

Dear referee,

Thanks for the constructive comments.

We address the reviewer's comments here below, in italics.

We have also updated the latest version of the manuscript of 10th December 2018 to address the reviewer's comments.

This review is focused specifically on the experimental design and interpretation of the NH₃ flux measurements during this study. It is well-known that NH₃ is susceptible to adsorption and desorption processes that are temperature and humidity-dependent and the authors acknowledge this issue throughout the manuscript. Several lines of evidence are provided to evaluate the potential for artefacts in the flux measurements. Given how rare direct measurements of NH₃ flux are, and how challenging they can be to obtain, I support the publication of the data in this manuscript, however I have some suggestions for the authors to slightly reframe their discussion of the results.

On lines 175 – 180, the authors state that calibration linearity at lower concentrations is assured by a 4-point linear calibration between 30 and 200 ppb, and the response to a 'zero air calibration'. However, several studies (e.g. Whitehead et al., 2008, Ellis et al., 2010) have reported that biases related to adsorption and desorption become relatively more important at lower concentrations, so one cannot simply interpret linearity and high concentrations and a clean background as evidence that there are no issues with quantification at lower concentrations. Given that the ambient mixing ratios are mostly below 10 ppb, the authors should be more cautious in their interpretation.

Authors response: Line 177, the following text has been added:

"The linearity of the response for low concentrations is tested by the response to zero air calibration, giving a $R^2=0.997$. However, at low mixing ratios (typically less than 100 ppb), a non linear increase of the interactions of NH₃ with the surface used in the inlet design has to be considered (Ellis et al., 2010, Whitehead et al., 2008). Therefore, an uncertainty in the quantification of low NH₃ concentrations has to be taken into account due to surface interactions. The global precision of the analyzer is ± 0.4 ppb according to the manufacturer's specification for a 0-500 ppbv range."

In Appendix B, the authors present a set of laboratory experiments to evaluate the potential impact of adsorption or desorption from the chamber walls under a narrow range of temperature and RH. I believe that the concentration differences between the through chamber (TC) and direct (D) measurements seen here would be a better way to estimate of the detection limit for fluxes than the 0.4 ppb instrument precision limit that the authors are currently using. For example, one could say that an absolute difference of 1.6 ppb or a relative difference of 12% can arise simply from the effects of adsorption or desorption. In this case, ambient flux measurements that generate such small differences in NH₃ should be viewed as not statistically different than zero.

Authors response: An error has been found in Table B1, line 3 column 4, [NH₃]D is 8.0 instead of 8.8.

According to Table B1 and Appendix B, an absolute difference of maximum 1.6 ppb or a relative difference of maximum 12% can be due to interactions of NH₃ with the surface. Instead of considering these maximum values, we calculated the average difference in concentration between TC and D measurements, which is 0.9 ppb.

Line 672, the following text has been added: “The average difference in concentration is 0.9 ppb, and should be considered as the detection limit for fluxes significantly different from zero (i.e. including potential effects of adsorption or desorption). 30% of the concentration differences are below 0.9 ppbv.”

Line 221, the following text has been added:” According to Appendix B, if the difference in NH₃ concentration used to calculate a flux is inferior to 0.9 ppb, the resulting flux may not be distinguished from a potential effect of adsorption or desorption onto the chamber walls. The precision of the analyzing device (analyzer + chamber + tube) may be defined at 0.9 ppb (corresponding to a flux of 0.55 ngN.m⁻².s⁻¹). In that case, low NH₃ fluxes comprised between 0.4 and 0.55 ngN.m⁻².s⁻¹ are considered as close to zero but are kept in the average daily flux calculation.”

Line 266, the following text was added: “. Among the 351 NH₃ valid fluxes, 30% are derived from a concentration difference of less than 0.9 ppb.”

A consequence of this more conservative approach will be that the number of NH₃ flux measurements above the threshold for quantification will decrease. The authors expand on this idea in the discussion in Appendix C, indicating that 23% of the fluxes measured were below a value of 0.5 ng N m⁻² s⁻¹. The authors go on to speculate about the consequence of excluding these low flux values from the database – the resulting average flux value is larger. I think that the low values should not be excluded – they contain important information about periods where the fluxes are small. Rather the authors should use the results from the experiments in Appendices B and C to derive a more realistic flux detection limit for NH₃ and then clearly report that for positive or negative fluxes of smaller magnitude, the flux is statistically indistinguishable from zero. These near-zero values are still meaningful – for example they can be compared to the predictions from the compensation point framework described in Section 2.7, or compared to the much larger fluxes measured over fertilized maize fields in other parts of the world.

Author response: The definition of a new detection limit has already been discussed in the previous response. We have modified Appendix C and added the following sentence:

Line 697: “Positive or negative fluxes inferior to 0.55 ngN.m⁻².s⁻¹ (corresponding to concentration differences less than 0.9 ppb in the flux calculation, as defined in Appendix B) represent 30% of the 350 measured fluxes”.

In the results and discussion part, the fluxes are daily averaged. It is therefore rather difficult to compare near zero individual fluxes with daily averages. To include their existence in the discussion part, the following sentence was added:

Line 494: “ As discussed in Appendices B and C, 30% of individual fluxes used to calculate the daily averages are very low and not distinguishable from adsorption or desorption of NH₃ on chamber walls. These very low fluxes are however meaningful and indicate that some periods of near zero measurements must be taken into account to represent the processes of exchanges in these ecosystems.”

Additional specific comment: The statement on lines 53-55 implies that application of synthetic fertilizer is the largest source of ammonia from agriculture whereas it is actually animal husbandry. The sentence should be modified accordingly.

Author response: Line 53, the sentence was changed into: “The largest sources of NH₃ emissions are animal husbandry, and agriculture via the application of synthetic fertilizer.”

Whitehead, J.D., M. Twigg, D. Famulari, E. Nemitz, M.A. Sutton, M.W. Gallagher, D. Fowler (2008) Evaluation of laser absorption spectroscopic techniques for eddy covariance flux measurements of ammonia, *Environmental Science and Technology*, 42, 2041-2046.

Ellis, R. A., Murphy, J. G., Pattey, E., van Haarlem, R., O'Brien, J.M., and Herndon, S. C. (2010) Characterizing a Quantum Cascade Tunable Infrared Laser Differential Absorption Spectrometer (QC-TILDAS) for measurements of atmospheric ammonia, *Atmos. Meas. Tech.*, 3, 397–406.

Measurements of nitric oxide and ammonia soil fluxes from a wet savanna ecosystem site in West Africa during the DACCIWA field campaign.

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Abstract.

Biogenic fluxes from soil at a local and regional scale are crucial to study air pollution and climate. Here we present field
measurements of soil fluxes of nitric oxide (NO) and ammonia (NH₃) observed over four different land cover types, i.e. bare
soil, grassland, maize field and forest, at an inland rural site in Benin, West Africa, during the DACCIWA field campaign in
20 June and July 2016. At the regional scale, urbanization and a massive growth in population in West Africa has been causing a
strong increase in anthropogenic emissions. Anthropogenic pollutants are transported inland and northward from the mega
cities located on the coast, where the reaction with biogenic emissions may lead to enhanced ozone production outside urban
areas, as well as secondary organic aerosol formation, with detrimental effects on humans, animals, natural vegetation and
crops. We observe NO fluxes up to 48.05 ngN m⁻² s⁻¹. NO fluxes averaged over all land cover types are 4.79 ± 5.59 ngN m⁻²
25 s⁻¹, maximum soil emissions of NO are recorded over bare soil. NH₃ is dominated by deposition for all land cover types. NH₃
fluxes range between -6.59 and 4.96 ngN m⁻² s⁻¹. NH₃ fluxes averaged over all land cover types are -0.91 ± 1.27 ngN m⁻² s⁻¹
and maximum NH₃ deposition is measured over bare soil. The observations show high spatial variability even for the same
soil type, same day and same meteorological conditions. We compare point daytime average measurements of NO emissions
recorded during the field campaign with those simulated by GEOS-Chem (Goddard Earth Observing System Chemistry
30 Model) for the same site and find good agreement. In an attempt to quantify NO emissions at the regional and national scale,
we also provide a tentative estimate of total NO emissions for the entire country of Benin for the month of July using two
distinct methods: upscaling point measurements and using the GEOS-Chem model. The two methods give similar results:

1.17 ± 0.6 GgN/month and 1.44 GgN/month, respectively. Total NH_3 deposition estimated by upscaling point measurements for the month of July is 0.21 GgN/month.

35 **1 Introduction**

Biogenic soil fluxes of nitric oxide (NO) and ammonia (NH₃) play an important role on tropospheric chemistry. Nitric oxide emitted by soil influences the concentration of nitrogen oxides (NO_x) in the atmosphere, consequently modifying the rates of ozone (O₃) production, where O₃ is a pollutant, harmful to humans and plants, and also a greenhouse gas (Steinkamp et al., 2009). The production and consumption of NO in soil is regulated by microbial activity, mainly nitrification/denitrification processes, and chemical reactions (Pilegaard et al., 2013). Measurements using soil chambers in the field and laboratory experiments show that nitrification/denitrification, and consequently NO emissions, vary greatly with climate and soil conditions, in particular they are strongly correlated with nitrogen (N) availability, temperature and soil moisture, making soil NO emissions dependent on regional temperature and precipitation patterns, and fertilizer management practices (e.g., Bouwman et al., 2002; Meixner and Yang, 2006; Hudman et al., 2010).

45 Soil NO emissions are about 20% of total NO sources to the atmosphere (IPCC, 2007) and almost of the same order of magnitude of fossil fuel NO emissions. Soil emission of biogenic NO plays a prominent role in the regional atmospheric chemistry of non-urbanized areas, where anthropogenic emissions are negligible (Pilegaard, 2013). The main inputs of N compounds onto semi arid uncultivated soils, like the savanna ecosystem, are biological nitrogen fixation, atmospheric wet and dry deposition and lightning. NO fluxes are considered as one way only, even if NO deposition exists in very specific conditions (Grote et al., 2009).

55 Soil N losses towards the atmosphere also involve NH₃. The largest sources of NH₃ emissions are animal husbandry, and agriculture, via the application of synthetic fertilizer. When released into the atmosphere, NH₃ increases the level of air pollution. In the atmosphere NH₃ has a relatively short life time of less than five days and high deposition rates, it is converted into ammonium (NH₄⁺) aerosols, which has a life time of the order of fifteen days, can travel long distances and it is relevant for air quality and climate (Fuzzi et al., 2015). The exchange of soil NH₃ is bi-directional as it also includes deposition. NH₃ returned to the surface by deposition can potentially cause eutrophication, reducing biodiversity and water quality (Sutton et al., 2009a).

60 The net flux of NH₃ is the combination of different exchange pathways between plant (cuticle and stomata), soil, leaf litter and atmosphere. The overall NH₃ flux for a given surface may switch from net emission to net deposition at sub-hourly, diurnal and seasonal scales. Moreover, NH₃ can be rapidly deposited onto cuticles due to its high solubility (e.g. Sutton et al., 2009b; Massad et al., 2010; Loubet et al., 2012).

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The direction and magnitude of NH_3 exchanges depend on the difference in NH_3 concentration between the canopy and the atmosphere, and on a large range of environmental factors, in particular air humidity, which influence surface wetness, and soil moisture conditions, but also vegetation cover and soil characteristics. The relationships between NO and NH_3 soil fluxes have been identified through the ammonium content in the soil (McCalley and Sparks, 2008). Ammonia is mainly
70 emitted by agricultural activities, and also by the decomposition of litter and volatilization of animal excreta (Sutton et al., 2009b; Massad et al., 2010).

Soil fluxes in West Africa have only been measured in a limited number of studies due to the challenging experimental conditions (remote sites, no power supply, very hot temperatures), and mainly with manual chamber techniques rather than
75 more complex micrometeorological techniques (Serça et al., 1998, Le Roux et al. 1995 for NO, Delon et al., 2017 for NO and NH_3). However, tropical savanna has been recognized as one of the ecosystems characterized by the largest NO emissions (Davidson and Kinglerlee, 1997, Hudman et al., 2012).

Anthropogenic emissions of pollutants from mega cities located on the Guinean coast in South West Africa have been
80 increasing, and are likely to keep increasing in the next decades, due to a strong anthropogenic pressure, land use change and urbanization. When transported northward on the African continent, polluted air masses meet biogenic emissions from rural areas which contributes to increased O_3 and secondary organic aerosol production, in high temperature and solar radiation conditions, highly favorable to enhance photochemistry (Knippertz et al., 2015a, 2015b).

The objectives of this study are to quantify soil fluxes of NO and NH_3 for the different land cover types typical of rural West Africa, suggest a tentative strategy to scale point measurements in the field to ecosystem and larger regional scale, and
85 provide data for inventories and model evaluation to improve air quality and climate modelling.

In this paper we present the soil fluxes of NO and NH_3 measured in a rural site near the city of Savè, Benin, West Africa,
90 during the DACCIWA (Dynamics-Aerosol-Chemistry-Cloud Interactions in West Africa) field campaign which lasted from 14th June to 30th July 2016 (wet season). The DACCIWA campaign was lead to investigate the possible role of local air pollution on climate change in West Africa, focusing on atmospheric composition, air pollution and cloud-aerosol interactions over several sites in the region (Knippertz et al., 2015a, 2015b, 2017). The Savè site is part of the savanna ecosystem, where grassland is intercut with crops and degraded forest. Biogenic soil fluxes measurements were taken using
95 the manual chamber technique, which is robust and of reduced costs (Delon et al., 2017). Along with these observations we also present measurements of soil characteristics and meteorological variables from the same site. We include the comparison of measured NO soil emissions with those simulated by the Hudman et al. (2012) process-based model for NO soil emission implemented into GEOS-Chem.

2 Material and method

100 2.1 Site description

The Savè site for ground-based observations is located in a hinterland area of Benin, 6 km south west from the city of Savè (8°02'03" N, 2°29'11" E, 166 m a.s.l.). The Savè ground-based observation site is located within the Gobè site managed by the Institut National des Recherches Agricoles du Bénin (INRAB).

105 The site is characterized by a wet savanna ecosystem. The climate of the region is Sudano-Guinean, with a rainy season from March to October and a dry season from November to February (Michels et al., 2000). The average annual rainfall is about 1100 mm (Savè weather station, data averaged from 1969 to 2004, Michels et al., 2000 and Säidou et al., 2004) and the average yearly temperature is about 27.5 °C with little variation from year to year (data averaged from 1984 to 2004, Säidou et al., 2004). Average minimum temperature, based on 1969-1990 data, is 21.5 °C and mean maximum temperature is
110 35.5 °C.

The tree coverage in the Savè region is low with most of the land occupied by subsistence agriculture and grassland (CILSS, 2016). Four land cover types are identified at the observation site: bare soil, grassland, maize field and degraded forest. Bare soil is defined as a patch of land of minimum five by five meters wide, without vegetation growing or hanging over the plot.

115 Ground photographs of the four land cover types are shown in Fig.1.

The most abundant tree species next to the grassland site and in the forest are: *Anacardium occidentale*, *Daniellia oliveri*, and *Pterocarpus erinaceus*; while the most abundant tree species next to the maize field are: *Mangifera indica*, *Cocos nucifera*, *Carica papaya L.*, *Tectona grandis*, and *Azadirachta indica*. The herbaceous vegetation is dominated by *Cleome*
120 sp., *Crotalaria* sp., *Mucuna* sp., *Imperata cylindrica* and *Rhynchelytrum repens* next to the grassland site and in the forest, and *Commelina benghalensis*, *Euphorbia* sp., *Boerhavia diffusa*, *Phyllanthus amarus*, *Digitaria horizontalis* by maize field. In the maize field, the main species, *Zea Mays*, is intercropped with *Sesamum indicum* and, to a lesser extent, with other species: *Dioscorea* sp., *Manihot esculenta*, *Arachis hypogaea*, *Vigna unguiculata*, *Gossypium* sp., *Sorghum* sp. and *Solanum lycopersicum*. The maize field was not treated with mineral fertilizer. The only livestock consist of a few dozens of domestic
125 fowls belonging to small subsistence-oriented family farms, mainly grazing in the maize field.

At the Savè site, the soil is sandy, with 87% of sand and 4.1% of clay (the rest being silt) for the 0-5 cm horizon. Surface pH ranges from 6.32 to 8.46, depending on the place where the measurement is done. Mean meteorological and average soil characteristics for the observation site are reported in Table 1, dominant vegetation species and soil composition for each
130 land cover type are given in Table 2 and 3, respectively. Sunrise and sunset UTC time at the beginning and at the end of the campaign are: 05:33 and 18:08 on 14th June, and 05:42 and 18:11 on 30th July.

2.2 Sampling sites

The samples were taken at the four land cover types (bare soil, grassland, maize field and forest), one location per day. Two to three samplings spots were chosen each day for each location, collecting eight to twenty-five flux measurements for both NO and NH₃ soil fluxes each day. Each location was sampled during daytime, approximately from 7 a.m. to 6 p.m., alternating measurements at the four different land cover types from one day to the other, over the entire campaign. Bare soil and the maize field were sampled for both NO and NH₃ soil fluxes on eight different, generally non-consecutive, days, grassland on ten days, and the forest site on four different days.

2.3 Chamber flux measurements

The technique used to measure NO and NH₃ soil fluxes makes use of a ThermoScientific 17i (ThermoFischer Scientific, MA, USA). This analyzer uses a chemiluminescence detector for NO. The air sample enters the reaction chamber and reacts with the O₃ generated by an internal generator. This reaction produces a luminescent radiation directly proportional to the NO concentration. The air sample is sequentially drawn through a molybdenum converter heated to 325°C which measures NO_x (NO+NO₂) by converting NO₂ to NO, and a stainless steel converter heated to 750°C which measures N_{total} (NH₃+NO_x) by converting NH₃ and NO₂ to NO. The detector hence measures rNO, then r(NO + αNO₂), and finally r(NO + βNO₂ + γNH₃), where r is the NO detection efficiency, α and β are the NO₂ conversion efficiency of the molybdenum and stainless steel converters and γ is the NH₃ conversion efficiency of the stainless steel converter. The efficiencies are determined by the calibration procedure. NH₃ concentration is therefore calculated from N_{total} – NO_x. The closed dynamic chamber technique is used to calculate fluxes. The details of this technique are fully described in Delon et al. (2017).

The remoteness of the study site limited installation of permanent structures and we were unable to automate our chamber measurements, thus all measurements were made manually. The instrument was powered by a generator (>100 m away) and carried around on a wheeled-table to reach the locations of the four soil types where the NO and NH₃ soil fluxes were measured. The analyzer was connected via a Teflon tube to the Teflon chamber which was put on the ground to detect the fluxes. The external sides of the chamber were covered with sand or soil to isolate it during the measurement. The soil under the chamber was left unperturbed. Adjustments were taken in order to make sure the analyzer did not reach temperatures that would invalidate the measurements.

The calibration of the NO sensor of the 17i analyzer was made before and after the campaign, with a reference NO air mixture, i.e. NO in N₂ diluted with zero air. The NO detection efficiency variation was 8% between the two calibrations (from 1.040 to 0.962). Two post-campaign calibrations were made: a first one to validate the efficiency of the NO₂ converter using a reference dilution of NO₂ in zero air, and a second one to validate the efficiency of the NH₃ converter with a NH₃/N₂

165 mixture diluted in pure air (Alphagaz 1, Airliquide). No change in the NO₂ conversion efficiency was necessary, and the NH₃
conversion efficiency variation was 3% (from 0.963 to 0.995). No drift in the conversion efficiencies was observed over
time, as from the first calibration when the analyzer was new until the post campaign calibration, changes never exceeded
±3%. The zero air for NO, NO₂ calibration was obtained by filtering ambient air, previously passed on charcoal and
desiccant cartridges. The dilution for all the calibration experiments was made with the 146i module (ThermoFischer
Scientific, MA, USA) and the dilution module, equipped with certified mass flow meters, on board of the ATR-42 research
170 aircraft during an inter-calibration with other NO_x instrumentation of the DACCIWA campaign (i.e. the instrumentation on
the Savè measurement site tower and the instrumentation on the ATR-42 aircraft; Brito et al., 2017, Derrien et al., 2016).
Reference NO, and NO₂ were ISO 6141:2015 certified at 8.73 and 8.58 ppm for NO, before and after the campaign,
respectively, and 9.28 ppm for NO₂, both with 5% precision. Reference NH₃ mixture was certified at 14.78 ppm with 2%
precision for NH₃. Multipoint (at least 4 points) calibrations between 50 to 250 ppb were done to ensure the linearity of the
175 response, obtaining regression coefficients over 0.9993 for both NO and NO₂. The dilution uncertainty was 10% for NO,
11% for NO₂ and 13% for NH₃ (see Appendix A for more detail). A multipoint calibration was done for NH₃, between 30 and
200 ppb and the regression coefficient was 0.997. The linearity of the response for low concentrations is ~~ensured~~-tested by
the response to zero air calibration, ~~with giving~~ a R²=0.997. ~~However, at low mixing ratios (typically less than 100 ppb), a
non linear increase of the interactions of NH₃ with the surface used in the inlet design has to be considered (Ellis et al., 2010,
Whitehead et al., 2008). Therefore, an uncertainty in the quantification of low NH₃ concentrations has to be taken into
account due to surface interactions.~~ The global precision of the analyzer is ±0.4 ppb ~~(according to the manufacturer's
specification for a 0-500 ppbv range).~~

185 The external volume of the chamber was 40 cm×20 cm×20 cm. The internal volume was 18×38×18 cm³, due to the thickness
of the Teflon walls. The air inlet is located on one side of the chamber, where a small vent of 4 mm in diameter provided the
pressure equilibrium between the inside and outside of the chamber. The air outlet on the other side is connected to the
analyzer with a 4 m Teflon tube (see picture displayed in Appendix B) The chamber is continuously swept with an air flow of
0.7 L min⁻¹ insured by the instrument pump, and the air flow is controlled inside the analyzer by a flow meter. The air
residence time in the chamber is approximately 20 min (Volume/flow), and the chamber is maintained in place for 10 min.
190 The Teflon chamber was cleaned (with a dry clean paper cloth) at the beginning of each day of measurement, and during the
day when the deposition of sand could potentially interfere with the measurements. Laboratory tests using different papers
for cleaning are displayed in Appendix C. According to these results, no clear tendency for potential adsorption or desorption
of NH₃ arises but these tests may be useful to warn for a potential pollution inside the chamber due to cleaning which would
interfere with low fluxes.
195 The opaque walls minimize photochemical reactions inside the chamber, which are therefore considered as negligible. The
chamber is placed on the soil for 10 min. After 10 min, the chamber is turned over to let the analyzer be swept by ambient air
for 5 min, then the chamber is placed again on the soil to begin a new cycle.

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200 The calculation of the fluxes is based on the closed dynamic chamber technique, with the following assumptions: the concentration in the chamber is equal to the concentration leaving the chamber to the analyzer, and potential deposition onto the Teflon walls of the chamber is assessed but considered as negligible. Vaittinen et al. (2013, and references therein) have demonstrated that the adsorption of ammonia on Teflon is negligible; However, the high NH₃ mixing ratios and the controlled conditions in Vaittinen experiment do not correspond to our field conditions. Therefore, experimental tests with and without the Teflon chamber attached to the analyzer were made in ambient air to verify that deposition on the walls of the Teflon chamber is negligible. These tests have been made in conditions comparable to in situ measurements, i.e. temperature (25 to 29°C) and humidity (46 to 54%), as well as NH₃ concentrations (8 to 35 ppb) close to the ones encountered in the field. They show that the concentrations measured with and without the chamber are equivalent. The results of this experiment are reported in Appendix B. Moreover, the temperatures of the chamber Teflon walls and Teflon tube have been measured in direct sunlight and the difference with air temperature is small (<1°C). We therefore assume that the Teflon walls and tube heating is small and does not affect the NH₃ and NO concentration measurements in the chamber. The results of this experiment are reported in Appendix D. All the details of the calculation are given in Delon et al. (2017). In brief:

$$F_x = \frac{V}{A_0} \frac{\delta C_x}{\delta t} \quad (1)$$

215 Where F_X is the flux (NO or NH₃) in nmol m⁻² s⁻¹, δC_X is the concentration variation in the chamber in nmol m⁻³ during the temporal interval δt. A₀=0.0684 m² is the surface of the ground covered by the chamber, V=0.0123 m³ is the volume of the chamber. This equation is similar to the one in Davidson et al. (1991). The flux is then converted to ngN m⁻² s⁻¹.

220 The linear regression is calculated over a 100 to 300 s time interval after the installation of the chamber on soil for both NO and NH₃. The dilution effect due to mixing of outside air in the chamber was evaluated based on our set up in which Q/V=8.13×10⁻⁴ s⁻¹. It is calculated for each flux separately and is in average 6.7(±1.6) % for NO and 7.7(±1.7) % for NH₃. Considering the precision of the analyzer (±0.4 ppbv), the detection limit is 0.4 ngN m⁻² s⁻¹ for NO and NH₃ fluxes.

225 According to Appendix B, if the difference in NH₃ concentration used to calculate a flux is inferior to 0.9 ppb, the resulting flux may not be distinguished from a potential effect of adsorption or desorption onto the chamber walls. The precision of the analyzing device (analyzer + chamber + tube) may be defined at 0.9 ppb (corresponding to a flux of 0.55 ngN.m⁻².s⁻¹). In that case, low NH₃ fluxes comprised between 0.4 and 0.55 ngN.m⁻².s⁻¹ are considered as close to zero but are kept in the average daily flux calculation.

230 The chemical reactions inside the chamber can determine NO consumption, and consequently an underestimation of the NO fluxes calculated with our method. This underestimation is taken into account and calculated following the method by Pape

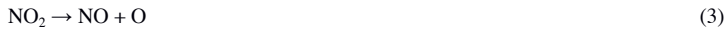
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et al. (2009) with the relation $k \cdot [\text{NO}] \cdot [\text{O}_3]$. In this relation k is the temperature-dependent reaction rate constant (Pape et al., 2009, Atkinson et al., 2004), $[\text{NO}]$ is measured by the ThermoScientific 17i at soil level just before positioning the chamber for the measurement of soil fluxes, and $[\text{O}_3]$ at soil level is derived by measurements of NO and NO_2 at soil level made with the ThermoScientific 17i and measurements of NO , NO_2 and O_3 taken on an 8 m high tower. On the 8 m high tower, NO and NO_2 were measured with a Model 42C TraceLevel $\text{NO-NO}_2\text{-NO}_x$ by Thermo-environmental Instruments Inc., calibrated with the same method as the ThermoScientific 17i and with 0.05 ppb (2-sigma) detection limit. Ozone was measured on the tower with a Model 49i Ozone Analyzer by Thermo-environmental Instruments Inc. with 1 ppb detection limit. The Model 49i Ozone Analyzer was calibrated by comparison with a Thermo Scientific Model 49PS reference instrument. The reference instrument is sent twice a year to the French Laboratoire national d'Essais (LNE) for comparison with a National Institute of Standards and technology (NIST). All data on the tower were sampled at 10 seconds. $[\text{O}_3]$ at soil level was then calculated considering the diurnal steady state of the reactions described in equation (2) and (3), using equation (4):



$$[\text{O}_3]_{sl} = \frac{[\text{NO}]_{tl} [\text{O}_3]_{tl} [\text{NO}_2]_{sl}}{[\text{NO}_2]_{tl} [\text{NO}]_{sl}} \quad (4)$$

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Where $[\]_{sl}$ is the concentration at the soil level and $[\]_{tl}$ is the concentration measured on the tower. In conclusion, we correct NO fluxes for the underestimation of NO fluxes due to chemical reactions inside the chamber with values ranging between 0 and 63% (8% on average for the whole campaign).

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As studied by Kristensen et al. (2010a) and Kristensen et al. (2010b), O_3 deposition can decrease O_3 concentration close to soil surface further. However, considering that O_3 concentrations calculated near the soil are already very low (1 ppb at soil level compared to 24 ppb at 8 m, averaged for the entire measurement campaign), O_3 deposition has been considered of secondary importance in this calculation and has not been included. If O_3 deposition were to be included it would possibly decrease the correction of NO fluxes and consequently slightly decrease NO emissions in a negligible proportion compared to the correction already applied for the chemical reactions inside the chamber.

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NH_3 measurements have not been corrected from a possible interaction with particulate matter (PM) as PM concentration (not measured at Savé) are supposed to be low because Savé is located in a rural area far from anthropogenic pollution influence. The walls of the Teflon chamber are cleaned daily to reduce any interference of NH_3 with PM deposition in the chamber. The effect of PM, even at low PM concentrations, may reduce the measurement accuracy and induce an uncertainty on the detection of the NH_3 flux from soil. This uncertainty has not been assessed quantitatively, but the reader must keep in mind that NH_3 fluxes may be estimated with less accuracy because of the presence of PM, especially for low fluxes.

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2.4 Data quality check

- 265 A quality check method based on the following criteria is used to select observed fluxes (Delon et al., 2017):
- The coefficient of determination for linear regression R^2 has to be higher than 0.4 (considered as a significant correlation) for NH_3 fluxes, and higher than 0.8 for NO fluxes. The variation of NH_3 is less stable than for NO because of potential interaction with PM in the chamber. However, 80% of the R^2 were superior to 0.6 for NH_3 , and 100% for NO. Examples of the variation in time of the concentration of NH_3 and NO in the chamber are shown in Appendix E for two different soils.
 - 270 - A flux error was estimated by calculating the dispersion of points around the linear regression's slope. According to this method, the dispersion for NO flux calculation is comprised between 5 and 12%, and the dispersion for NH_3 flux calculation is comprised between 15 and 20%.
 - The concentration difference between the last and the first NH_3 measurement point has to be more than 0.4 ppb (sensitivity of the analyzer). R^2 was generally lower than 0.4 for concentration differences below 0.4 ppb.
 - 275 Finally, 351/488 (72%) NH_3 flux measurements and 459/488 (94%) NO flux measurements are considered valid. Among the 351 NH_3 valid fluxes, 30% are derived from a concentration difference of less than 0.9 ppb.

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Uncertainty of the NH_3 flux calculation

- Despite all precautions to reduce adsorption on the chamber walls and/or interaction with PM in air, (see Appendix B, C D and E), R^2 are less good for NH_3 , due to potential chemical or physical interaction of material with NH_3 (whereas considered as negligible in this study). However, no absolute correction for adsorption can be calculated in field conditions. Teflon remains the more reliable material to measure NH_3 , as shown in Sauren et al. (1989) who find that Teflon has the lowest adsorption affinity for NH_3 (as compared with aluminium, parafine and gold), but a passivation time lag remains for NH_3 detection in measurements systems (Yokelson et al., 2003).
- 280

285 2.5 Meteorological station

- Continuous in situ observations of meteorological variables, including air and soil temperature and moisture, rainfall, wind speed, wind direction, radiation and energy balance components were taken at the Savè site as part of the DACCIWA campaign. Data are provided as 1 min averages, apart from energy fluxes which are given as 30 min averages (Derrien et al., 2016, Kohler et al., 2016, Handwerker et al., 2016, Wieser et al., 2016). An overview of the complete set of instrumentation and measurements is given by Brooks et al. (2017), while a summary of the available ground-based meteorological observations is given by Kalthoff et al. (2017). In this study we present soil moisture measured in two distinct locations of the Savè site by the Karlsruhe Institute of Technology (KIT) instrumentation at 5cm depth on grassland, and average soil moisture, between 0 and 30 cm, measured by the Université Paul Sabatier (UPS) instrumentation in the maize field. Details of the instrumentation is given by Brooks et al. (2017). We include soil moisture measured with both systems as the inter-
- 295 comparison of the two methods is out of the scope of this study.

2.6 Soil characteristics (texture, pH, N content)

Soil samples were collected with a cylinder of known volume (290 cm³) during the measurement campaign to analyze the biogeochemical characteristics of the site. Soil samples (0-5 cm) were taken for each land cover type where NO and NH₃ fluxes were measured. Fifteen samples were collected at the four different land cover types, three to four times during the campaign.

Samples were dried in ambient conditions (mean day-time temperature is approximately 26 °C, Kalthoff et al., 2017), and stored in the dark. After drying, the weight of the samples was measured to determine the bulk density (d_a = dry soil mass / total volume), which was found to be $1.24 \pm 0.14 \text{ g cm}^{-3}$. Assuming a density of soil particles (d_p) of 2.6 g cm^{-3} , the Water Filled Pore Space (WFPS) is calculated with the following equation 5:

$$\text{WFPS} = \text{SM} / (1 - d_a / d_p) \quad (5)$$

where SM is soil moisture in %

Soil samples were analyzed for the determination of texture, ammonium concentrations [NH₄⁺], C/N ratio, total C, total N, and pH at the GALYS Laboratoire (<http://www.galys-laboratoire.fr>, NF EN ISO/CEI 17025: 2005). The analyses were performed two months after sampling. We assume that the ammonium content in litter or soils is not modified by volatilization or chemical transformation during transport and storage, because of the very low soil moisture level in samples. Indeed, when collected, WFPS of the samples ranged between 6 and 14% (mean=8.5±3.5%) and soil temperature between 35 and 38 °C (data obtained from the databases described in Brooks et al., 2017). Bai et al. (2013, and references therein) have found that significant changes in nitrification and net mineralization (influencing the ammonium content) may occur when soil temperature raises until 35°C (the optimum for nitrification), for optimal soil moisture conditions (WFPS =20%, Oswald et al., 2013). In the present study, soil temperatures when sampling were equal or above the optimum, and WFPS was below the optimum, reducing the nitrification efficiency and the change in ammonium content. Several authors have published results of ammonium concentrations measured in soils dried in ambient air. For example, Dick et al., (2006) collected top soil after the wet season in two sites in Senegal. The authors state that their soils were considered dry when collected and were air-dried in the mid-day sun immediately after collection. The protocol used in our study is identical. Other studies (Bai et al., 2010, Cassity-Duffrey et al., 2015, Vanlauwe et al., 2002) also published ammonium measurements made on air dried soils from seasonally dry climates with comparable textures to the soil in Savé. Soil texture is determined following norm NF X 31e107. Clay (<2 µm), fine silt (2 to 20 µm), coarse silt (20 to 50 µm) and total sand (50 to 2000 µm) are determined without decarbonation. Organic carbon and total carbon are determined following norm NF ISO 10694. The whole carbon of the sample is transformed into CO₂. Then CO₂ is measured by thermal conductivity. NF ISO 13878 is used for Total N. Mineral nitrogen is determined following an internal method MT-AZM adapted from norm NF ISO 14256-2. This method uses a potassium chloride solution and is COFRAC certified. The sample is heated at 1000°C with O₂. Products

of combustion or decomposition are reduced in N_2 . N_2 is then measured by thermal conductivity (catharometer). pH is
330 determined according to norm NF ISO 10390, with soil samples stirred with water (ratio 1/5).

2.7 Soil ammonia emission potential Γ_g and compensation point χ_g

Measurements of soil pH and ammonium concentrations $[NH_4^+]$ are used to quantify the soil emission potentials for the
different land cover types at the measurement site. The soil emission potential Γ_g is the ratio of $[NH_4^+]$ to $[H^+]$ concentrations
in the water solution of the soil (mol L^{-1}). A large Γ_g indicates that the soil has a high propensity to emit NH_3 , considering
335 that the potential emission of NH_3 depends on the availability of ammonium in the soil and on pH.

The soil compensation point (χ_g) has been calculated from the emission potential Γ_g , as a function of soil surface temperature
(T_g in K) according to Wentworth et al. (2014):

$$340 \quad \chi_g \text{ (ppb)} = 13\,587 \cdot \Gamma_g \cdot e^{-(10\,396\text{K}/T_g)} \times 10^9, \quad (6)$$

The soil compensation point indicates the equilibrium between gaseous NH_3 in the soil pore space and $[NH_4^+]$ in the soil
solution, i.e. the concentration of NH_3 for which the NH_3 flux switches from emission to deposition (or vice versa).

2.8 Stepwise multiple regression analysis

345 A stepwise linear multiple regression analysis was performed between daytime averaged gas fluxes of NO and NH_3 and
relevant available daily averaged variables such as wind speed, soil temperature at 5 cm, soil moisture at 5 cm, soil heat flux,
outgoing longwave radiation and incoming shortwave radiation. Soil parameters such as mineral nitrogen, total N and
organic C, soil texture and pH could not be used for this regression analysis since their relative measurements did not have
the same temporal resolution as the other parameters. The R software (<http://www.R-project.org>) was used to provide the
350 results of this linear regression analysis.

2.9 GEOS-Chem

GEOS-Chem is a global three-dimensional model of tropospheric chemistry driven by meteorological input from the NASA
Goddard Earth Observing System (www.geos-chem.org, Bey et al., 2001). In this study we use GEOS-Chem Version 10-01
which includes the process-based parameterization of soil NO emission by Hudman et al. (2012). This parameterization
355 represents available nitrogen (N) in soils using biome specific emission factors, online wet and dry deposition of N, and
fertilizer and manure N derived from a spatially explicit dataset, distributed using seasonality derived from data obtained by
the Moderate Resolution Imaging Spectrometer (MODIS). Emissions are a smooth function of soil moisture and temperature
consistent with point measurements and ecosystem scale experiments. This parameterization also included pulsing following

soil wetting by rain or irrigation, represented as a function dependent on dry spell length. The parameterization by Hudman et al. (2012) was successfully evaluated for pulsing events in central Sahel (0–30° W, 12–18° N).

Boundary conditions for our experiment are generated from a global GEOS-Chem simulation at 4° x 5° horizontal resolution. The regional GEOS-Chem model for West Africa runs at a horizontal resolution of 0.25° x 0.3125° (latitudes 6°S–16°N, longitudes 18.125°W–26.875°E) and a vertical resolution of 47 levels (up to 0.01hPa). Meteorology is driven by the NASA GMAO (Global Modeling and Assimilation Office) GEOS-FP (Forward Processing) assimilated meteorological data. The global model is spun-up from 1st May 2015 to 1st May 2016. The global simulation is then run from 1st May 2016 to 1st August 2016, outputting boundary condition files for West Africa. The regional West Africa simulation is then run from 1st May 2016 to 1st August 2016 using the 4° x 5° boundary conditions from the global simulation. All simulations use the GEOS-FP meteorology which has a three-hour time resolution. We used the same MODIS/Koppen land cover map as in Hudman et al. (2012; <http://glcf.umd.edu/data/lc>) which includes 24 land cover types. In this simulation we use EDGAR v4.2 (EC-JRC/PBL, 2011) for anthropogenic emissions, GFED4 (Giglio et al., 2013) for biomass burning emissions and MEGAN v2.1 (Guenther et al., 2012) for biogenic emissions of volatile organic compounds. The same emission inventories are used for both the boundary conditions and the West Africa simulation.

3 Results and discussion

3.1 Meteorological data

Mean air temperature averaged over the whole campaign was 25.4 ± 2.6 °C, mean wind speed was 1.3 ± 0.6 m s⁻¹, mean relative air humidity is 86.3 ± 10.5 %, mean soil temperature was 25.2 ± 3.4 °C, mean KIT soil moisture at 5 cm was 7.1 ± 3.6 %, while mean UPS soil moisture averaged between 0 and 30 cm was 4.5 ± 2.8 %. Total KIT precipitation was 198 mm for the whole campaign, and total UPS precipitation was 215 mm.

Median diurnal cycles of air temperature, specific humidity and precipitation are reported in Kalthoff et al., (2017). Knippertz et al. (2017) distinguish four different phases of the monsoon season during the DACCIWA campaign (14th June to 30th July 2016) over the DACCIWA focus region (5–10° N, 8° W - 8° E), which covers a wide area of West Africa (see Fig. 1, Knippertz et al. (2017)). The division into phases is mainly based on the north–south precipitation difference between the coastal zone (0–7.5° N) and the Sudanian–Sahelian zone (7.5–15° N), both averaged across the longitude range 8° W–8° E. Savè (8.03° N) is located very close to the border between the two zones, with a rainfall pattern that seems to follow more closely that of the coastal zone rather than that of the northern inland Sudanian–Sahelian zone. These four phases are: the pre-onset phase characterized by a rainfall maximum near the coast (before 21st June, phase 1); the post-onset phase during which the rainfall maximum occurred inland (22nd June - 20th July, phase 2); the wet westerly regime when the rainfall maximum shifted back to the coast (21st – 26th July, phase 3); and the recovery of the monsoon with a shift of the rainfall

maximum inland (27th July until the end of the campaign, phase 4). A specific period within phase 2 is indicated “vortex”, during which an unusual development occurred (09th – 16th July): in the north, a cyclonic feature slowly propagated from eastern Mali to Cape Verde and in the south, an anticyclonic vortex tracked in the west-northwesterly direction along the Guinean coast (see Knippertz et al. 2017 for a more detailed description). At the Savè site the most intense rainfall events happened the day before the first soil fluxes observation, on 15th June 2016, and towards the end of the measurement campaign between 20th and 23rd July 2016. Other minor rainfall events are recorded on 19th and 27th June, 8th, 12th, 13th, 24th and 26th July. Daily rainfall measurements are reported in Figs. 2 to 5.

3.2 Soil texture, soil organic carbon, total nitrogen, pH and ammonium content

Bare soil recorded the lower amount of total sand (83.75 ± 1.82 %) and the higher amount of clay (5.13 ± 0.63 %), fine (5.13 ± 0.96 %) and coarse silt (5.98 ± 0.51 %). Grassland recorded the higher amount of total sand (89.20 ± 0.71 %) and the lower amount of clay (3.15 ± 0.50 %) and fine silt (2.93 ± 0.32 %), while intermediate values were found for the maize field and forest (Table 3). These values determine the classification of sandy soil for all measurements sites.

Soil organic carbon (C) and total nitrogen (N) are respectively 12.2 ± 5.7 g kg⁻¹ and 0.95 ± 0.51 g kg⁻¹, averaged for all land cover types over the entire campaign. Table 4 gives soil characteristics for each land cover types, including individual values of C/N ratio, soil organic C and total N for the entire field campaign. The highest average soil organic C was measured for bare soil (17.3 ± 5.9 g kg⁻¹) and the lowest soil organic C was measured for grassland (6.2 ± 1.3 g kg⁻¹), while the maize field and forest site accounted for 14.1 ± 2.9 g kg⁻¹ and 11.4 ± 4.5 g kg⁻¹ soil organic C, respectively. The highest average total N was measured for bare soil (1.44 ± 0.51 g kg⁻¹) and the lowest total N was measured for grassland (0.44 ± 0.04 g kg⁻¹), while the maize field and forest site accounted similar amounts of total N, 0.99 ± 0.19 g kg⁻¹ and 0.94 ± 0.48 g kg⁻¹, respectively. Values of C/N, soil organic C and total N recorded for grassland at the Savè site compare closely to those reported by Delon et al. (2017, table 2) for the semi-arid site of Dahra (15°24' N 15°25' W), Senegal. Our values of C/N and total N for grassland are also close to those reported by Le Roux et al. (1995, table 1) and Lata et al. (2004) for the wet savanna ecosystem of Lamto (6°13' N, 5°20' W), Ivory Coast, although we observe lower values of soil organic C compared to these studies. Values of C/N and soil organic C recorded for the maize field at the Savè site are slightly higher than those recorded by Barthes et al. (2004) in a maize field at Agonkanmey (6°24' N, 2°20' E), near Cotonou in southern Benin.

All the sites listed in the comparison in the previous paragraph are sandy, as the Savè site. The Dahra site (Delon et al., 2017) also shows similar pH than our site (Table 5), while lower pH (acidic or near-neutral) was recorded at the sites of Lamto (Le Roux et al., 1995, Lata et al., 2004) and Agonkanmey (Barthes et al., 2004). Table 5 provides individual values of pH, [NH₄⁺], Γg and χ_g for the entire field campaign. The highest average pH was observed for bare soil (8.23) and the lowest for the forest site (7.07), while measured average pH was 7.27 for grassland and 7.70 for the maize field. The [NH₄⁺] content averaged for all land cover types over the entire campaign is 5.33 ± 4 mg kg⁻¹. The highest average [NH₄⁺] was recorded for

the maize field ($7.9 \pm 6 \text{ mg kg}^{-1}$) and the lowest for grassland ($2.0 \pm 0.3 \text{ mg kg}^{-1}$). Average $[\text{NH}_4^+]$ is $6.2 \pm 5 \text{ mg kg}^{-1}$ and $7.0 \pm 2.2 \text{ mg kg}^{-1}$ for forest and bare soil, respectively. Dick et al. (2006) have found NH_4^+ concentrations between 2 and 8 mgN.kg^{-1} in Senegalese soils, which is very close from our results. Vanlauwe et al. (2002) have found values between 0.8 and 1.4 mgN.kg^{-1} in West African moist savanna soils (in Togo and Nigeria).

Higher soil organic C and N over bare soil could be due to the fact that these bare soil patches experienced recent burning (Santín and Doerr, 2016). The higher $[\text{NH}_4^+]$ over the maize field can be caused by chicken excreta, as chickens were roaming over the maize field (Paillat et al., 2005, Tiquia and Tam, 2000).

3.3 Soil emission potential Γ_g and compensation point χ_g

The mean soil emission potentials for the Savè site is $43\,714 \pm 58\,077$, with values ranging from 380 to 159 343. The highest values of soil emission potentials are observed for bare soil ($113\,672 \pm 67\,788$), followed by maize field ($33\,880 \pm 20\,680$), forest ($11\,982 \pm 11\,061$) and grassland ($4\,929 \pm 4\,409$). The ammonia compensation point ranges between 5 to 2 215 ppb, with soil temperatures between 25 and 29 °C. The highest values of χ_g are observed for bare soil ($1\,607 \pm 993$), followed by maize field (473 ± 317), forest (175 ± 167) and grassland (58 ± 47). Our values of soil emission potential for bare soil and maize (no fertilization) are comparable with those presented in Massad et al. (2010, table 4), although those data come from measurements taken on different ecosystems. Both Γ_g and χ_g values recorded at the Savè site exceed those recorded by Delon et al. (2017) over a grazed semi-arid Sahelian ecosystem in Senegal.

3.4 NO fluxes

NO fluxes from soil measured during the field campaign range between 0 and $48.05 \text{ ngN m}^{-2} \text{ s}^{-1}$. NO fluxes averaged over all land cover types are $4.79 \pm 5.59 \text{ ngN m}^{-2} \text{ s}^{-1}$, while average NO fluxes for each land cover type are: $8.05 \pm 3.49 \text{ ngN m}^{-2} \text{ s}^{-1}$ for bare soil, $3.73 \pm 1.76 \text{ ngN m}^{-2} \text{ s}^{-1}$ for the maize field, $2.87 \pm 1.49 \text{ ngN m}^{-2} \text{ s}^{-1}$ for forest and $2.82 \pm 3.46 \text{ ngN m}^{-2} \text{ s}^{-1}$ for grassland. Soil emissions of NO from the different land cover types provide similar values, NO emissions from bare soil are higher on average, but have a larger standard deviation (Table 6).

Other measurements of biogenic NO soil emissions from the West African wet savanna can be found in Delon et al. (2012, table 7). We find that our measured NO soil emissions averaged over all land cover types are higher than those measured from other wet savanna sites. Our measurements are in better agreement with emissions from dry savanna grasslands (Delon et al., 2012), and with measurements from a semi-arid savanna, with over 80% sandy soil, in South Africa (Parsons et al., 1996, Scholes et al., 1997). However, these studies measured NO emissions during different seasons and soil moisture conditions compared to our study. For example, Parsons et al. (1996) recorded NO emissions up to $20 \text{ ngN m}^{-2} \text{ s}^{-1}$ over an open savanna during the period going from the end of the dry season to the beginning of the wet season. Nitric oxide

455 emissions of the same magnitude as in our study were also recorded over a grazed semi-arid Sahelian ecosystem in Senegal during the month of July by Delon et al. (2017): $5.7 \pm 3.1 \text{ ngN m}^{-2} \text{ s}^{-1}$ in July 2012 and $5.1 \pm 2.1 \text{ ngN m}^{-2} \text{ s}^{-1}$ in July 2013.

Daytime means of NO concentrations are measured close to the soil (0.1m, half height of the chamber) and reported in fig. 2 to 5. Daytime means of NO concentration vary from 1.28 to 5.40 ppb for all sites. The average concentration during the whole campaign on all sites is 2.70 ± 1.03 ppb. Average NO concentration is 2.97 ± 1.49 ppb on bare soil, 2.57 ± 0.96 ppb on grassland, 2.55 ± 0.83 on maize, and 2.76 ± 0.65 ppb on forest soil (Table6). The concentrations are quasi equivalent for all sites. As these concentrations are low, they do not lead to NO deposition on soil and the NO flux stays positive. In fact, NO deposition has been measured in other studies only in the case of high NO concentrations (>60 ppb, Laville et al., 2011).

465 Figures 2 to 5 show daytime averaged NO and NH₃ fluxes (± 1 standard deviation) for each land cover type, along with precipitation and soil moisture. The spatial variability of NO fluxes is high, especially for bare soil, forest and the maize field where underground roots, not visible at the surface, are heterogeneously distributed. These roots are likely to influence the ammonium content of the soil, and the subsequent NO flux measurement. Standard deviation is generally smaller for grassland (except for two days, July 9th and 13th), where the vegetation (and the root distribution) is more homogeneous.

470 The variation of soil moisture is consistent with the presence of rain events, showing a sharper increase of soil moisture at 5 cm, especially after rainfall following dry periods.

NO emissions from bare soil and grassland show an increase, sharper for grassland, one to two days after the rain event on 8th July. The longer rain event between 20th and 24th July does not seem to produce an increase in NO emissions (data available only for maize field and forest). This might be linked with the non-linear relationship between NO biogenic soil emissions and soil water content (Oswald et al., 2013). In fact, a light precipitation event (5-15 mm) occurring on dry soils can result in a large flux of NO (Meixner & Yang, 2006, Hartley & Sclesinger, 2000). However, when soil moisture stays at an equivalent level, after several rain events, pulse emissions do not occur (Millet et al., 2004). Due to this non linear character of the NO fluxes, no direct correlation was found between NO fluxes and environmental variables such as soil moisture or soil temperature taken individually. Moreover, soil temperature and soil moisture were not measured on the same soil parcel where the soil fluxes were measured and the location of the soil flux measurements was not kept constant even for the same land cover type on the same measurement day. This measurement protocol was designed to give an estimate of soil fluxes at a large ecosystem scale, rather than reproducing the relationships between soil fluxes and meteorological variables, like soil temperature and soil moisture.

485 A multiple linear regression analysis was performed between daytime mean NO fluxes and the following variables: wind speed, soil temperature at 5 cm, soil moisture at 5 cm, soil heat flux, upward longwave radiation and downward shortwave radiation. This regression gives $R^2=0.49$ (p-value=0.004), indicating a weak but existing relationship between those variables

and NO soil emissions, while the regression was weak between NO fluxes and each individual variable. This correlation shows the influence of these environmental variables considered collectively on NO fluxes, highlighting the underlying mechanisms responsible for NO release to the atmosphere. Our experiment does not show the details of microbial and physical processes driving soil fluxes at a single point because measurements are done at different locations every day, but aims to estimate the spatial variability of fluxes at the ecosystem scale.

The NO flux estimated in this study does not consider the impact of vegetation on the net ecosystem flux, as we focus on soil fluxes only. However, the net emission to the atmosphere should take into account the oxidation of NO to NO₂ and the eventual re-deposition of NO₂ on the vegetation, i.e. what is called Canopy Reduction Factor and is assumed to be a linear function of the Leaf Area Index (e.g. Yienger and Levy, 1995, and Ganzeveld et al., 2002).

3.5 NH₃ fluxes

NH₃ fluxes measured during the field campaign range between -6.59 and 4.96 ngN m⁻² s⁻¹. Ammonia fluxes averaged over all land cover types are -0.91 ± 1.27 ngN m⁻² s⁻¹, showing a predominance of NH₃ deposition over emission, which is verified for every land cover type, with an average value of: -1.33 ± 0.86 ngN m⁻² s⁻¹ for bare soil, -0.75 ± 0.31 ngN m⁻² s⁻¹ for the maize field, -0.48 ± 0.55 ngN m⁻² s⁻¹ for grassland, and -0.30 ± 0.38 ngN m⁻² s⁻¹ for forest (Table 6). Low positive ammonia fluxes, indicating average NH₃ emission, are only recorded during three days, between 6th and 8th July, after the longest dry period of the measurement campaign (Figures 2 to 5).

As discussed in Appendices B and C, of 30% of individual fluxes used to calculate the daily averages are very low and not distinguishable from adsorption or desorption of NH₃ on chamber walls. These very low fluxes are however meaningful and indicate that some periods of near zero fluxes must be taken into account to represent the processes of exchanges in these ecosystems.

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To our knowledge, NH₃ soil fluxes from ~~west~~West African wet savanna are not available in the scientific literature. In Delon et al. (2017) NH₃ soil fluxes measured in Dahra (15°24' N 15°25' W), Senegal, on a dry savanna ecosystem, show low fluxes with a predominance of NH₃ emission: 1.3 ± 1.1 ngN m⁻² s⁻¹, -0.1 ± 1.1 ngN m⁻² s⁻¹ and 0.7 ± 0.5 ngN m⁻² s⁻¹ over three different measurements campaigns. However, Sutton et al. (2007) shows how pre-cut grassland is characterized by NH₃ deposition, as in our study, in contrast to post-cut grassland, which is marked by NH₃ emission. It is interesting to notice that the literature provides up to about 700 ngN m⁻² s⁻¹ NH₃ emission for fertilized Zea Mays fields (Walker et al., 2013) while in our study site NH₃ deposition was recorded for the maize field, which is not treated with mineral fertilizer.

As for NO concentrations, NH₃ concentrations are reported in fig. 2 to 5. Daytime means of NH₃ concentration vary from nearly 0 to 12.46 ppb for all sites, and the average concentration is 4.42 ± 3.23 ppb during the whole campaign. Average NH₃

concentration is 6.28 ± 3.90 ppb for bare soils, 3.28 ± 1.79 ppb for grassland, 4.36 ± 3.99 for the maize field, and 3.68 ± 2.13 ppb for forest (Table 6). The largest deposition fluxes are found on bare soils, where the largest concentrations are measured.

525 A multiple linear regression analysis was performed between daytime mean NH_3 fluxes and the following variables: wind speed, soil temperature at 5 cm, soil moisture at 5 cm, soil heat flux, outgoing longwave radiation and incoming shortwave radiation). This regression gives a weak but existing relationship, with $R^2=0.37$ (p-value=0.03). This correlation highlights the link between NH_3 fluxes and relevant environmental parameters. However, the same considerations explained in Sect. 3.4 for NO emissions are also valid for the correlation between NH_3 fluxes and meteorological variables.

530 According to the current parameterization of soil ammonia emission potential (Sect. 2.7), high values of pH and $[\text{NH}_4^+]$ in the liquid phase will determine high values of Γ_g indicating that the soil has a high propensity to emit NH_3 . However, despite the high values of Γ_g recorded, our measurement site remains a net sink for NH_3 . The reasons for this can be manifold. One explanation could be that soil particles on our site may have a high adsorption capacity limiting the amount of soil gaseous NH_3 concentrations (Neftel et al., 1998) and the largest part of the estimated ammonium content in the soil may not be in the
535 liquid phase, but adsorbed by solid soil particles. In these conditions ammonium will not be available for gas exchange to open porosity and the atmosphere (Flecharth et al., 2013). Another explanation could be given by the presence of a water film at the soil surface (linked to high air humidity at the site), which will increase the net deposition process. David et al. (2009) conclude from their measurements that the bare soil can be a significant source of NH_3 only for a limited period and only when the cut vegetation is removed, but not if the soil surface remains covered by grass. Measurements in Ferrara et al.
540 (2014) show other occurrences of high soil ammonia emission potential and NH_3 deposition.

Our measurements were conducted without vegetation inside the chambers, but vegetation was present in the fields. It is important to mention that the role of vegetation on NH_3 bidirectional fluxes is essential, especially during the wet season (time of the experiment), when deposition on the vegetation through stomata and cuticles dominate the exchange (during rain
545 events, the cuticular resistance becomes small and cuticular deposition dominates), due to an increase of the deposition velocity of NH_3 (consecutive to the humidity response of the surface) and a decrease of the canopy compensation point, sensitive to the surface temperature and the surface wetness (Wichink-Kruit et al., 2007).

3.6 Comparison of observed and modelled NO soil emissions

We have compared observed daytime averaged (8 a.m. to 6 p.m.) soil NO emissions with those modelled by GEOS-Chem
550 for the entire period of the campaign over the model grid box including the measurement site. The model grid box is positioned at latitude $8.0^\circ \text{ N} - 8.25^\circ \text{ N}$ and longitude $2.19^\circ \text{ E} - 2.5^\circ \text{ E}$. The area of this grid box is 958 km^2 . The land cover type within this grid box is classified as “Savannah (Warm)” but the surrounding area also consists of “Woody Savannah”,

while the observations were taken over the four land cover types representative of the region: bare soil, grassland, maize field and forest.

555

The model is able to reproduce mean air temperature (25.3 ± 0.7 °C) and the main rain events. Soil emissions of NO are well simulated in magnitude. Simulated NO emissions are often higher than those recorded over the grassland areas, however, simulated NO emissions are often within the error bars of measurements (Fig. 6). The model uses land cover and vegetation types to simulate the highly variable land and vegetation cover of the observation site, for this reason we do not expect the model to reproduce the site-to-site variability of the measured soil fluxes, but to at least reproduce their average magnitude and behaviour. It appears that when the model is able to reproduce the length and the intensity of the rain events, NO emissions are especially well simulated, e.g. the model is able to reproduce the longest rain period (from 20th to 26th July 2016) and the decrease of emissions at the end of the measurement campaign.

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3.7 Estimate of total NO soil emissions and NH₃ deposition for Benin

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In order to give a tentative estimate of NO and NH₃ soil fluxes for Benin we have used the land use/land cover map of Benin provided by the US Geographical Survey Atlas: Landscapes of West Africa – A Window on a Changing World (CILSS, 2016, Fig. 7a). The method of mapping land use/land cover used in this atlas was based on Landsat imagery and expert visual interpretation. In particular, these maps provide an accurate indication of cropland distribution using visual interpretation. According to CILSS (2016) Benin's present-day (2013) land cover is mainly savanna, almost 60%, followed by agricultural land, 31%, while forest is only a small fraction under 1% (the rest of the surface is mainly gallery forest and, on a smaller extent, settlements). In the Atlas (CILSS, 2016) bare soils are defined as those surfaces that are bare even in the green/rainy season. For Benin, the amount of bare soil estimated by CILSS (2016) is very small, not big enough to appear on the Atlas' maps. We have multiplied average NO emissions measured at the Savè site for each land cover type by an estimate of the land cover area of each class given by the Atlas. We have made some approximations: as the land use/land cover maps do not distinguish between shrub savanna, tree savanna, and wooded savanna, we have considered NO soil emissions from Savè's grassland savanna to be representative of the general savanna category in CILSS (2016). Moreover, the Atlas has a crop category that does not distinguish the type of crop and we only have observations of NO soil emissions from an intercropped maize field. We have taken NO soil emissions from the maize field as representative of NO emissions of Benin's agricultural land, but other cultures are present in other parts of the country, e.g. oil palm plantations, where possibly stronger fertilization could determine higher NO soil emissions. This tentative calculation gives that Benin's NO soil emissions for the month of July (wet season) is 1.17 ± 0.6 GgN/month, i.e. 0.09% of the average global monthly NO soil emissions as given by Davidson et al., (1997).

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We have also calculated Benin's total monthly NO soil emissions with GEOS-Chem adding together the NO soil emissions from the grid boxes where 50% or greater of the box lies within Benin. Benin's total monthly NO soil emissions calculated

with GEOS-Chem for the month of July are 1.44 GgN/month and agree with the tentative calculation given above ($1.54 \pm 0.8 \text{ GgN/month}$). However, the land cover types covering Benin in GEOS-Chem differ from those in the US Geographical Survey Atlas (CILSS, 2016). In GEOS-Chem Benin is cover by 60.9% savanna, 31.4% woody savannah, 4.5% grassland, 1.3% mixed forest and 0.6% urban and built-up lands. Benin's total monthly NO soil emissions calculated with GEOS-Chem
590 for the months of May and June are higher, 3.51 GgN/month and 2.59 GgN/month , respectively, given those months are at the beginning of the wet season and are characterized by more predominant pulse emissions. Using the same method described above we have upscaled point measurements of NH_3 fluxes with relevant land cover surfaces from CILSS (2016) and obtained that total NH_3 dry deposition for the month of July is $0.21 \pm 0.11 \text{ GgN/month}$ ($0.22 \text{ kgN ha}^{-1} \text{ yr}^{-1}$). This value is about ten times smaller than the estimation of NH_3 dry deposition given in Adon et al. (2013) for the wet savanna site of
605 Djougou (Benin, 9.7°N , 1.7°E) for the month of July, which is around $2.5 \text{ kgN ha}^{-1} \text{ yr}^{-1}$.

4. Conclusion

We provide soil flux measurements along with soil characteristics for a land cover type, savanna, that is considered to have large NO emissions (Davidson and Kinglerlee, 1997), and for an area of the world, West Africa, with little observations. The aim of this study is to contribute to our knowledge in biogenic soil nitrogen exchanges, provide data for inventories and
600 model evaluation to improve air quality and climate modelling.

In situ measurements were made in a wet savanna site in central Benin from mid-June to the end of July 2016. Complementary to these exchange fluxes, soil N and C content, as well as soil pH, soil moisture, soil temperature and meteorological data were measured. Soil fluxes of NO and NH_3 were measured over four different land cover types in order
605 to give a tentative estimate of regional soil fluxes.

Given the set up of the experiment, the known relationships between soil fluxes, soil temperature and soil moisture were not reproduced. Rather than looking at the microbial and physical processes behind soil fluxes, we are able to provide observations that are representative of a bigger surface area and that represent the spatial variability of fluxes. However, we
610 observe that while shorter rain events determine an increase in NO soil emissions, the longer rain event at the end of the campaign (20th to 24th July 2016) is accompanied by a decrease in NO soil emissions, in agreement with the fact that the relationship between NO soil emissions and soil moisture is not univocal. Soil emissions of NO increase until an optimum value of soil moisture is reached and then decrease (Oswald et al., 2013).

615 NH_3 emissions measured in this study probably underestimate total NH_3 emissions for the entire country, as possibly higher localized NH_3 emissions are present in the south of the country where industrial scale agriculture would probably deploy mineral N fertilization.

620 Soil NO emissions simulated by GEOS-Chem are in good agreement with the local observations taken at the site of Savè,
providing a good baseline for simulating local atmospheric chemistry. Moreover, GEOS-Chem is also in good agreement
with the tentative total monthly NO soil emission estimate for Benin for the month of July made with local observation in
Savè and US Geographical Survey Atlas (CILSS, 2016). All these elements contribute to improve our confidence in the
results of modelling studies of local and regional air quality and climate over this region.

625 Agriculture is the first form of economic activity in Benin, occupying a majority of the active population. The most obvious
recent change in land cover is the major expansion of agricultural land across most regions of Benin. Agricultural areas
(including plantations and irrigated agriculture) progressed from 9.2 to 27.1 % of the total country area between 1975 and
2013, improving food security. Oil palm trees are the main crop, and oil palms farmland already covered most of the
southern Terre de Barre plateau of Benin by 1975, and increased by about 28 percent over the following 38-year period. A
630 century or more ago, Benin was covered by dense, biologically diverse forest. Since then, Benin has lost nearly all of that
forest cover, by 2013, 58 % of the 1975 forest cover had been lost, leaving only 0.2 percent of the country covered with
dense forest. Savanna area has also decreased by 23 percent since 1975, but it still remains the dominant land cover type in
Benin and covers more than half of the country (CILSS, 2016).

635 More measurements of NO and NH₃ exchanges between soil-vegetation-atmosphere in areas of Benin (or West Africa)
interested by land-use change could improve our estimate of the impact of biogenic soil emissions on air quality and climate,
as biogenic soil fluxes influence for example the amount of aerosol and tropospheric O₃, a greenhouse gas and pollutant, in
the atmosphere. Management practices of agriculture affect biogenic soil emissions. Moreover, loosing savanna to oil palm
plantations or other crop would have different impacts on air quality, carbon budget and climate than the conversion of forest
640 into crop or oil palm plantation. Furthermore, oil palm plantations are generally closer to the coast and likely to be more
influenced by anthropogenic emissions from industry and coastal cities (Knippertz et al., 2015a, 2015b). Oil palm trees are
also a strong isoprene emitters. Isoprene emissions influence ozone concentration and the oxidizing capacity of the
atmosphere, and it is a source of secondary organic aerosol, thus affecting local air quality and global climate. Large-scale
land use change in the tropics – specifically the conversion of tropical rain forest to oil palm plantations in Malaysia – were
645 shown to cause changes in atmospheric composition and chemistry (Hewitt et al., 2009), indicating that the management of
the emissions of reactive nitrogen species is essential to prevent damaging levels of ground-level ozone in those regions.

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Appendix A

The dilution uncertainty is calculated based on the uncertainties of standard concentration, standard flow and dilution flow. The uncertainty of standard concentration is 5% for NO and NO₂, 2% for NH₃. The maximum uncertainty of dilution flow is 1% of the plain scale (10 L.min⁻¹) for the three standards divided by the flow used in the diluter (3.2 L.min⁻¹ maximum), which gives 0.1/3.2=3.1%. The uncertainty of standard flow is 1% of the plain scale standard flow (50 mL.min⁻¹) divided by the standard flow used to obtain the needed concentration (50 ppb for NO or 30 ppb for NH₃).

Standard flow=(needed concentration/standard concentration)*dilution flow.

660 **For NO**, dilution flow=3.2 L.min⁻¹, needed concentration=50 ppb, standard concentration=8.73 ppm, standard flow=18.4 mL.min⁻¹. Uncertainty of the standard flow = 1%*50/18.4=2.71%. Total uncertainty is therefore 5%+3.1%+2.7%=**10.8%**.

For NO₂, dilution flow=3.2 L.min⁻¹, needed concentration=50 ppb, standard concentration=9.28 ppm, standard flow=17.2 mL.min⁻¹. Uncertainty of the standard flow = 1%*50/17.2=2.9%. Total uncertainty is therefore 5%+3.1%+2.9%=**11%**

For NH₃, dilution flow=3.2 L.min⁻¹, needed concentration=30 ppb, standard concentration=14.78 ppm, standard flow=6.5 mL.min⁻¹. Uncertainty of the standard flow = 1%*50/6.5=7.7%. Total uncertainty is therefore 2%+3.1%+7.7%=**12.8%**.

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Appendix B

We ran a laboratory experiment to verify that deposition on the walls of the Teflon chamber is negligible.

Ambient air concentrations were measured by the analyzer, inside the room where the analyzer and the chamber were placed.

675 Measurements of NH₃ concentrations were made in ambient air with and without the Teflon chamber attached to the analyzer. The Teflon chamber was placed on a Teflon frame, and they were sealed together with Teflon tape. Measurements of NH₃ concentrations with the Teflon chamber attached to the analyzer were followed by measurements without the chamber 30 to 60 minutes later. The two sets of measurements were made under similar conditions of temperature and humidity. Average values of NH₃ concentrations were calculated for 10 to 30 minutes before and after connecting the chamber. Average NH₃ concentrations during this time interval varied between 8 and 36 ppb, with a variation between 1.5 to 13% around the mean. The lowest NH₃ concentrations correspond to air samples previously passed through charcoal and desiccant cartridges (NO and NO₂ zero air). Measured NH₃ concentrations are reported in Table B1, along with temperature, humidity and the ratio between average concentration with and without the Teflon chamber attached to the analyzer.

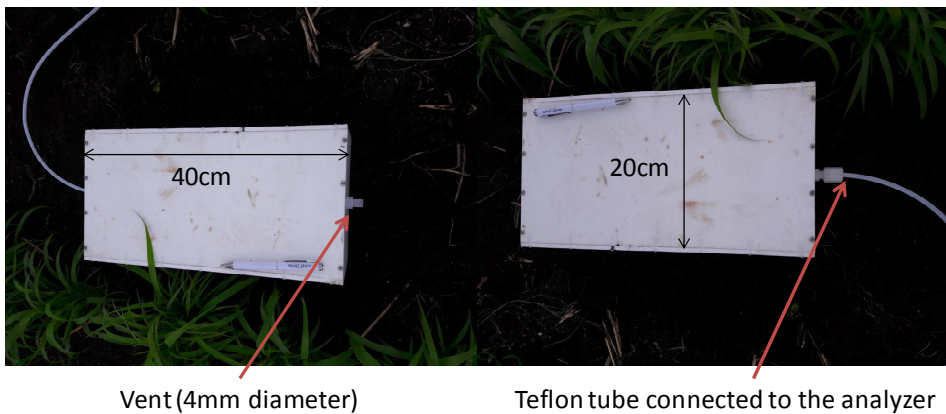
685 This test was made at different times of the day on different days: air humidity varied between 46 and 54%, temperature varied between 25 to 29°C, while pressure varied between 1006 and 1008 hPa (not reported).

Results show negligible variation between concentrations of air reaching the analyzer via the chamber or going directly to the analyzer; The average difference in concentration is 0.9 ppb, and should be considered as the detection limit for fluxes significantly different from zero (i.e. including potential effects of adsorption or desorption). 30% of the concentration differences are below 0.9 ppbv.

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Temperature (°C)	Humidity (%)	[NH ₃] TC	[NH ₃] D	Ratio [NH ₃]TC / [NH ₃]D
25	54	26.4±0.4	25.9±0.5	1.02
27	49	9.0±0.8	8.08±1.2	1.12
29	46	35.2±2.4	36.0±1.4	0.98
28	46	25.6±0.7	24.0±1.0	1.07
27	45	26.2±0.9	27.4±0.9	0.96
28	46	23.4±0.3	24.1±0.4	0.97
26	50	19.2±0.6	18.4±0.8	1.04

Table B1 Measurements of NH₃ concentrations (ppb) through the chamber (TC) or directly (D) to the analyzer.



Side of the chamber: vent and tube connection located in the middle of the square

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Picture B1: Description of the Teflon chamber.

To verify if mixing the air in the chamber by a fan would change the shape of the increase in concentration or the concentration itself in the chamber, a test was made with a syringe simulating the action of a fan (i.e. we have mixed the air inside the chamber by sucking and releasing the same air with a syringe through the small vent, while letting outside air entering the chamber by the vent as usual to ensure pressure equilibrium between outside and inside air). The comparison between a flux measurement with and without mixing gives similar slopes of the concentration increase (or decrease).

Appendix C

NH₃ concentration was recorded continuously inside the Teflon chamber (placed on a Teflon material as in the tests summarized in Appendix A), and the chamber was cleaned successively with three different papers, referred to as A, B and C. The concentration was recorded at least for 30 minutes between every cleaning. Table C1 summarizes the averaged concentrations (and standard deviations) for every period. Results show a variation of concentration when different papers are used, but this variation is not reproducible and is difficult to differentiate from a natural variation of the NH₃ concentration in the room. As a matter of fact, the effect of cleaning on NH₃ adsorption or desorption is not clear, but questions about the potential pollution of the chamber arise. Results in Table C1 may lead to the conclusion that if the difference in concentration during a flux measurement is inferior to a certain threshold, it is not necessarily a flux from the ground but could be due to ~~an~~ adsorption or desorption of NH₃ by the chamber walls due to cleaning. Only low fluxes are concerned. ~~To set ideas down and take an example, Positive or negative~~ fluxes inferior to 0.55 ngN.m⁻².s⁻¹ (corresponding to concentration differences less than 0.9 ppb in the flux calculation, as defined in Appendix B) represent 23.30% of the 350 measured fluxes. If those low fluxes were removed from the database, the resulting average would be slightly larger in magnitude (-1.1 instead of -0.9 ngN.m⁻².s⁻¹). As a conclusion, these tests may help to warn the reader that caution must be kept for low NH₃ fluxes because of possible pollution in the chamber.

Paper used for cleaning	30 minutes average (standard deviation) in ppb	Difference between two successive averages in ppb
First day of test		
Before cleaning	8.13±0.58	
A	7.43±0.47	-0.69

B	8.55±0.79	1.12
C	9.73±0.81	1.18
B	9.75±0.88	0.02
Second day of test		
B	16.35±0.92	
C	16.77±0.60	0.42
A	17.30±0.76	0.53
A	18.94±0.72	1.64
B	18.96±0.62	0.02
C	18.79±0.79	-0.17

720 Table C1: 30 minutes averaged concentrations in the Teflon chamber after cleaning with different dry papers.

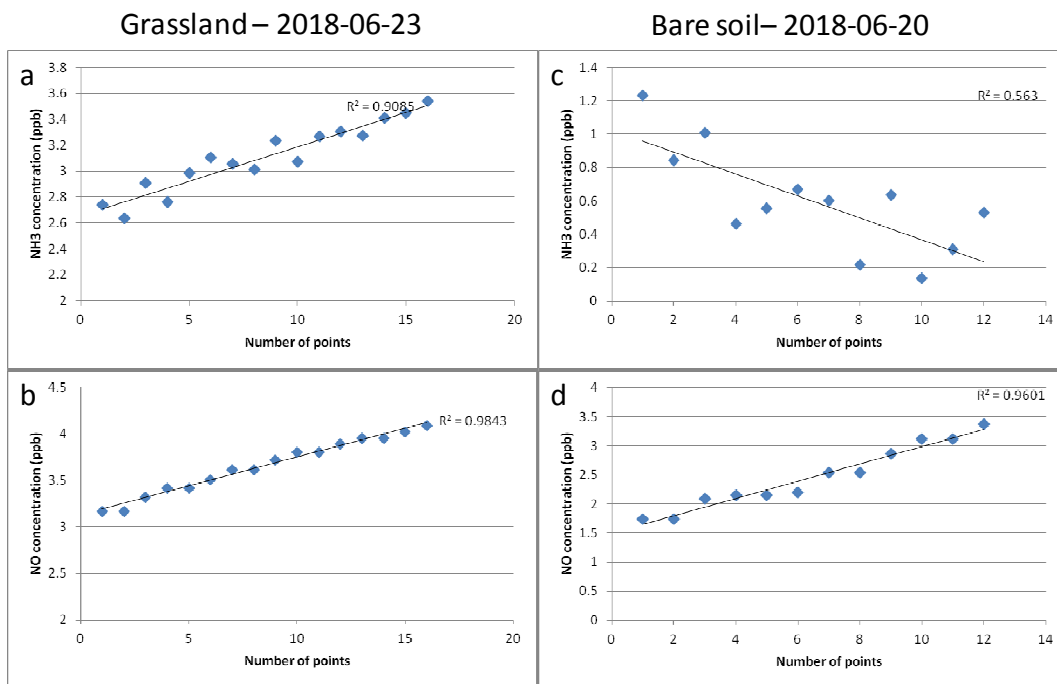
Appendix D

Location of the temperature measurement	Temperature (°C)
Air	32.7
Soil	34.2
Chamber: outside wall	33.1
Chamber: inside wall	33.3
Chamber: outside top	33.8
Tube: outside close to the chamber	30.5
Tube : outside close to the analyzer	30.3
Tube: inside	30.9

725 Table D1 : Temperature measured on the Teflon chamber and on the Teflon tube. These measurements have been made after the field campaign in direct sunlight at 3:30 PM. Measurements were made with a calibrated thermometer HI 98509 with stainless steel probe (-50 → +150 °C).

Appendix E

730 NO and NH₃ fluxes are calculated from the slope of their concentration increase (or decrease) in the chamber through time. Two examples are given in figures E1 to illustrate the larger instability of NH₃ detection compared to NO detection, due to possible interaction of NH₃ with chamber walls, or particulate matter in the chamber.



735 Figure E1: NH₃ (a,c) and NO concentration (b,d) variation with time (one point every 10 second) inside the chamber on grassland (left) and bare soil (right).

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	Savè ground-based observation site
Location	8°02'03" N, 2°29'11" E
Elevation	166 m a.s.l.
Mean annual precipitation	1100 mm
Mean annual temperature	27.5 °C
Soil type	sandy
Sand percentage	87%
Clay percentage	4.1%

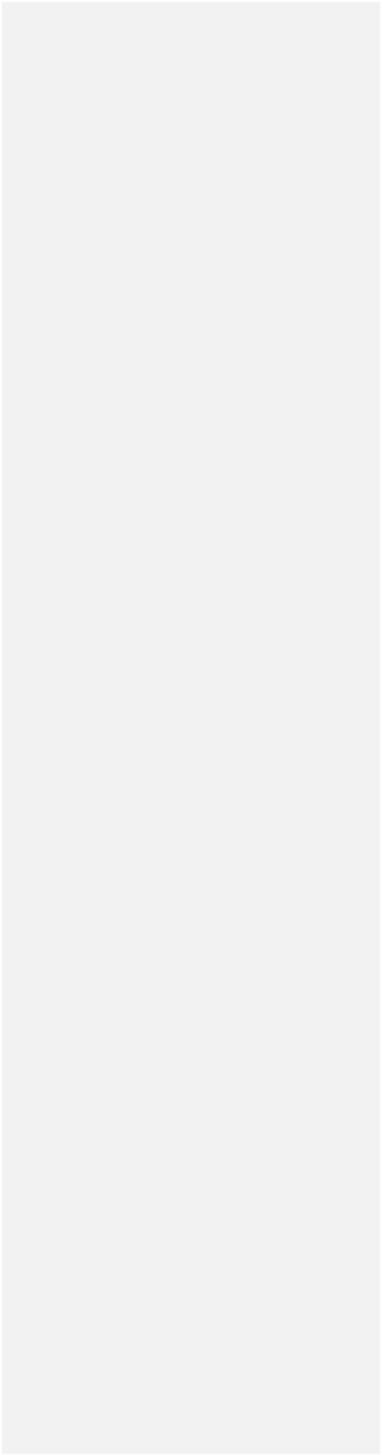
945 Table 1
Main characteristics of the Savè site.

Soil type		Plant family	Plant species	Common name/s	
Next to grassland and forest	Dominant tree species	Anacardiaceae	<i>Anacardium occidentale</i>	cashew tree	
		Fabaceae	<i>Daniellia oliveri</i>	African copaiba balsam tree	
			<i>Pterocarpus erinaceus</i>	barwood, muninga, vène, mukwa	
	Dominant ground species	Cleomaceae	<i>Cleome</i> sp.	spider flowers, spider plants	
		Fabaceae	<i>Crotalaria</i> sp.	rattlepod or rattlebox	
			<i>Mucuna</i> sp.	velvet bean	
		Poaceae	<i>Imperata cylindrica</i>	cogon grass, cotton wool grass, kura-kura	
		<i>Rhynchelytrum repens</i>	rose natal grass		
Next to maize field	Dominant tree species	Anacardiaceae	<i>Mangifera indica</i>	mango	
		Arecaceae	<i>Cocos nucifera</i>	coconut tree	
		Caricaceae	<i>Carica papaya</i> L.	papaya	
		Lamiaceae	<i>Tectona grandis</i>	teak	
		Meliaceae	<i>Azadirachta indica</i>	neem, nintree, Indian lilac	
	Dominant ground species	Commelinaceae	<i>Commelina benghalensis</i>	benghal dayflower, tropical spiderwort	
		Euphorbiaceae	<i>Euphorbia</i> sp.	spurge	
		Nyctaginaceae	<i>Boerhavia diffusa</i>	punarnava, red spiderling	
		Phyllanthaceae	<i>Phyllanthus amarus</i>	gale of the wind, stonebreaker	
	Crops		Poaceae	<i>Digitaria horizontalis</i>	Jamaican crabgrass
			Dioscoreaceae	<i>Dioscorea</i> sp.	yam
			Euphorbiaceae	<i>Manihot esculenta</i>	cassava
			Fabaceae	<i>Arachis hypogaea</i>	peanut
				<i>Vigna unguiculata</i>	cowpea
			Malvaceae	<i>Gossypium</i> sp.	cooton
			Pedaliaceae	<i>Sesamum indicum</i>	sesame
			Poaceae	<i>Zea mays</i>	maize
		<i>Sorghum</i> sp.	sorghum		
	Solanaceae	<i>Solanum lycopersicum</i>	tomato		

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Table 2

List of plant species at the Savè site. The list of common names is not considered to be exhaustive.



	Bare Soil	Grassland	Maize Field	Forest
Clay (<2µm) (%)	5.13±0.63	3.15±0.50	4.40±0.35	3.70±1.25
Fine Silt (2 to 20µm) (%)	5.13±0.96	2.93±0.32	4.13±1.00	3.40±1.21
Coarse Silt (20 to 50µm) (%)	5.98±0.51	4.78±0.66	4.37±0.38	4.67±1.05
Total Sand (50 to 2000 µm) (%)	83.75±1.82	89.20±0.71	87.13±0.99	88.20±3.50

Table 3

List of soil characteristics for each land cover type at the Savè site, including standard deviation.

Soil Type	Date	C/N ratio	Organic C (g kg ⁻¹)	Total N (g kg ⁻¹)
Bare soil	06/07/2016	14.90	24.28	1.63
	09/07/2016	12.20	10.47	0.86
	19/07/2016	12.30	15.05	1.22
	28/07/2016	9.50	19.30	2.04
Grassland	07/07/2016	15.20	5.78	0.38
	09/07/2016	16.00	7.36	0.46
	19/07/2016	14.90	7.16	0.48
	28/07/2016	10.90	4.56	0.42
Maize	09/07/2016	14.80	17.33	1.17
	19/07/2016	16.40	13.08	0.8
	28/07/2016	11.80	11.83	1.00
Forest	06/07/2016	14.80	7.98	0.54
	19/07/2016	11.90	9.62	0.81
	28/07/2016	11.30	16.56	1.47

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Table 4

List of soil characteristics for each land cover type at the Savè site for each soil sampling day: Carbon-to-Nitrogen ratio (C/N), organic carbon (g kg⁻¹) and total nitrogen (g kg⁻¹). The accuracy for the C/N ratio is 14%. The measurement accuracy for organic carbon and total nitrogen is 14 and 13%, respectively.

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Soil Type	Date	pH	[NH ₄ ⁺] (mg kg ⁻¹)	Γ _g [NH ₄ ⁺]/[H ⁺]	X _g (ppb)
Bare soil	06/07/2016	8.38	6.82	136 334	1891
	09/07/2016	7.73	2.90	12 978	134
	19/07/2016	8.46	6.63	159 343	2188
Grassland	28/07/2016	8.34	8.01	146 033	2215
	07/07/2016	7.15	1.96	2 307	29
	09/07/2016	7.91	1.55	10 499	108
	19/07/2016	7.52	2.28	6 291	86
	28/07/2016	6.51	2.30	620	9
Maize	09/07/2016	7.46	4.40	10 575	109
	19/07/2016	7.61	14.74	50 040	687
	28/07/2016	8.04	4.49	41 027	622
Forest	06/07/2016	6.32	2.18	380	5
	19/07/2016	7.51	4.88	13 159	181
	28/07/2016	7.37	11.47	22 407	340

975 Table 5

List of soil pH, ammonium concentrations [NH₄⁺] (mg kg⁻¹), soil emission potential Γ_g and soil compensation point χ_g (ppb) for each land cover type at the Savè site for each soil sampling day. The measurement accuracy for pH is 0.15 when pH ≤ 7 and 0.20 when pH > 7. The accuracy for ammonium concentrations [NH₄⁺], soil emission potential Γ_g and soil compensation point χ_g is 25%.

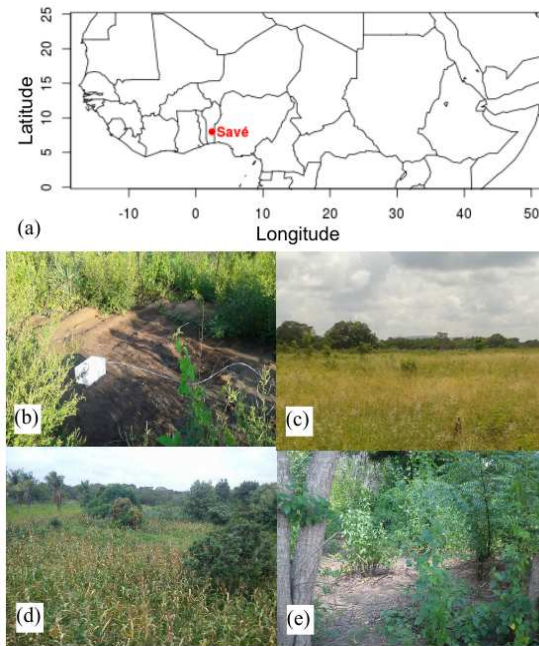
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	Bare soil	Grassland	Maize field	Forest	Mean over all land cover types
NO fluxes (ng.m ⁻² .s ⁻¹)	8.05 ± 3.49	2.82 ± 3.46	3.73 ± 1.76	2.87 ± 1.49	4.79 ± 5.59
NH ₃ fluxes (ng.m ⁻² .s ⁻¹)	-1.33 ± 0.86	-0.48 ± 0.55	-0.75 ± 0.31	-0.30 ± 0.38	-0.91 ± 1.27
NO concentration (ppb)	2.97 ± 1.49	2.57 ± 0.96	2.55 ± 0.83	2.76 ± 0.65	2.70 ± 1.03
NH ₃ concentration (ppb)	6.28 ± 3.90	3.28 ± 1.79	4.36 ± 3.99	3.68 ± 2.13	4.42 ± 3.23

Table 6: List of average NO and NH₃ fluxes (ngN.m⁻².s⁻¹) and concentrations (ppb) for bare soil, grassland, maize field and forest sites, and for all cover types.

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1020 Fig. 1(a) Location of the Savè site in West Africa, (b) one of the bare soil sampling sites, (c) the grassland sampling site, (d) the maize field sampling site and (e) the forest sampling site at the Savè site.

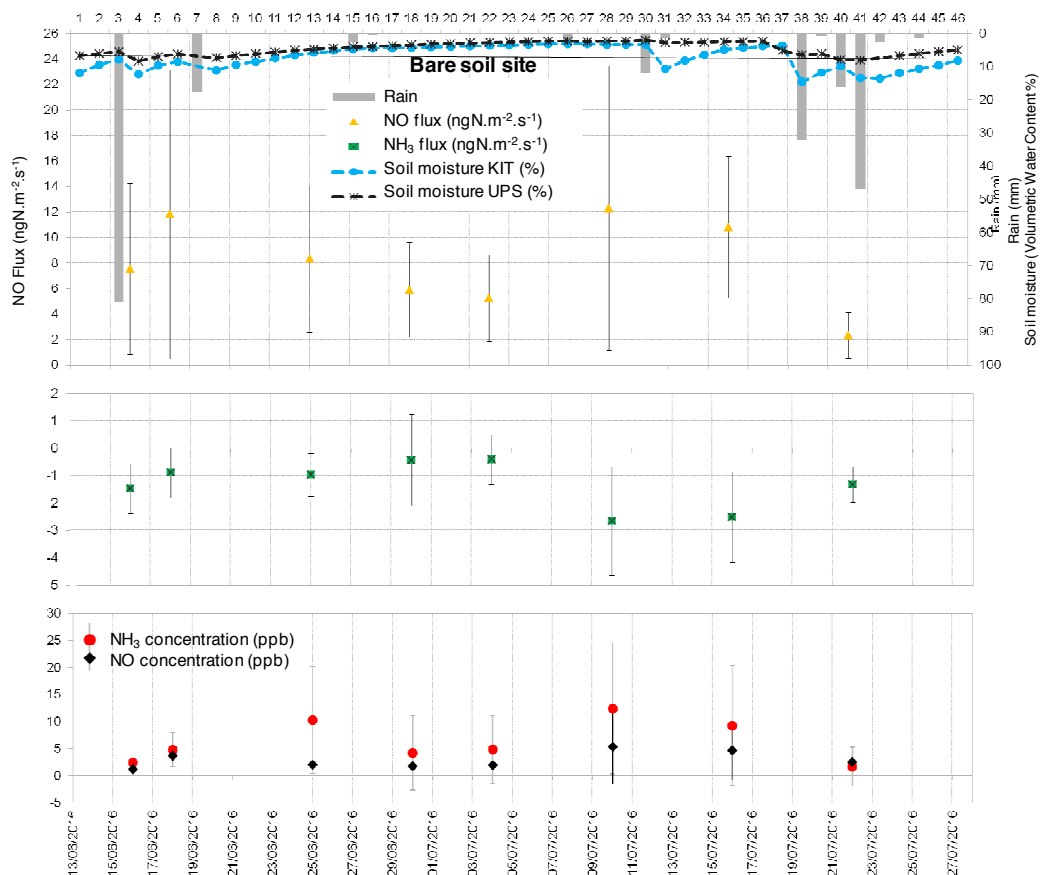
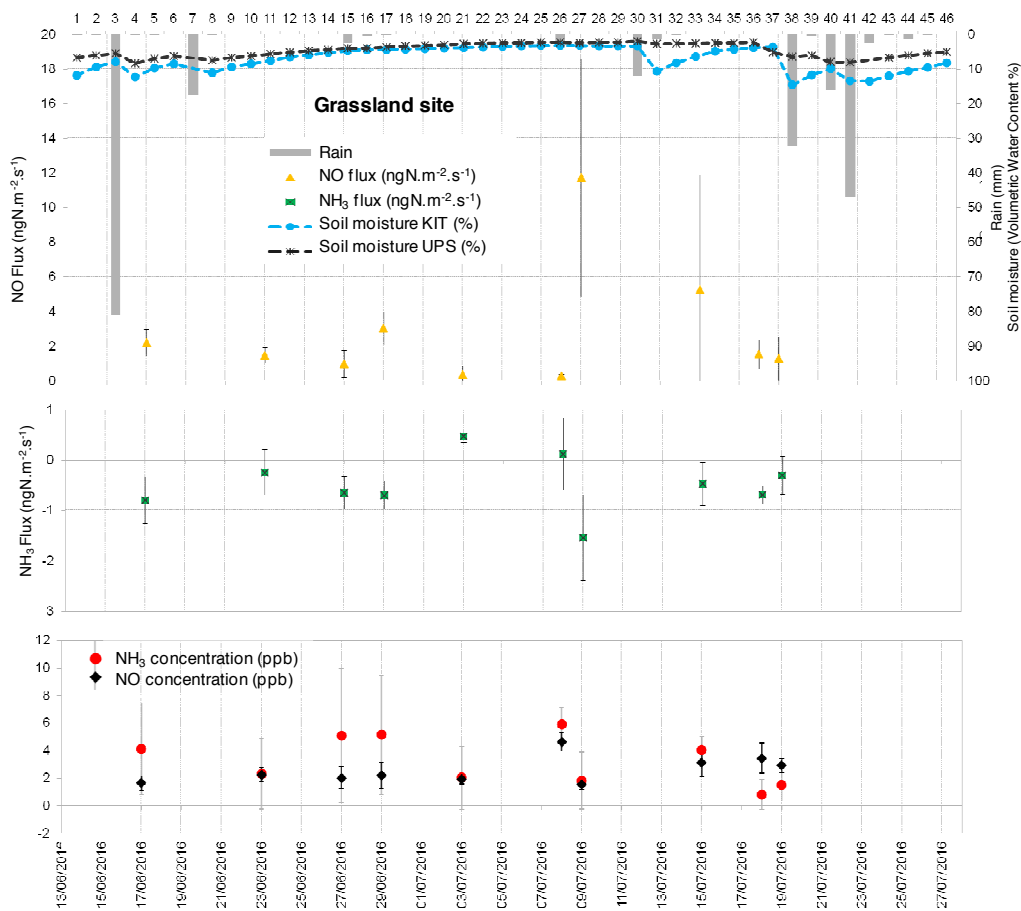


Fig. 2 Upper panel: Daily total precipitation (mm), daily mean soil moisture at 5 cm (%) measured by the Karlsruhe Institute of Technology (KIT), daily mean soil moisture averaged between 0 and 30 cm measured by the Université Paul Sabatier (UPS) instrumentation, daytime mean NO fluxes in $\text{ngN}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ measured at the bare soil site; Middle panel: daytime mean NH₃ fluxes in $\text{ngN}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$; Lower panel: daytime mean NO and NH₃ concentrations in ppb. Vertical bars show the standard deviation from individual fluxes and concentrations



1035 Fig. 3 Upper panel: Daily total precipitation (mm), daily mean soil moisture at 5 cm (%) measured by the Karlsruhe Institute of Technology (KIT), daily mean soil moisture averaged between 0 and 30 cm measured by the Université Paul Sabatier (UPS) instrumentation, daytime mean NO fluxes in $\text{ngN m}^{-2} \text{s}^{-1}$ measured at the grassland site; Middle panel: daytime mean NH₃ fluxes in $\text{ngN m}^{-2} \text{s}^{-1}$; Lower panel: daytime mean NO and NH₃ concentrations in ppb. Vertical bars show the standard deviation from individual fluxes and concentrations.

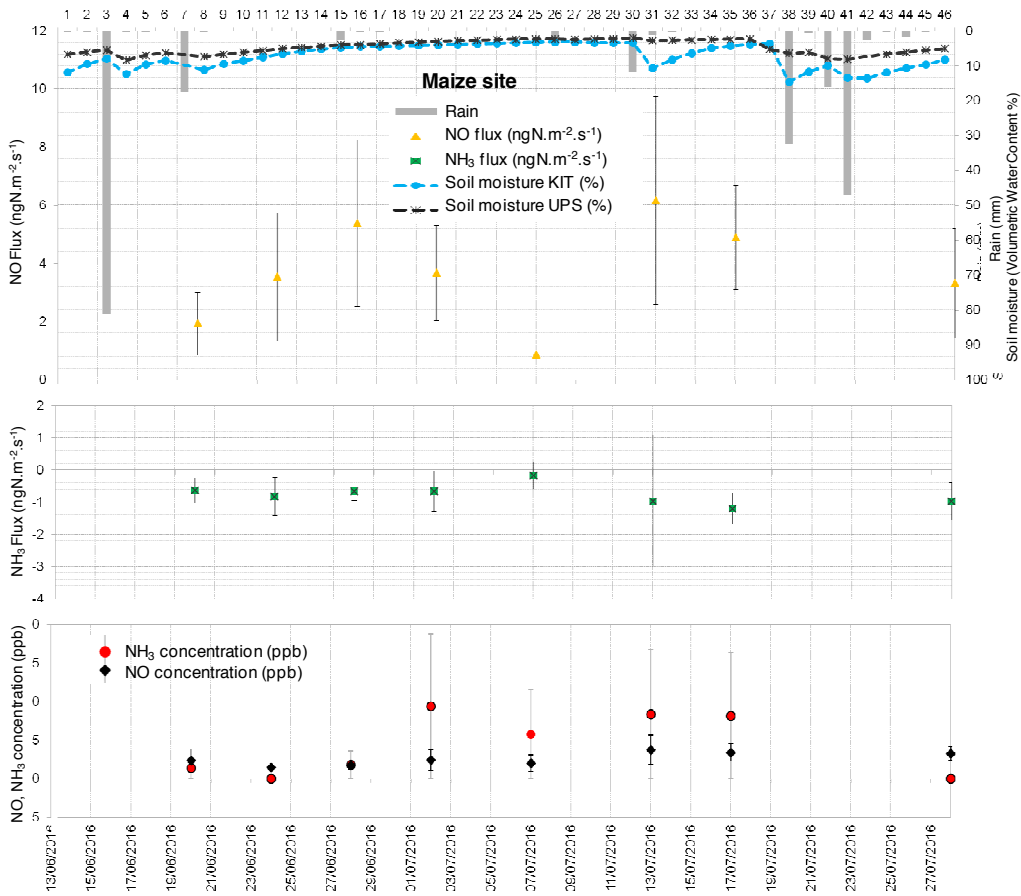
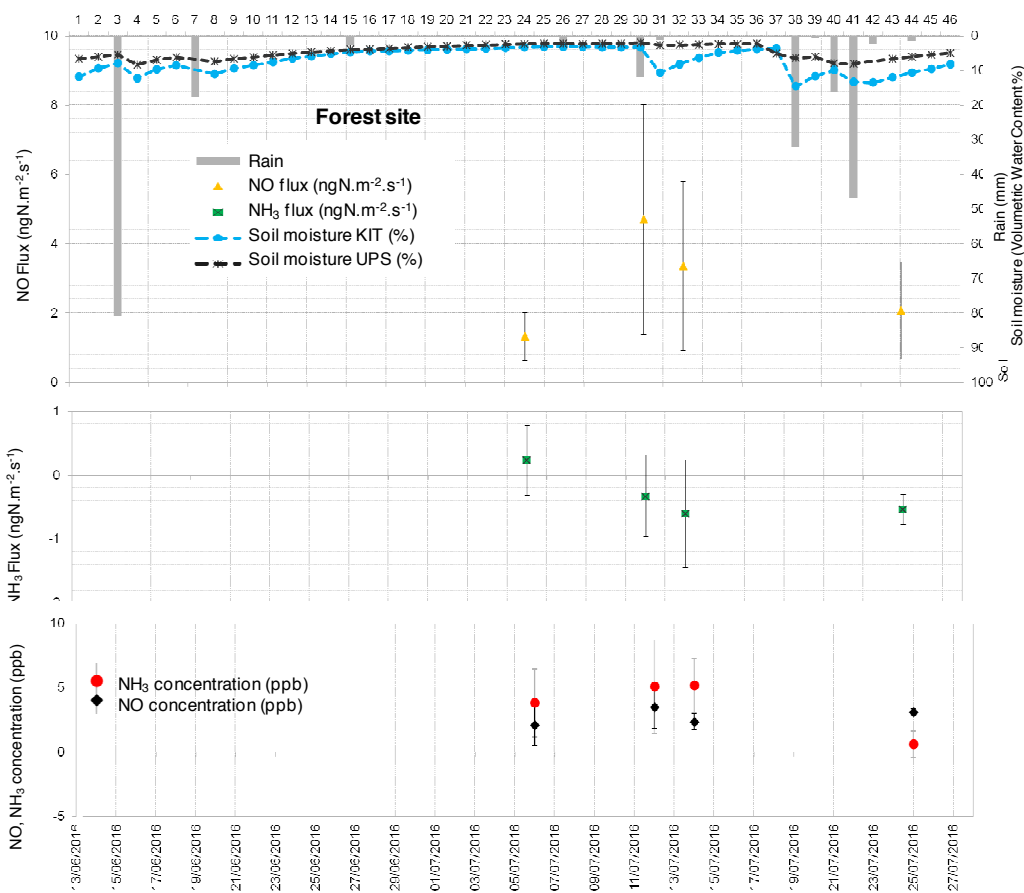
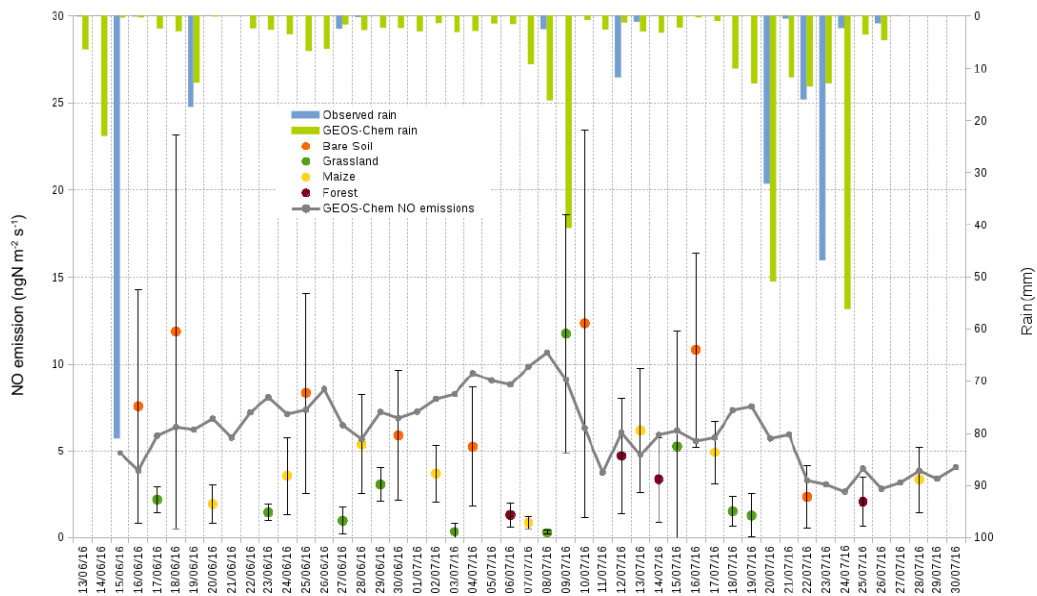


Fig. 4 Upper panel: Daily total precipitation (mm), daily mean soil moisture at 5 cm (%) measured by the Karlsruhe Institute of Technology (KIT), daily mean soil moisture averaged between 0 and 30 cm measured by the Université Paul Sabatier (UPS) instrumentation, daytime mean NO fluxes in $\text{ngN m}^{-2} \text{s}^{-1}$ measured at the maize field site ; Middle panel: daytime mean NH_3 fluxes in $\text{ngN m}^{-2} \text{s}^{-1}$; Lower panel: daytime mean NO and NH_3 concentrations in ppb. Vertical bars show the standard deviation from individual fluxes and concentrations.

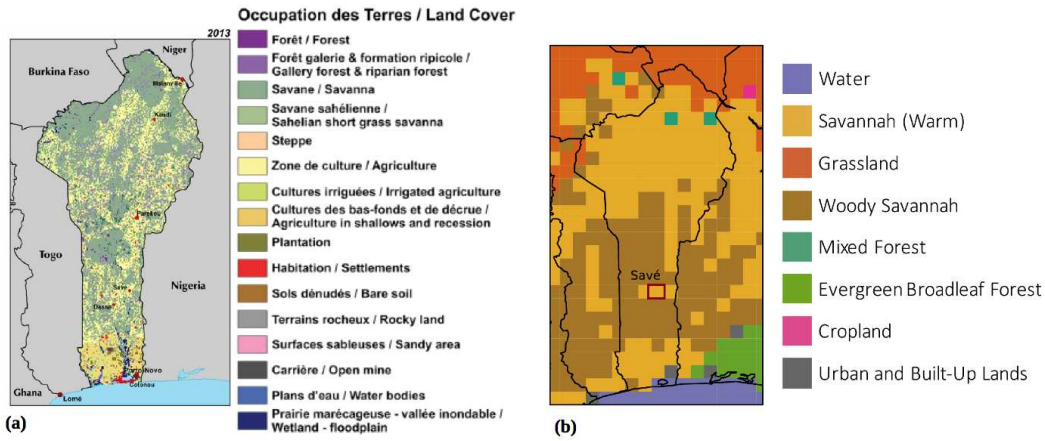


1050 Fig. 5 Upper panel: Daily total precipitation (mm), daily mean soil moisture at 5 cm (%) measured by the Karlsruhe Institute of Technology (KIT), daily mean soil moisture averaged between 0 and 30 cm measured by the Université Paul Sabatier (UPS) instrumentation, daytime mean NO fluxes in $\text{ngN}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ measured at the forest site ; Middle panel: daytime mean NH₃ fluxes in $\text{ngN}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$; Lower panel: daytime mean NO and NH₃ concentrations in ppb. Vertical bars show the standard deviation from individual fluxes and concentrations.



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Fig. 6 Nitric oxide emissions in $\text{ngN m}^{-2} \text{s}^{-1}$ measured over each land cover type (orange dot for bare soil, green for grassland, yellow for the maize field and brown for forest) and simulated with GEOS-Chem, along with rainfall measured and modelled with GEOS-Chem. Soil NO emissions are daily average between 8 a.m. and 6 p.m..



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Fig. 7 (a) Land cover map of Benin for 2013 from the US Geographical Survey Atlas: Landscapes of West Africa – A Window on a Changing World (CILSS, 2016) and (b) land cover map of Benin used in the GEOS-Chem simulation.