



Supplement of

NO₃ reactivity during a summer period in a temperate forest below and above the canopy

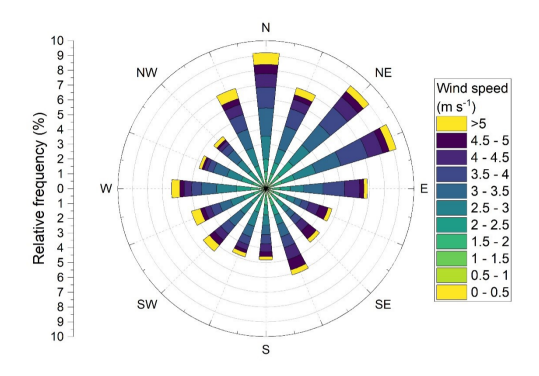
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Supplement

S1 Wind rose

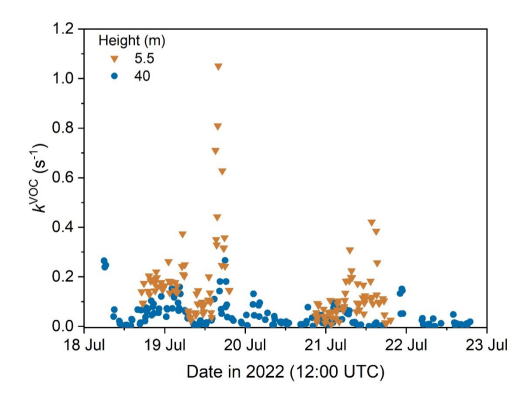


5 Figure S1: Wind rose coloured according to wind speed as measured at a height of 40 m during the ACROSS campaign.

S2 NO₃ reactivity measurements above and below the canopy

Figure S2 shows the time-series of k^{VOC} measured at 5.5 m (every odd hour) and at 40 m (every even hour) between July 18 and July 23. Note that the data coverage of k^{VOC} (5.5 m) is reduced due to missing NO measurements. During the daytime

- 10 period of July 21, k_{day}^{VOC} is about (0.05 ± 0.03) s⁻¹ at both heights. From ca. 17:00 UTC on, k^{VOC} (5.5 m) and k^{VOC} (40 m) diverge with an increase in both the magnitude and variability of the former and a continuous decrease in the latter to the instrument LOD of 0.006 s⁻¹ at ~22:30 UTC. On this night, there was a moderate temperature inversion ($\Delta T = -2.9$ K, Fig. 2) and the inlet at 40 m (ca. 20 m above canopy level) was thus effectively decoupled from any sources of biogenic emissions, explaining the low reactivity at this height. Furthermore, contrary to the daytime when the variability in k^{VOC} was similar at both heights, at
- 15 night k^{VOC} (40 m) is less affected by short-term fluctuations, presumably a result of enhanced mixing above canopy. It is also observed that $k^{\text{VOC}}(5.5 \text{ m})$ and $k^{\text{VOC}}(40 \text{ m})$ show a similar diel profile on July 19-20. However, k_{day}^{VOC} between the two heights differ by a factor of 2 and a weaker nocturnal gradient in k_{night}^{VOC} (only a factor of 4) occurs. The latter presumably results from a weaker, less stable temperature inversion ($\Delta T \approx -0.5 \text{ K}$, Fig. 2) during that night.



20 Figure S2: Time series of k^{VOC} (10 min averages) at 5.5 m (dark orange triangles) and 40 m (blue dots) between July 18 and July 23 during the ACROSS campaign. Major and minor ticks on the x-axis represent 12:00 and 00:00 UTC, respectively.

S3 Impact of OH, XO2 and heterogeneous reactions of N2O5 on NO3 loss

of 0.003 s⁻¹, found to be insignificant compared to photolysis and VOC-induced losses.

A chemical ionization mass spectrometer (CIMS) measured the sum of peroxy radicals ($XO_2 = RO_2 + HO_2$) as well as OH radicals ca. 3 m above ground level via their conversion to sulphuric acid (H_2SO_4) by sulphur dioxide (SO_2). Nitrate ions (NO_2^-)

serve as ionization agent. The associated measurement uncertainty of XO₂ is ca. 30 % (2σ) with an LOD of 2 x 10⁶ molecule 25 cm⁻³ (4 min). A detailed description of the instrument can be found in Kukui et al. (2008). IUPAC-recommended rate coefficients (IUPAC, 2024) were used to derive pseudo first-order NO₃ loss rate coefficients. In case of XO₂, the reaction rate coefficient of RO₂ was set to the value of HO₂ (4 x 10^{-12} cm³ molecule⁻¹ s⁻¹), since isoprene-derived RO₂ are suspected to be a factor of two more reactive (4.6 x 10^{-12} cm³ molecule⁻¹ s⁻¹) towards NO₃ (Dewald et al., 2020) than for instance ethyl peroxy radicals. Losses of NO₃ via its reaction with OH, HO₂ and RO₂ (Fig. S3) were, with a campaign-averaged mean contribution

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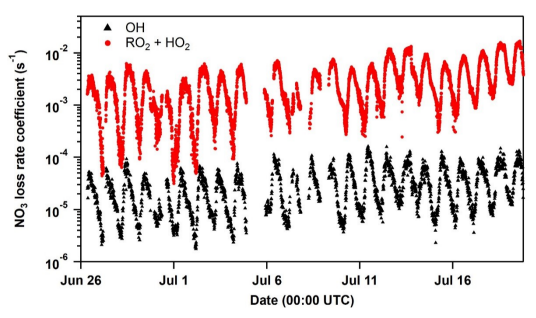
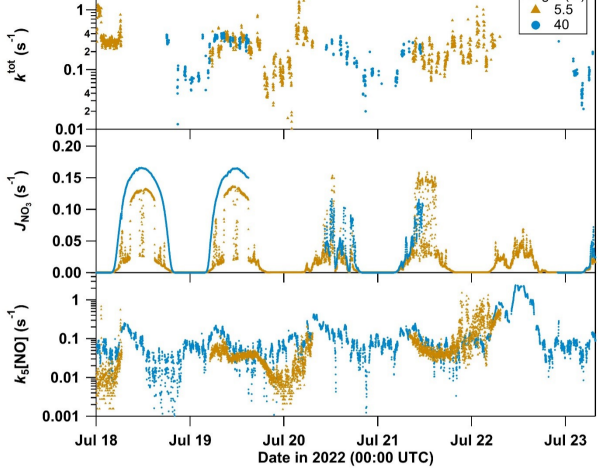


Figure S3: Time series of pseudo first-order NO₃ loss rate coefficients from OH (black) and RO₂ + HO₂ (red).

In addition, neither direct nor indirect heterogeneous losses (via N_2O_5) of NO₃ are expected to contribute significantly to the NO₃ loss term. Assuming a large uptake coefficient γ of 0.03 for N₂O₅ (featuring a mean molecular velocity c of 26233 cm s⁻ 35 ¹ at 298 K) and a surface area (A_S) of $1.5 \times 10^{-6} \text{ cm}^2 \text{ cm}^{-3}$ in a semi-rural environment (Phillips et al., 2016) together with the equilibrium constant (Keq) for reactions R3 and R4 of 2.8 x 10⁻¹¹ cm³ molecule⁻¹ at 298 K (IUPAC, 2024) would result in a k_{het} $= 0.25\gamma cA_s K_{eq} [NO_2] = 4 \times 10^{-4} s^{-1}$ in case of 2 ppby NO₂ at noon (see Fig. 1). Heterogeneous uptake is thus only of very minor importance below the canopy.

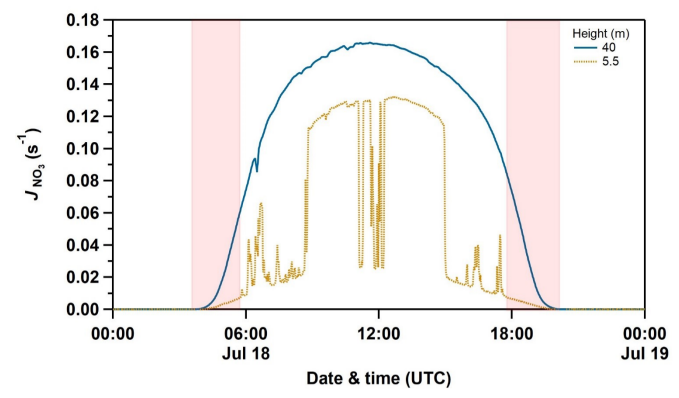
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S4 NO₃ loss rate coefficients (sum, NO, photolysis) above and below the canopy between July 18 and July 23

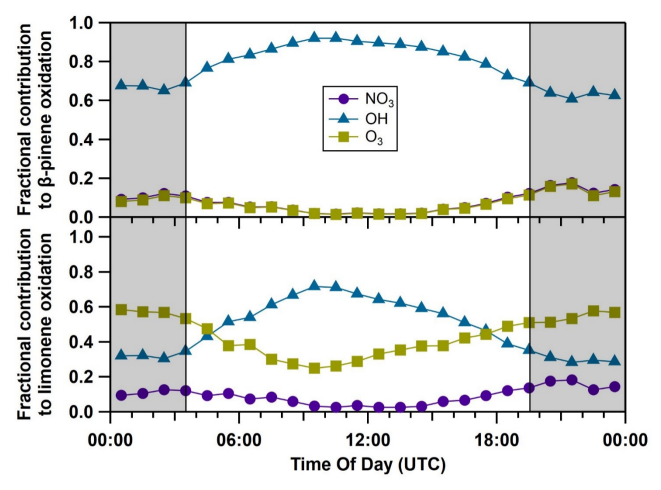


45 Figure S4: Time series of $k_5[NO]$, J_{NO_3} as well as k^{tot} at 5.5 m (dark orange dots) and 40 m (blue dots) between July 18 and July 23 during the ACROSS campaign. Major and minor ticks on the x-axis represent 00:00 UTC.





50 Figure S5: Exemplary diel profile of NO₃ photolysis frequencies (J_{NO_3}) measured in the clearing below the canopy (orange line, 5.5 m) versus those measured on top of the tower above the canopy (blue line, 40 m) during the ACROSS campaign on a clear day. The red shades mark periods in the early morning and the late afternoon when only diffuse sunlight reached the spectral radiometer below the canopy. As a consequence, $J_{NO_3}(5.5 \text{ m})$ was on average a factor of 12.5 lower than $J_{NO_3}(40 \text{ m})$ during these periods.



60 Figure S6: Median diel cycles (1h, circles) of fractional contributions of NO₃ (violet), OH (green) and O₃ (dark yellow) to the oxidation of β-pinene (upper panel) and limonene (lower panel) during the ACROSS campaign. The nighttime period is grey-shaded.

70 References

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