, for comparative measurements. McCalley and Sparks also found a 100% transformation for HONO, however only 78 to 99% for NOy (PAN, alkyl nitrate, HNO3 and other forms of NOy).

Kliner et al. (1997) also used a Au catalyst to convert NOx compounds to NO and discovered a gas-phase process that contributes to a more readily conversion of HNO3 than e.g. NO2. Furthermore, they investigated some non-NOx species (reduced N2 species) such as hydrogen cyanide (HCN) and acetonitrile (CH3CN) for potential interferences in the NOy detection scheme. They found an almost full conversion of HCN and a partial conversion (\( \sim 40 \) to 65%) of CH3CN at 300°C. In the TRANC with the additional 870°C unit we expect a full conversion of all organic N2 compounds. In Kliner et al. (1997) and also in a similar study by Fahey et al. (1985), N2O conversion was found to be negligible. In our tests, we performed an in-field comparison between the sum of individually measured NO, NO2, NH3, HNO3, HONO, NH4+, NO3- by commercially available systems and the \( \sum N_r \) concentration by TRANC-CLD. Excellent agreement was found in the temporal course \( (R^2 = 0.96) \) with the TRANC showing on average slightly (7%) higher values. This small systematic difference might be due to minor N2 compounds not measured individually (e.g. organic compounds like PAN) or small errors in the calibration of any of the used analysers. The concentrations of HONO and HNO3, for which we did not explicitly check the conversion efficiency, were very small during the field experiment and thus their contribution to the \( \sum N_r \) measurements by the TRANC was not statistically significant. However, based on previous reports in the literature (e.g. Fahey et al., 1985; McCalley and Sparks, 2009) we assume that they were nearly fully converted under the conditions in the TRANC.

Regarding the response time for step concentration changes, we determined e-folding times of \( \leq 0.35 \) s during different laboratory and field tests. As this value is in large part influenced by the flushing of the CLD chamber, it demonstrates that the TRANC is able to quickly and accurately detect the variability in \( \sum N_r \) concentrations. For a similar system with a lower flow rate that was used for NOy deposition measurements over a 24-m high forest Munger et al. (1996) found an e-folding time of less than 1 s.

The usage of the TRANC-CLD system within an EC setup allows for the determination of the net ecosystem exchange of \( \sum N_r \) continuously over longer periods, thereby providing a parameter of major interest when assessing the productivity of ecosystems by its nutrient N input. Also, there is no need to correct for chemical interactions and phase changes of the N2 compounds, keeping the measurements at a relatively low maintenance and reasonable cost level. However, one of the major benefits of the TRANC is at the same time one of its biggest constraints, i.e. the fact, that it can only measure \( \sum N_r \) (or net N exchange) and a differentiation into single N2 compounds is not possible with one converter system. But even for process studies and mechanistic modeling, the \( \sum N_r \) measurement can provide important constraints and validation data in combination with additional selective converters or detectors.

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References


