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Influence of local air pollution on the deposition of peroxyacetyl nitrate to a nutrient-poor natural grassland ecosystem

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Dry deposition of peroxyacetyl nitrate (PAN) is known to have a phytotoxic impact on plants under photochemical smog conditions, but it may also lead to higher productivity and threaten species richness of vulnerable ecosystems in remote regions. However, underlying mechanisms or controlling factors for PAN deposition are not well understood and studies on dry deposition of PAN are limited. In this study, we investigate the impact of PAN deposition on a nutrient-poor natural grassland ecosystem situated at the edge of an urban and industrialized region in Germany. PAN mixing ratios were measured within a 3.5 months summer to early autumn period. In addition, PAN fluxes were determined with the modified Bowen ratio technique for a selected period. The evaluation of both stomatal and non-stomatal deposition pathways was used to model PAN deposition over the entire summer-autumn period. We found that air masses at the site were influenced by two contrasting pollution regimes, which lead to median diurnal PAN mixing ratios ranging between 50 and 300 ppt during unpolluted and between 200 and 600 ppt during polluted episodes. The measured PAN fluxes showed a clear diurnal cycle with maximal deposition fluxes of $\sim -0.1 \text{ nmol m}^{-2} \text{ s}^{-1}$ (corresponding to a deposition velocity of 0.3 cm s^{-1}) during daytime and a significant non-stomatal contribution was found. The ratio of PAN to ozone deposition velocities was found to be ~ 0.1 , which is much larger than assumed by current deposition models. The modelled PAN flux over the entire period revealed that PAN deposition over an entire day was $333 \mu\text{g m}^{-2} \text{ d}^{-1}$ under unpolluted and $518 \mu\text{g m}^{-2} \text{ d}^{-1}$ under polluted episodes. Besides, thermochemical decomposition PAN deposition accounted for 32 % under unpolluted episodes and 22 % under polluted episodes of the total atmospheric PAN loss. However, the impact of PAN deposition as a nitrogen source to the nutrient-poor grassland was estimated to be only minor, under both unpolluted and polluted episodes.

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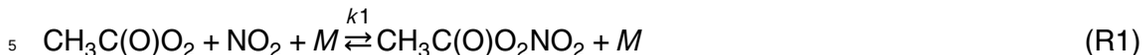
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1 Introduction

Originating from both anthropogenic and natural sources, peroxyacetyl nitrate ($\text{CH}_3\text{C}(\text{O})\text{O}_2\text{NO}_2$, PAN) is primarily known as an atmospheric pollutant. Both, the peroxyacetyl radical ($\text{CH}_3\text{C}(\text{O})\text{O}_2$, PA) and nitrogen dioxide (NO_2), which form PAN via



have anthropogenic sources. Due to its thermal instability via the back reaction of (R1) and subsequent reaction of PA with nitric oxide (NO),



long range transport of PAN in cold layers of the upper troposphere may constitute a significant source of reactive nitrogen (N_r) in remote regions. Consequently, it affects e.g. the production of ozone (O_3) and links the atmospheric and biospheric nitrogen cycle through dry deposition (Singh, 1987). Besides, locally produced PAN may also impact on ecosystems downwind of pollution sources. While high PAN mixing ratios (> 15 ppb), prevailing under strong photochemical smog conditions, PAN is known to be phytotoxic and may harm plant tissues significantly (Temple and Taylor, 1983), the impact of PAN deposition under less extreme conditions and for lower PAN mixing ratios is not yet clear. As a nitrogen source, PAN deposition may also lead to higher productivity and may threaten species richness especially in vulnerable ecosystems (Stevens et al., 2010).

Previous studies on the surface-atmosphere exchange of PAN showed that PAN is deposited to vegetation. On the one hand, chamber experiments on PAN uptake (Okano et al., 1990; Sparks et al., 2003; Teklemariam and Sparks, 2004) found a direct relationship between PAN uptake and stomatal conductance. They suggest that stomatal uptake is the major pathway of PAN into leaves. On the other hand, previous studies have also shown the existence of non-stomatal deposition of PAN, mainly associated with the uptake by the leaf cuticles (Teklemariam and Sparks, 2004; Turnipseed

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fluxes via the modified Bowen ratio (MBR) technique (Businger, 1986). The PAN flux ($F_{\text{MBR}_{\text{PAN}}}$) was estimated by the ratio of the PAN and O_3 mixing ratio difference between the upper and lower measurement height, ΔC_{PAN} and ΔC_{O_3} , multiplied by the eddy covariance flux of O_3 ($F_{\text{EC}_{\text{O}_3}}$):

$$F_{\text{MBR}_{\text{PAN}}} = F_{\text{EC}_{\text{O}_3}} \cdot \frac{\Delta C_{\text{PAN}}}{\Delta C_{\text{O}_3}} \approx F_{\text{EC}_{\text{O}_3}} \cdot \frac{\Delta \chi_{\text{PAN}}}{\Delta \chi_{\text{O}_3}} \quad (1)$$

We used O_3 as a proxy scalar due to its similarity to PAN in the sink and source distribution. On the one hand, the production of both PAN and O_3 is linked to photochemical processes and, on the other hand, both compounds are known to deposit to vegetation. Instead of using concentration differences in Eq. (1), we used the mixing ratio differences of PAN ($\Delta \chi_{\text{PAN}}$) and O_3 ($\Delta \chi_{\text{O}_3}$) since the differences in the molar air density between the two heights were negligible. The correction of PAN fluxes for the loss by thermochemical decomposition of PAN is presented in Sect. 2.5. The storage term (see e.g., Rummel et al., 2007) of PAN was estimated using a logarithmically interpolated vertical profile of PAN and was found to be negligible. Further details on the flux measurements, including necessary modifications of the GC-ECD and the inlet system, and an extensive error analysis are given in Moravek et al. (2014). Flux values with random errors larger than 100 % were regarded as below the flux detection limit. Furthermore, under conditions with low friction velocities ($u_* < 0.07 \text{ m s}^{-1}$) the application of the MBR methods is prone to larger errors (Liu and Foken, 2001).

2.3 Additional measurements

For the determination of O_3 eddy covariance fluxes, required for the application of the MBR method, a closed-path fast response O_3 detector (Enviscope GmbH, Germany) was employed together with a sonic anemometer (CSAT3, Campbell Scientific Inc., USA) at $z_{\text{ref}} = 3 \text{ m a.g.l.}$ (see Moravek et al., 2014, for details on eddy covariance set up and O_3 calibration). In addition, CO_2 and latent heat fluxes were determined with

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an open-path CO₂/H₂O analyser (LI-7500A, LI-COR, USA). All turbulent fluxes were calculated using the eddy covariance software TK3.1 (Mauder and Foken, 2011), applying state-of-the-art corrections methods as listed in Foken et al. (2012). Additionally, the O₃ flux was corrected for high frequency loss of the 2.5 m long inlet tube (Moravek et al., 2013), for the storage effect and for chemical production from NO₂ photolysis and loss by reaction with NO (Rummel et al., 2007). The nighttime removal pathway of O₃ via the reaction with NO₂ forming NO₃ (see e.g., Brown and Stutz, 2012) was found to be insignificant as the reaction rate is about one order of magnitude lower than of the reaction of O₃ with NO. The quality scheme of Foken and Wichura (1996) was used to exclude periods with significant non-stationarity or poor developed turbulence. Data for which the footprint area of the flux measurement (calculated with a Lagrangian forward stochastic model from Rannik et al., 2000) included less than 80 % of the natural grassland area were omitted.

The mixing ratio difference of O₃ between 4.0 and 0.8 m a.g.l. was determined using a differential UV absorption O₃ analyser (49i, Thermo Environmental, USA, modified according to Cazorla and Brune (2010); see Moravek et al. (2014) for details on operation). Absolute O₃ mixing ratios at both heights were derived from a vertical profiles system, which also measured NO and NO₂ mixing ratios (O₃ analyser: 49i, Thermo Environmental, USA; NO/NO₂ analyser: CLD 780 TR, Eco-Physics, Switzerland). A vertical profile of temperature, humidity and wind speed was retrieved at 0.2, 0.8, 1.5, 2.5 and 4.0 m a.g.l.

Meteorological parameters used in this study are global radiation (CNR1, Kipp&Zonen, the Netherlands), NO₂ photolysis frequency (j_{NO_2}) (Meteorology Consult GmbH, Germany), rainfall (AGR100, Environmental Measurements) and surface wetness (Campbell Scientific Inc., USA). All additional measurements were performed during the entire experiment period from 29 June to 21 October 2011.

Finally, g_{ns} , representing all non-stomatal deposition pathways, e.g., to leaf cuticles, soil or water surfaces, was derived by the difference between g_c and g_s (Eq. 3).

The findings on g_{nsPAN} from the partitioning of $F_{\text{MBR}_{\text{PAN}}}$ were used to model PAN deposition fluxes for the entire period from 29 June to 21 October. Applying the resistive scheme given in Eq. (3), the modelled PAN flux ($F_{\text{model}_{\text{PAN}}}$) was derived as

$$F_{\text{model}_{\text{PAN}}} = - \left(R_a + R_{\text{d}_{\text{PAN}}} + \frac{1}{g_{\text{s}_{\text{PAN}}} + g_{\text{ns}_{\text{PAN}}}} \right)^{-1} \cdot \rho_m \cdot \chi_{\text{PAN}} \quad (4)$$

where R_a , $R_{\text{d}_{\text{PAN}}}$ and $g_{\text{s}_{\text{PAN}}}$ were determined as described above over the entire period. Here, ρ_m and χ_{PAN} represent the molar air density and the PAN mixing ratio, respectively, at the height of the eddy covariance measurements (z_{ref}).

2.5 Determination of PAN loss by thermochemical decomposition

Next to dry deposition process, other sink terms impact the measured surface PAN mixing ratios. While PAN photolysis and reaction with the hydroxyl radical (OH) are expected to be very low at altitudes below 7 km (Talukdar et al., 1995), thermochemical decomposition of PAN (back reaction of R1) has to be considered. Thermochemical decomposition of PAN increases exponentially with temperature and is more efficient at high NO/NO₂ ratios as PA reacts faster with NO than with NO₂ to reform PAN. Hence, the time scale of PAN towards thermochemical decomposition (τ_{chem}) is given by (Orlando et al., 1992; Shepson et al., 1992) as

$$\tau_{\text{chem}}(\text{PAN}) = \frac{1}{k_{1r}} \left(1 + \frac{k_{1f}[\text{NO}_2]}{k_2[\text{NO}]} \right) \quad (5)$$

To evaluate whether PAN loss by thermochemical decomposition significantly impacted the MBR fluxes (Sect. 2.2), the chemical flux divergence between z_{ref} and z_0 due to thermochemical decomposition of PAN ($F_{\text{chem}_{\text{surf}}}$) was determined according to Doskey

these periods occurred sometimes as very isolated events and were associated with an unstable boundary layer during daytime and a stable stratification during nighttime.

For the further evaluation, entire days were selected and classified according to wind speed and wind direction. In total 20 days were classified as low NO_x and 27 days as high NO_x conditions. The diurnal averages of the meteorological conditions and micrometeorological characteristics during these days are displayed in Fig. 2a–f and mixing ratios of O_3 , NO and NO_2 are shown in Fig. 2g–i. For both low and high NO_x conditions photolysis of NO_2 contributed to the steady increase of NO mixing ratios after sunrise, which peaked between 08:00 and 10:00 CET and then declined with the growth of the daytime boundary layer. During high NO_x conditions both advection of freshly emitted NO from nearby sources and generally higher NO_2 levels lead to high NO mixing ratios exceeding sometimes 10 ppb. Biogenic NO emission from the grassland ecosystem, determined with the dynamic chamber method, were found to be insignificant (Plake et al., 2014). NO_2 mixing ratios showed a high variability during high NO_x conditions also indicating local sources. The daytime NO_2 decline was caused by both dilution due to the growing boundary layer and photolysis. It was anti-correlated with the increase of O_3 mixing ratios. The development of a shallow nocturnal inversion layer during high NO_x conditions caused increased O_3 removal rates. As a result, nighttime O_3 mixing ratios were lower than during low NO_x conditions. During daytime, both the higher insolation and the presence of pollutants under high NO_x conditions resulted in higher O_3 mixing ratios during the afternoon.

3.2 Characterisation of PAN under low and high NO_x conditions

The diurnal cycle of PAN mixing ratios was closely linked to the diurnal cycle of O_3 . As for O_3 , PAN mixing ratios increase after dawn to the maximum in the afternoon, with median values of 300 ppt under low and of 600 ppt under high NO_x conditions, respectively (Fig. 2j). The maximum is followed by a steady decrease over night to

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median values just before dawn of about 50 ppt under low NO_x and 200 ppt under high NO_x conditions.

The major reason for the much higher PAN levels during high NO_x conditions, are the elevated NO₂ mixing ratios, which occurred especially during nighttime and declined with the onset of photolysis after dawn and the clearing of the nocturnal boundary layer. Comparing the diurnal evolution of PAN and O₃ mixing ratios, we find a higher PAN/O₃ ratio under high NO_x conditions at all times throughout the diurnal cycle. During peak PAN and O₃ mixing ratios in the afternoon, the PAN/O₃ ratio was 0.003 and 0.006 during low and high NO_x, conditions, respectively. Since photolytic production of O₃ from NO₂ was similar for both conditions, a large PAN/O₃ ratio implies a higher abundance of PA as a precursor of PAN (Zhang et al., 2009). Although no direct measurements of PA were available, the very low abundance of volatile organic compounds measured at the site (e.g., isoprene < 0.7 ppb, monoterpene < 0.3 ppb, J. Kesselmeier, personal communication, 2013) suggests that these higher levels of PA during high NO_x conditions primarily originated from anthropogenic non-methane hydrocarbons (NMHCs). Hence, PAN mixing ratios at the site were mainly influenced by advection from nearby pollution sources from north easterly directions.

The timescale for thermochemical decomposition of PAN, τ_{chem} , ranged for both low and high NO_x conditions mainly between 4 and 20 days at night (Fig. 2k). During daytime, τ_{PAN} ranged between 2 h and nearly one day (median ~ 5 h) for low NO_x conditions, but were significantly lower during high NO_x conditions (ranging between 30 min and 5 h; median ~ 2 h) caused by both on average higher NO/NO₂ ratios in the morning and higher temperatures in the afternoon.

3.3 Evaluation of PAN flux measurements

3.3.1 Deposition fluxes and canopy conductance

During the period of the PAN flux measurement mainly high NO_x conditions prevailed. The PAN fluxes showed a clear diurnal cycle with maximum deposition fluxes at midday

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and very small fluxes during nighttime (Fig. 3). Although the random flux errors were large compared to the observed fluxes (median $\pm 0.035 \text{ nmol m}^{-2} \text{ s}^{-1}$, see Moravek et al., 2014) a daytime PAN deposition was clearly visible on most days. The gaps in the time series are due to extended instrument calibrations and maintenance of the GC-ECD. For the further evaluation PAN fluxes below the flux detection limit (34 % of data, see Sect. 2.2 for definition) were neglected, aside from data where $u_* < 0.07 \text{ m s}^{-1}$ (28 % of data) as this criterion would have eliminated most of the nighttime values.

The diurnal median values of the PAN and O_3 fluxes are shown in Fig. 4a and b. A diurnal course of the PAN flux is observed with maximal deposition fluxes of $\sim -0.1 \text{ nmol m}^{-2} \text{ s}^{-1}$ during daytime, which corresponds to $v_{\text{D}_{\text{PAN}}}$ at z_{ref} of $\sim 0.3 \text{ cm s}^{-1}$. The visible short-term peaks are mostly attributed to the low number of data values ($\sum n = 255$) and also caused by the uncertainty of the MBR method. For O_3 , this feature was much less pronounced due to the higher number of data points used ($\sum n = 639$). Both measured PAN fluxes and $v_{\text{D}_{\text{PAN}}}$ values were between the observations by Wolfe et al. (2009) (midday averages $-0.04 \text{ nmol m}^{-2} \text{ s}^{-1}$; $v_{\text{D}} \approx 0.1 \text{ cm s}^{-1}$) and fluxes by Turnipseed et al. (2006) (midday averages $\sim -0.35 \text{ nmol m}^{-2} \text{ s}^{-1}$; $v_{\text{D}} \approx 1 \text{ cm s}^{-1}$) measured at two different pine forest sites in the USA during summer (Table 1). Daytime flux measurements at a grassland site by Doskey et al. (2004) resulted in an average $v_{\text{D}_{\text{PAN}}}$ of 0.13 cm s^{-1} . The magnitude of the daytime PAN flux at our site was about two orders of magnitude lower than the O_3 flux, yielding a median $v_{\text{D}_{\text{PAN}}}/v_{\text{D}_{\text{O}_3}}$ ratio of 1.03. Comparison with experimentally derived PAN fluxes in the past (Table 1) reveals that $v_{\text{D}_{\text{PAN}}}/v_{\text{D}_{\text{O}_3}}$ ratios vary considerably, which might be attributed to a large extent to the error of the applied measurement methods and the assumptions made. It has to be noted that v_{D} is height dependent, which can make its comparison between different studies difficult. However, the ratio $v_{\text{D}_{\text{PAN}}}/v_{\text{D}_{\text{O}_3}}$ is largely independent from height in case the vertical profiles of PAN and O_3 concentrations are similar as it was shown at least for the data from our site.

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The chemical flux divergence between z_{ref} and z_0 due to thermochemical decomposition of PAN (Eq. 6) was found to be very small with the highest median value of $0.007 \text{ nmol m}^{-2} \text{ s}^{-1}$ at noon (Fig. 4a). In contrast, for the O_3 flux, the loss term due to reaction with NO and the production by NO_2 photolysis were significantly higher between 06:00 and 11:00 CET and led to a small net production of O_3 during daytime, which was corrected for in the presented fluxes.

The overall canopy conductance for PAN (g_{cPAN}), representing the flux normalized by the concentration at z_0 , shows a mean diurnal cycle with its maximum during daytime (Fig. 4c and d). The midday median values were around 0.4 cm s^{-1} and were similar to g_{c} values observed for O_3 .

3.3.2 Stomatal uptake

During nighttime g_{sPAN} values were zero due to stomata closure (Fig. 4c). With the onset of radiation in the morning g_{sPAN} increases and reaches its maximum of 0.26 cm s^{-1} at 11:00 CET. As both g_{sPAN} and g_{sO_3} differ only by the PAN and O_3 diffusivities (see Sect. 2.4), they show the same pattern, while g_{sO_3} is larger by a factor of 1.6 due to the faster diffusivity of O_3 . Due to an increased vapour pressure deficit in the afternoon the maximum values of g_{sPAN} and g_{sO_3} are slightly skewed towards the morning.

The existence of a mesophyllic resistance limiting the stomatal uptake of PAN, as it was found by Teklemariam and Sparks (2004) or by Sparks et al. (2003) at high stomatal conductance, cannot be validated from our data. Only if the modelled g_{sPAN} values exceeded the experimentally determined g_{cPAN} values, a limitation could be suspected. It is suggested that the mesophyllic uptake of PAN is lower than for O_3 , as there are less reaction sites for PAN within the plant cell and its reaction with proteins is slower, although the mesophyll biochemistry for PAN assimilation is not clearly understood (Doskey et al., 2004).

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smallest root mean square error was obtained with an optimal g_{ns} value of 0.28 cm s^{-1} for the whole dataset, which corresponds to $R_{\text{nsPAN}} = 360 \text{ s m}^{-1}$. This resistance value is in the range of findings by other studies, e.g., Turnipseed et al. (2006) observed a median value of 244 s m^{-1} under dry and 125 s m^{-1} under wet conditions.

To obtain another independent estimate of the non-stomatal deposition, we employed the nocturnal boundary layer budget (NBLB) method according to Shepson et al. (1992), which compares the nocturnal PAN decay to that of O_3 and yields a ratio of PAN to O_3 deposition velocities. The main assumption, that the decline of both PAN and O_3 mixing ratio during nighttime was only due to deposition, is valid as chemical reactions are negligible at night for both PAN (see Fig. 2l) and O_3 (no photolysis and low NO mixing ratios). We analysed in total 16 nights, where a clear logarithmic decline of both PAN and O_3 mixing ratios was observed and where the main wind direction was within the “clean” sector in the south west. Using only nights with a good correlation between both PAN and O_3 decline ($R^2 > 0.7$), we obtain an average value for $v_{\text{DPAN}}/v_{\text{DO}_3}$ of 1.75, which is similar to the average value obtained by Shepson et al. (1992). As stomatal uptake is assumed to be zero at night, we obtained according to Eq. (3) a bulk g_{nsPAN} of 0.5 cm s^{-1} ($R_{\text{nsPAN}} = 200 \text{ s m}^{-1}$). This value of g_{nsPAN} is slightly larger than the value obtained from the MBR measurement.

In currently applied deposition models (e.g., Simpson et al., 2012), g_{nsPAN} is often derived according to Wesely (1989) as a composite between the non-stomatal conductance of sulphur dioxide (g_{nsSO_2}) and g_{nsO_3}

$$g_{\text{nsPAN}} = 10^{-5} \cdot H_{\text{PAN}}^* \cdot g_{\text{nsSO}_2} + f_{0\text{PAN}} \cdot g_{\text{nsO}_3} \quad (9)$$

SO_2 represents a very soluble substance (effective Henry constant $H_{\text{SO}_2}^* = 10^5 \text{ M atm}^{-1}$) and O_3 a compound that reacts fast with substances in the leave cuticles such as protein thiols (Mudd, 1982). Due to the poor solubility of PAN in water ($H_{\text{PAN}}^* = 4.1 \text{ M atm}^{-1}$, see Kames and Schurath, 1995) the first term of the right side of Eq. (9) can be neglected and only the reactivity index, f_0 , is of significant importance.

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According to Wesely (1989) $f_{0_{\text{PAN}}} = 0.1$, which suggests the non-stomatal deposition of PAN would be about one order of magnitude lower than for O_3 . Zhang et al. (2002) suggest a $f_{0_{\text{PAN}}} = 0.6$ based on first studies on PAN deposition by Hill (1971) and Garland (1977). This contradicts our findings by both the MBR and the NBLB method, which observed at least equal or even higher non-stomatal deposition for PAN than for O_3 , and supports the statement by Turnipseed et al. (2006) that current deposition models may significantly underestimate PAN non-stomatal deposition.

3.4 PAN deposition fluxes for low and high NO_x conditions

To evaluate the PAN deposition under both low and high NO_x conditions as well as its potential influence on the natural grassland ecosystem and its role for the atmospheric N_r budget, the PAN deposition flux was modelled for the entire period from 29 June to 21 October (see Sect. 2.4). For this, we used the bulk value for $g_{\text{ns}_{\text{PAN}}}$ of 0.28 cm s^{-1} (Sect. 3.3.3) for both low and high NO_x , as we found this to be the best estimate from our data. The obtained median diurnal cycles of $F_{\text{model}_{\text{PAN}}}$ for low and high NO_x conditions (Fig. 5) reveal that the total deposition (i.e. stomatal + non-stomatal) was more than twice as high during high NO_x ($\sim -0.1 \text{ nmol m}^{-2} \text{ s}^{-1}$) than during low NO_x ($\sim -0.05 \text{ nmol m}^{-2} \text{ s}^{-1}$) conditions, which is mainly attributed to the higher PAN mixing ratios during high NO_x conditions. Median midday deposition velocities were very similar during both episodes ($v_{\text{D}_{\text{PAN}}} \approx 0.5 \text{ cm s}^{-1}$). As already discussed in Sect. 3.3.3, the non-stomatal pathway was significant, which is reflected by a daytime fraction of $g_{\text{ns}_{\text{PAN}}}/g_{\text{c}_{\text{PAN}}}$ of 0.7 during low NO_x and 0.6 during high NO_x conditions. As about half of the grassland vegetation was senescing or was already dead, reaction on plant surfaces may be a reason for the large non-stomatal fraction.

The importance of PAN deposition as a loss process of PAN from the atmosphere is determined by comparison to the magnitude of the thermochemical decomposition of PAN in the boundary layer (Eq. 7). Due to the lower temperatures and the lack of NO at night, the nocturnal thermochemical loss was insignificant during both low and high

as well as detailed studies on the role of non-stomatal uptake mechanisms to improve current deposition models are desirable tasks for future research on PAN deposition.

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Table 1. Comparison of PAN fluxes and deposition velocities at the Mainz-Finthen grassland with previous studies. All field experiments were performed in the mid-latitudes of the Northern Hemisphere within the period of late spring to early autumn.

Method	Vegetation/ location	LAI (onesided)	Flux ($\text{nmol m}^{-2} \text{s}^{-1}$)		v_D (cm s^{-1})		$v_{D_{\text{PAN}}}/v_{D_{\text{O}_3}}$ (-)		Study
			day	night	day	night	day	night	
EC (CIMS)	pine forest	3.5	-0.35	-0.05	~ 0.7	~ 0.3	-	> 1	Turnipseed et al. (2006) Wolfe et al. (2009) ^a
	pine forest	5.1	-0.04	-0.03	0.30	0.10	1.25	0.5	
Gradient/MBR	corn field	-	-	-	-	0.54	-	-	Schrimpf et al. (1996) Doskey et al. (2004) ^b this study
	grassland grassland	- 4.8	- -0.07	- -0.01	- 0.30	- 0.06	- 1.03	- 1.25	
NBLB	n.s.	-	-	-	-	-	-	0.31	Garland and Penkett (1976) ^b Shepson et al. (1992) ^b McFadyen and Cape (1999) ^b this study (see Sect. 3.3.3)
	forest, agric.	-	-	-	-	-	-	2.38	
	rural, sea-side	-	-	-	-	-	-	1.1–6.2	
	grassland	4.8	-	-	-	-	-	1.75	
Laboratory	alfalfa	-	-	-	-	-	-	0.37	Hill (1971) Garland and Penkett (1976) Sparks et al. (2003) Teklemariam and Sparks (2004)
	grass	-	-	-	-	0.25	-	0.5	
	trees, crops	-	-0.02 to	-0.06	-	-	0.00–1.50	-	
	trees, herb.	-	-0.01 to	-0.02	-	-	0.11–0.34	-	

^a Net flux; in-canopy production of PAN was observed.

^b O_3 flux not was not measured directly.

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Table 2. Modelled PAN deposition and thermochemical loss for low and high NO_x conditions at the Mainz-Finthen grassland site. The description of the shown parameters is given in the text.

	PAN (ppt)	<i>T</i> (°C)	NO/NO ₂ (–)	<i>h</i> _{BL} (m)	τ_{dep} (d)	τ_{chemBL} (d)	k_{dep} (s ⁻¹)	k_{chemBL} (s ⁻¹)	L_{dep} ($\mu\text{g m}^{-2} \text{d}^{-1}$)	L_{chemBL} ($\mu\text{g m}^{-2} \text{d}^{-1}$)	L_{dep} (%)
Low NO _x											
daytime	182	17.6	0.18	567	1.50	0.88	1.8×10^{-4}	3.2×10^{-4}			
nighttime	147	13.6	0.01	200	0.42	16.08	6.7×10^{-4}	0.17×10^{-4}			
all									333	698	32
High NO _x											
daytime	405	20.0	0.23	641	1.54	0.54	1.8×10^{-4}	5.1×10^{-4}			
nighttime	334	15.9	0.01	200	0.83	12.67	3.3×10^{-4}	0.21×10^{-4}			
all									518	1840	22

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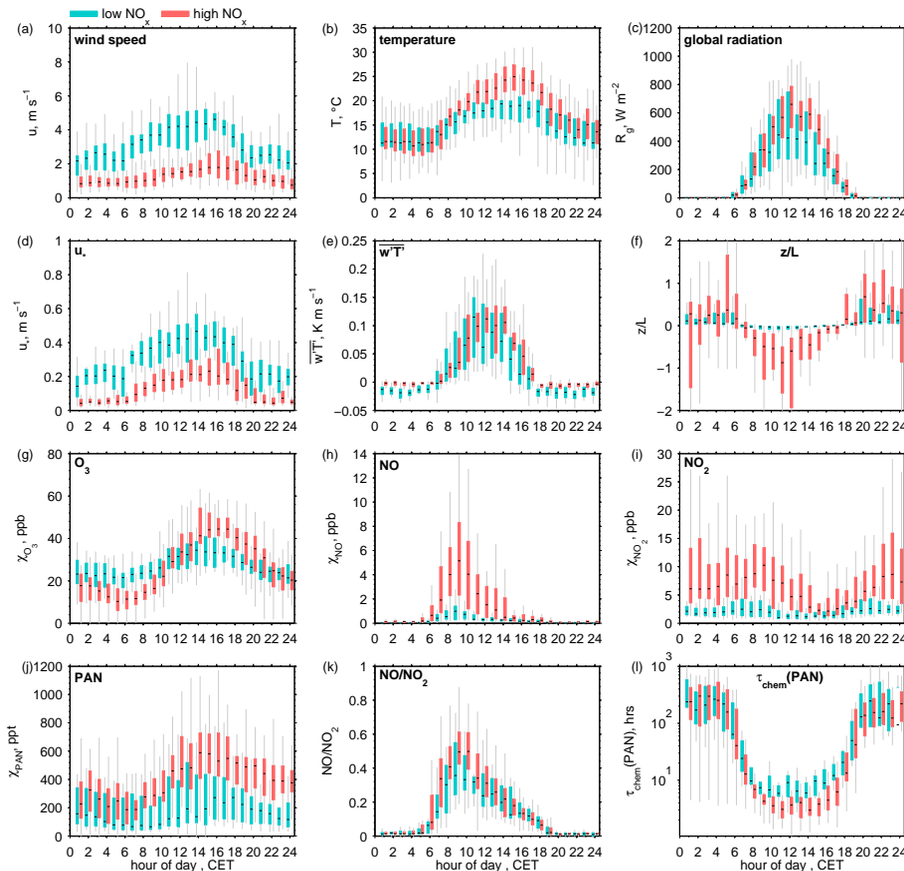


Figure 2. Diurnal boxplot statistics for the period from 29 June to 21 October 2011 at the Mainz-Finthen grassland site, characterising low and high NO_x periods according to the prevailing meteorological conditions (a–f), mixing ratios of the O_3 – NO – NO_2 triad (g–i) and PAN mixing ratios including the NO/NO_2 ratio used for the calculation of τ_{chem} (j–l).

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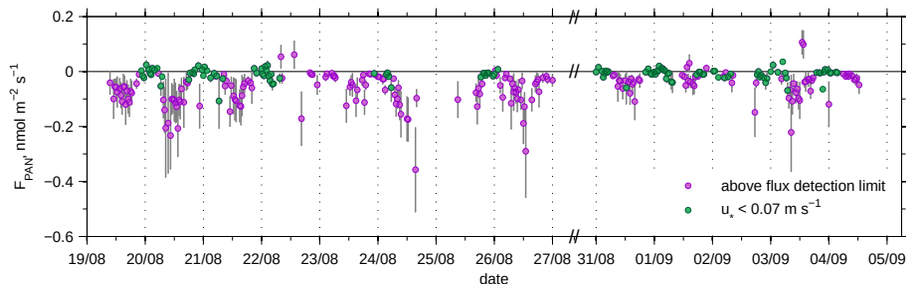


Figure 3. Overview of PAN flux measurements using the MBR method from 19 August to 4 September 2011 at the Mainz-Finthen grassland site after applying quality criteria as described in Moravek et al. (2014). Error bars represent the random flux error and green values indicate periods with weak turbulent exchange ($u_* < 0.07 \text{ m s}^{-1}$).

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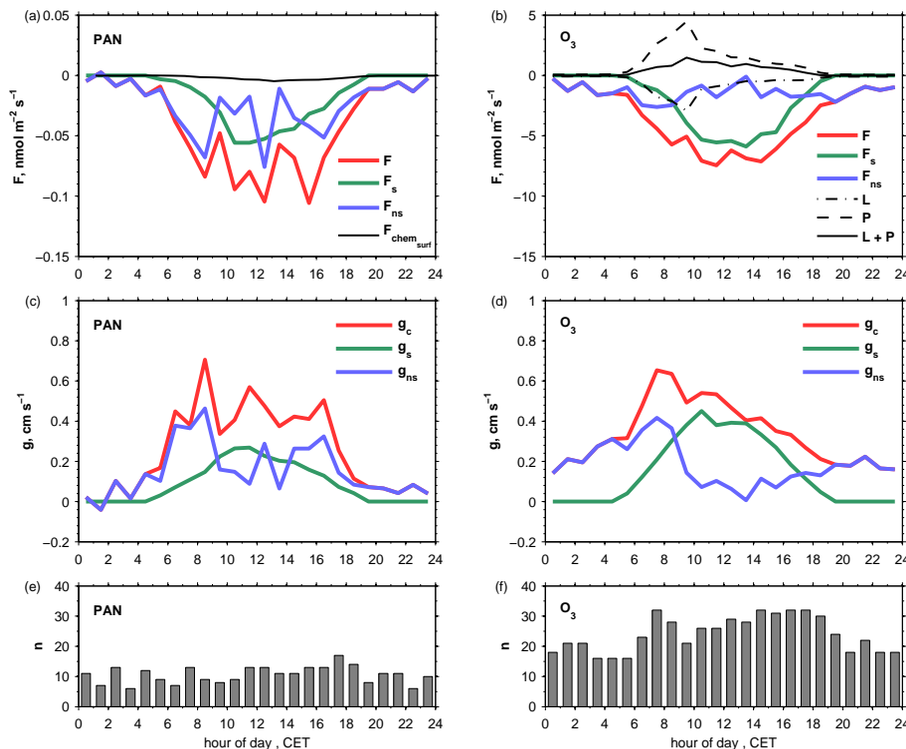


Figure 4. Flux partitioning of PAN (left column) and O₃ (right column) deposition into stomatal and non-stomatal pathways at the Mainz-Finthen grassland site. Shown are diurnal median values for the period from 19 August to 4 September 2011. Panels (a) and (b) indicate the deposition fluxes including the thermochemical flux term (F_{chem_surf}) for PAN and the loss (L) and production (P) terms for O₃. Panels (c) and (d) show the respective conductances, while (e) and (f) show the number data points used for every hourly interval.

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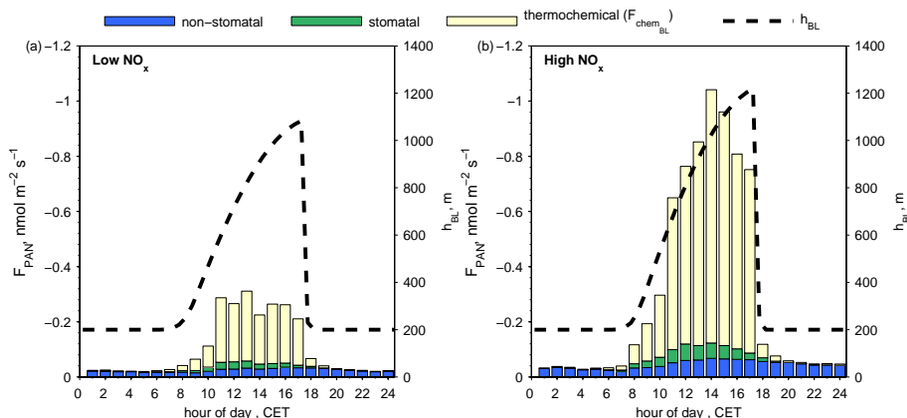


Figure 5. Modelled stomatal and non-stomatal PAN deposition fluxes and PAN loss due to thermochemical decomposition for (a) low and (b) high NO_x periods at the Mainz-Finthen grassland site. The dashed line marks the theoretical boundary layer height used for the calculation of the PAN decomposition (for details see text).

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