The following text contains the reviewer’s comments (black), our replies (blue) and the changes made to the manuscript (red).

**Reviewer 1**

This manuscript reports a careful lab study on the reactive quenching of electronically excited atmospheric NO2* and NO3* radicals by water vapour as potential primary sources of atmospheric HOx radicals. The topic is relevant for ACP.

The manuscript is divided into two parts. Firstly, the reactive quenching rate of NO2* by water vapour is re-investigated in order to elucidate the controversially discussed role of this photoreaction as a relevant primary source of atmospheric OH. From their results the authors inferred a new upper limit of 6x10^-6 (at an excitation wavelength of 562 nm) about an order of magnitude lower than previously measured. Extended measurements across the whole non-dissociative part of the absorption spectrum of NO2 lead to the conclusion that atmospheric OH formation by NO2* + H2O is insignificant.

Secondly, the reactive quenching of NO3* was studied. Although daytime concentration of NO3 are usually very low the authors estimated that under certain conditions the formation rate of OH by the title reaction could compete with the primary formation of OH from ozone photolysis. The measured quenching rate constant for the bath gas N2 compares well with the literature value of Nelson et al. The measured total quenching rate constant for H2O for which only one measurement is available in the literature was found to be a factor of four lower. In an elegant series of well-thought-out experiments it is shown that the majority of excited NO3* is deactivated by water vapour via radiationless processes and only an upper limit of <3% was found to react with water to form OH.

The experiments were conducted in a well-proven and tested apparatus. Concentrations of reactants are directly measured by UV/VIS absorption spectroscopy reducing the uncertainties to <10%. All relevant experimental details are reported, measurement uncertainties and experimental limitations are always well defined. Figures are meaningful and support the thorough and critical analysis and the discussion of the results.

I recommend the well written manuscript for publication once the following minor comments are addressed.

We thank the reviewer for this positive assessment of our manuscript.

Is there a deeper reason for the application of two different OH formation pathways (R13, R14) to convert the LIF raw signal to OH concentrations? In the experimental section both methods are described but it is pointed out later in the text (page 7, line 9) that in the presence of NO2 193 nm photolysis (R14) would produce small amounts of O1D atoms from NO2 in a two-photon process which finally rules out this reaction for the NO2* study.

On the other hand Figure 3 shows OH formation from the photolysis of water vapour. Why not from HNO3 photolysis as applied in the NO2* experiment? Is there a corresponding figure?

This point certainly needs some clarification and I would suggest to re-write the first paragraph of page 6 accordingly.

As evident in Table 1, both R13 and R14 can be (and were) used to calibrate the OH signal. The point we make is that use of 193 nm (H2O photolysis, R14) can not be used in the presence of NO2. In the first paragraph of page 6 we already explain this:

When using (R14) to calibrate the OH signal, the NO2 supply to the experiment was replaced with N2, and 193 nm light used to dissociate OH from the H2O already present (in unchanged conditions of pressure, temperature and [H2O]).

We have amended the caption to Figure 3 to remove any ambiguity:

The open circles are OH-calibrations obtained by the 193 nm photolysis of H2O (in the
absence of NO\textsubscript{2}) at different laser fluences (mJ cm\textsuperscript{-2}).

In order to enhance the concentration of excited NO\textsubscript{3}* radicals to a maximum of 50% the absorption transition at 662 nm was “saturated” using very high laser power intensities. How can saturation be experimentally verified and which laser fluence was applied? The laser fluences (in photons cm\textsuperscript{-2} per pulse) are listed in Table 2. These were used to calculate, via the Beer-Lambert law, which fraction of NO\textsubscript{3} would have been promoted to the excited state. If this exceeded 50%, the transition was considered saturated. No attempt was made to verify this experimentally.

The discussion of the potential OH formation rate from NO\textsubscript{3}* + H\textsubscript{2}O closes with the words “… it may still represent an important contribution to OH formation in some environments.” I am not happy with this general and more or less meaningless statement. It should be either substantiated or omitted.

We have substantiated this statement: …it is non-negligible compared to OH production rates from photolysis of O\textsubscript{3} (see section 1.2) and may still represent an important contribution to OH formation in environments where OH generation via traditional routes involving absorption of UV radiation is suppressed e.g. at high-latitudes in winter. We recognise that this statement is still very qualitative, but feel that a more detailed analysis (requiring global-scale model calculation of OH formation) is not presently warranted.

Page 3, line 30 – I calculate an excitation energy of 180.85 kJ/mol from the absorption of 662 nm photons. Hence, the given values of the reaction enthalpies require correction as well. The calculation of the reaction is infect correct even though the energy of 662 nm photon was wrongly listed. We have corrected the text and write:

Absorption of a 662 nm photon (the wavelength of maximum absorption by NO\textsubscript{3}, see Figure 1), provides an excitation energy of \( \approx 181 \text{ kJ mol}^{-1} \). Using compilations of enthalpies of formation (Wagman et al., 1982; Davis et al., 1993; Ruscic et al., 2004; Ruscic et al., 2005; Ruscic et al., 2006) we calculate that formation of radical products from NO\textsubscript{3}* is exothermic:

- by 110 kJ mol\textsuperscript{-1} for OH + HNO\textsubscript{3} (R12b)
- by 81 kJ mol\textsuperscript{-1} for HO\textsubscript{2} + HONO (R12c).

Page 5, line 25 – add type of light source, presumably low pressure Hg lamp
Information added. We now write:

…Hg-line at 184.95 nm,

Page 7, line 5 – add absorption cross section of H\textsubscript{2}O at 193 nm
Addition made. We write:

The OH concentration was calculated using a 193 nm cross section for H\textsubscript{2}O of \( 2.1 \times 10^{-21} \text{ cm}^2 \text{ molecule}^{-1} \) (Sander et al., 2011).

line 28 – typo: OH
Correction made

Page 9, line 1 – typo: excited state
Correction made

line 15 – typo: 1.6 x 10-10 / factor of four…
Correction made

Page 12 – reference Dillon et al.: two names in the list of co-authors are missing (Vereecken, Peeters)
Correction made

Page 17, Fig 3 – add concentration of H\textsubscript{2}O: 1.5x1017 molec cm\textsuperscript{-3} – correct ‘the the’
Correction made
**Reviewer 2**

The authors have carried out a thorough and comprehensive study and showed that there is no evidence that NO2 excited by visible light can react with H2O to form OH + HONO. The present study has lowered the upper limit for OH formation and implies that this reaction has no atmospheric impact. In addition, the possibility that the reaction between NO3 excited by visible light and H2O can form OH was investigated. No evidence for OH formation from this reaction was observed. However, the assigned upper OH yield for this reaction does not wholly rule it out from having some atmospheric impact. The other potential reactive channel HO2 + HONO is discussed but not investigated.

This paper is fine for publication with just a few minor corrections.

We thank the reviewer for this positive assessment of our manuscript.

**Reviewer 3**

The authors present new data that help to better constrain reactions of excited state NO2* and NO3* with H2O. These are challenging experiments to conduct and interpret, and the authors have done a great job. This is a very interesting paper and should be published after my suggestions below have been addressed.

We thank the reviewer for this positive assessment of our manuscript.

1 - Error bars /estimates.

Errors estimates should be added throughout the manuscript (especially to tables and graphs), if possible

The only Figure which can usefully present error bars is Figure 6. This already has error bars.

We now describe the source of these error bars in the caption:

The error bars are statistical uncertainty (2σ) from the fits to OH-decays as exemplified in Figure 5.

We feel we have adequately addressed sources of uncertainty when deriving (upper limits to) the yields we present. We write, for example:

The upper limits were calculated from the minimum observable OH-signal (assumed to be twice the RMS noise levels on the OH-signal) and accounting for uncertainty in parameters such as laser fluence (30%), NO2 concentration (10%) and concentration of H2O (10%).

And

The major sources of uncertainty in the calculated OH-yield are uncertainty in the measurement of laser fluences (30%) required to calculate the initial OH and NO3* concentrations and assumptions related to the (unmeasured) NO3 time profile.

2 - Manuscript organization.

The results and discussion section gives (too many) experimental details. For example, pg 6, line 25, begins with "A Nd-YAG pumped dye-laser was used to generate 532 and 567 - 647 nm light ... Reagent concentrations and conditions for these experiments ..." These are experimental details. Consider some re-organization (shifting of text) to improve the organization of the paper.

In the results and discussion section we have removed reference to the lasers

NO2 was excited at a number of different wavelengths, 532 and 567 - 647 nm; reagent concentrations and conditions for these experiments are given in Table 1

As several different experiment type using different excitation wavelengths, calibration schemes and conditions were used, we prefer to keep some experimental details in the results and discussions section (i.e. close to the results being discussed) rather than moving it to the experimental section.

3 - The authors expanded the wavelength regions over which the title reactions have been studied to above 532 nm and below 647 nm. It would be useful if the authors could comment on this chemistry occurring at lower wavelengths (in the intro and future work sections).
We see no real benefit in giving a more detailed description of chemistry at shorter excitation wavelengths and prefer to keep the discussion focused.

Our value of $k_{12}$ is more than a factor of four smaller than one previously reported value.

Most atmospheric OH is believed to be generated via a combination of primary photolytic processes involving e.g. $O_3$ ($\lambda \leq 370$ nm, (IUPAC, 2018)) (R1, R2) and HONO ($\lambda$: 280-370 nm, (IUPAC, 2018)).

We now refer only to 27°: (calculated using the TUV program (http://cprm.acom.ucar.edu/Models/TUV/Interactive_TUV/) for 50°N at a zenith angle of 27°, overhead $O_3$ column of 300 Du, a surface albedo of 0.1 and an aerosol optical depth of 0.235).

In experiments where both HNO$_3$ and H$_2$O were present, they were added sequentially so first the optical density due to a single component was measured before the second was added and the resultant total optical density monitored.

Pg 7 line 17. “it is immediately apparent, that our data are not consistent with”. Can you add error bars to make the line of reasoning more convincing?
As we present an upper limit, error bars on our data are not appropriate. The uncertainty is discussed in subsequent text. We now have removed “immediately apparent” and write:

Within experimental uncertainty (see below) our data are clearly not consistent with the large yield of OH reported by Li et al. (2008).

Pg 10 line 13 – replace “is” with “are”

Correction applied

Pg 11 discussion of HO2+HONO pathway, line 22 “Given that our experiments were blind to formation of HO2 or HONO”. This was very interesting to read, though speculative. It may be worth pointing that HO2 reacts with NO3, but (probably) too slowly to matter in these experiments.

Yes, reaction between these two radicals would not be a source of OH.

Pg 20 Figure 5. There are data points between the “2” and “3” line, and it is not clear what data set they belong to. Can you color-code the odd and even data differently, perhaps?

We have now used different symbol types to separate the individual decays.
Reactive quenching of electronically excited NO$_2^*$ and NO$_3^*$ by H$_2$O as potential sources of atmospheric HOx radicals

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Abstract. Pulsed laser excitation of NO$_2$ (532 – 647 nm) or NO$_3$ (623 - 662 nm) in the presence of H$_2$O was used to initiate the gas-phase reactions NO$_2^*$ + H$_2$O $\rightarrow$ products (R5) and NO$_3^*$ + H$_2$O $\rightarrow$ products (R12). No evidence for OH production in (R5) or (R12) was observed and upper-limits for OH production of $k_{5b}/k_5 < 1 \times 10^{-5}$ and $k_{12b}/k_{12} < 0.03$ were assigned. The upper limit for $k_{5b}/k_5$ renders this reaction insignificant as a source of OH in the atmosphere and extends the studies (Crowley and Carl, 1997; Carr et al., 2009; Amedro et al., 2011) which demonstrate that the previously reported large OH yield by (Li et al., 2008) was erroneous. The upper limit obtained for $k_{12b}/k_{12}$ indicates that non-reactive energy transfer is the dominant mechanism for (R12), though generation of small but significant amounts of atmospheric HOx and HONO cannot be ruled out. In the course of this work, rate coefficients for overall removal of NO$_3^*$ by N$_2$ (R10) and by H$_2$O (R12) were determined: $k_{10} = (2.1 \pm 0.1) \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ and $k_{12} = (1.6 \pm 0.3) \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$. Our value of $k_{12}$ is more than a factor of four smaller than the single, previously reported value.

1 Introduction

The capacity of the atmosphere to oxidise trace gases released at the Earth’s surface is sensitively dependent on the concentration of the hydroxyl radical, OH (Lelieveld et al., 2008). Most atmospheric OH is believed to be generated via a combination of primary photolytic processes involving e.g. O$_3$ ($\lambda \leq 370$ nm, (IUPAC, 2018)) (R1, R2) and HONO ($\lambda$: 280-370 nm, (IUPAC, 2018)) as well as in reaction of NO with HO$_2$, the latter being formed in the troposphere via the oxidative degradation of organic trace gases.

\[
\begin{align*}
O_3 + h\nu & \rightarrow O(^1D) + O_2 \quad \text{(R1)} \\
O(^1D) + H_2O & \rightarrow 2 \text{OH} \quad \text{(R2)} \\
\text{HONO} + h\nu & \rightarrow \text{OH} + \text{NO} \quad \text{(R3)}
\end{align*}
\]

As a large fraction of the oxidation of organic trace gases is initiated by reaction with OH, the conversion of HO$_2$ back to OH (e.g. via reaction with NO) is often referred to as recycling; the relative importance of direct OH formation and recycling depending on the concentrations of organics and NO. Together, OH and HO$_2$ are referred to as HOx.
Any reaction that can generate OH or HO₂ directly or indirectly (e.g. via generation of a short lived OH-precursor such as HONO), thus contributes to atmospheric oxidation capacity. Processes that form HONO (both gas-phase and heterogeneous) are therefore of great interest to atmospheric science and have been the subject of many studies (see e.g. (Stemmler et al., 2007; Li et al., 2014; Meusel et al., 2016). Two processes that may potentially generate HOₓ and HONO are the gas-phase reactions of H₂O with electronically excited nitrogen dioxide (NO₂ \( A \ ^2B_2 \) henceforth NO₂*) and electronically excited nitrate radical (NO₃ \( A \ ^2E^{''} \) and \( B \ ^2E^{'}, \) henceforth NO₃*).

1.1 NO₂* + H₂O

The potential for this reaction to generate both OH and HONO was first discussed and evaluated by Crowley and Carl (1997) who highlighted a possible role in increasing OH production rates in the weakly illuminated winter troposphere. It was argued that non-dissociative absorption by NO₂ (R4) could lead to formation of OH and HONO in a process (R5b) that is exothermic for excitation wavelengths across visible absorption spectrum of NO₂ which extends to \( \approx 650 \) nm.

\[
\begin{align*}
\text{NO}_2 + h\nu (\lesssim 650 \text{ nm}) & \rightarrow \text{NO}_2^* \quad \text{(R4)} \\
\text{NO}_2^* + \text{H}_2\text{O} & \rightarrow \text{NO}_2 + \text{H}_2\text{O} \quad \text{(R5a)} \\
\text{NO}_2^* + \text{H}_2\text{O} & \rightarrow \text{OH} + \text{HONO} \quad \text{(R5b)} \\
\text{NO}_2^* + \text{N}_2 / \text{O}_2 & \rightarrow \text{NO}_2 + \text{N}_2 / \text{O}_2 \quad \text{(R6)}
\end{align*}
\]

The rate of OH-formation following NO₂ excitation in the atmosphere depends on the OH yield (\( k_{5b} / k_5 \)) and on the relative rates of NO₂* deactivation by H₂O (R5) and by N₂ and O₂ (R6). For details of the NO₂ cross-sections, quantum yields, quenching rate constants and associated photo-physics for these processes we refer to our previous publication (Crowley and Carl, 1997).

Crowley and Carl (1997) used 532 nm pulsed-laser excitation of NO₂, to determine an upper limit to the OH-yield of (\( k_{5b} / k_5 \)) \( \leq 7 \times 10^{-5} \). Crowley and Carl (1997) also identified routes to O(\(^1\)D) at shorter wavelengths that involved two-photon excitation of NO₂, and which lead indirectly to OH formation via reaction of O(\(^1\)D) with H₂O. Whilst of some utility in the laboratory, such processes that require multi-photon excitation are generally of no consequence for the atmosphere.

More than ten years later, Li et al. (2008) carried out similar experiments but at longer wavelengths (560 – 640 nm) and reached very different conclusions, deriving a yield of OH (and thus also HONO) close to 1 \( \times 10^{-3} \), a factor of 14 times larger than the upper limit of Crowley and Carl (1997). Calculations of the impact of (R4-R5) using the large yield reported by Li et al. (2008) led to the conclusion that (R5) is important for air-quality under highly polluted conditions; use of the lower yield from Crowley and Carl (1997) resulted in minimal impact (Wennberg and Dabdub, 2008; Ensberg et al., 2010). Subsequent to the work of Li et al. (2008), two further experimental studies (Carr et al., 2009; Amedro et al., 2011) appeared to confirm the conclusions of Crowley and Carl (1997), and suggested that the high yield reported by (Li et al., 2008) was an experimental artefact, resulting from multi-photon laser-excitation of NO₂ in their focussed laser-beam (Amedro et al., 2011). However, the experiments of Amedro et al. (2011) at 565 nm and Carr et al. (2009) at 563.5 and 567.5 nm used NO₂*
prepared at wavelengths that covered only a small portion of the 560 – 630 nm range from Li et al. (2008). The single wavelength (532 nm) used by Crowley and Carl (1997), whilst interrogating the same excited states of NO2, was outside of the range of wavelengths covered by Li et al. (2008). The principal goal of the experiments on (R5) described in this work was therefore to measure OH yields ($k_{5b} / k_5$) using a range of photoexcitation wavelengths similar to those employed by Li et al. (2008) but avoiding potential complications related to multi-photon excitation.

1.2 NO$_3$* + H$_2$O

The NO$_3$ radical is generated throughout the atmospheric diel cycle via the oxidation of NO$_2$ by O$_3$:

$$\text{NO}_2 + \text{O}_3 \rightarrow \text{NO}_3 + \text{O}_2 \quad (R7)$$

At night, NO$_3$ can acquire mixing ratios of 100s of pptv. The high-reactivity of NO$_3$ towards unsaturated, organic trace gases (especially biogenically emitted ones in forested regions (Liebmann et al., 2018a; Liebmann et al., 2018b)) make it an important nocturnal oxidant. NO$_3$ is generally considered to be unimportant during daytime due to rapid photolysis. Rapid photodissociation (R8a & R8b) following absorption of visible light, reduces the daytime NO$_3$ lifetime to only a few seconds, and usually limits mixing ratios to less than 1 pptv.

$$\text{NO}_3 + h\nu \rightarrow \text{NO}_2 + \text{O}(^3\text{P}) \quad (R8a)$$
$$\text{NO}_3 + h\nu \rightarrow \text{NO} + \text{O}_2 \quad (R8b)$$

NO$_3$ photo-physics has been the subject of many studies, up to 1991 reviewed by (Wayne et al., 1991). Briefly, the NO$_3$ absorption spectrum ($\approx 400 – 665$ nm) is broad and diffuse with an extended excited-state lifetime of several hundred µs (Nelson et al., 1983) for excitation beyond the photo-dissociation limit. The extended lifetime results from coupling between ro-vibrational levels of the ground ($X^2\text{A}_2$) state and the excited ($A^2\text{E}^-'$ and $B^2\text{E}^-'$) electronic states, so that excitation into the strongest feature (centred at $\approx 662$ nm) can be considered to populate a manifold of mixed ground and excited electronic states (Carter et al., 1996). For simplicity, we refer to excited state NO$_3$ as NO$_3$*.

NO$_3$* can dissociate (R8, dominant at excitation wavelengths < 630 nm), fluoresce (R9) and return to the ground-state or be quenched in collisions with the main atmospheric bath gases N$_2$, O$_2$ and H$_2$O (R10-R12). Fluorescence and collisional quenching are important only at wavelengths longer than $\approx 630$ nm.

$$\text{NO}_3^* \rightarrow \text{NO}_3 + h\nu \quad (R9)$$
$$\text{NO}_3^* + \text{N}_2 \rightarrow \text{NO}_3 + \text{N}_2^# \quad (R10)$$
$$\text{NO}_3^* + \text{O}_2 \rightarrow \text{NO}_3 + \text{O}_2^# \quad (R11)$$
$$\text{NO}_3^* + \text{H}_2\text{O} \rightarrow \text{NO}_3 + \text{H}_2\text{O}^# \quad (R12a)$$
$$\rightarrow \text{OH} + \text{HNO}_3 \quad (R12b)$$
$$\rightarrow \text{HO}_2 + \text{HONO} \quad (R12c)$$

where # denotes formation of vibrationally hot products following energy transfer from NO$_3$*. Absorption of a 662 nm photon (the wavelength of maximum absorption by NO$_3$, see Figure 1), provides an excitation energy of $\approx 181$ kJ mol$^{-1}$. 


Using compilations of enthalpies of formation (Wagman et al., 1982; Davis et al., 1993; Ruscic et al., 2004; Ruscic et al., 2005; Ruscic et al., 2006) we calculate that formation of radical products from NO$_3^*$ is exothermic: by 110 kJ mol$^{-1}$ for OH + HNO$_3$ (R12b) and by 81 kJ mol$^{-1}$ for HO$_2$ + HONO (R12c).

The net result of NO$_3$ formation in (R7) and photolysis via the main channel, (R8a), is no change in NO$_X$ (NO$_X$ = NO + NO$_2$) or O$_3$. The net effect of formation in (R7) and photolysis via the minor (20%) channel (R8b) is conversion of NO$_2$ to NO (i.e. no net loss of NO$_X$) and conversion of O$_3$ to O$_2$ (loss of odd-oxygen). Reaction of NO$_3^*$ with H$_2$O to form OH + HNO$_3$ (R12b) changes this picture dramatically. As illustrated in Figure 2, if NO$_3^*$ reacts with H$_2$O to form OH + HNO$_3$ (R12b), the net effect is conversion of NO$_2$ to HNO$_3$ (i.e. loss of NO$_X$) and conversion of O$_3$ and H$_2$O to OH. This process (R7, R12b) therefore allows formation of atmospheric OH from O$_3$ in the absence of actinic UV radiation normally required to generate O(¹D) from O$_3$ (R1). If NO$_3^*$ reacts with H$_2$O to form OH + HONO + O$_2$, as in (R12c), the net effect is conversion of NO$_2$ to NO (no loss of NO$_X$) and formation of two HO$_X$ molecules, again bypassing the need for the actinic radiation in the UV.

Using literature values for the wavelength dependent NO$_3$ absorption cross-sections (Yokelson et al., 1994) and photolysis quantum yields (Orlando et al., 1993) as well as actinic flux (calculated using the TUV program ([http://cprm.acom.ucar.edu/Models/TUV/Interactive_TUV/] for 50 °N at a zenith angle of 27°, overhead O$_3$ column of 300 Du, a surface albedo of 0.1 and an aerosol optical depth of 0.235) we calculate that, averaged over the NO$_3$ absorption spectrum, 60% of actinic photons absorbed result in dissociation of NO$_3$. The residual 40% results in formation of NO$_3^*$ which can then undergo chemical and photo-physical transformation. Figure 1 gives an example of the relative rates of photo-dissociation and (non-dissociative) photo-excitation across the NO$_3$ absorption spectrum.

The relative importance of fluorescence and the collisional deactivation processes depends on the fluorescence lifetime and the rate constants for quenching. Nelson et al. (1983) report two components to the NO$_3$ fluorescence decay they observed following excitation at 661.9 nm, with collision-free fluorescence-lifetimes of 27 and 340 µs.

The longer lived component (accounting for > 85% of the total fluorescence) was quenched by N$_2$ and O$_2$ with rate coefficients of $k_{10} = (1.7 \pm 0.2) \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ and $k_{11} = (2.1 \pm 0.02) \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, respectively. Nelson et al. (1983) did not report a quenching rate coefficient for H$_2$O, but determined large quenching coefficients for propane ($1.09 \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$) and nitric acid ($3.07 \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$), presumably resulting from more efficient energy transfer due to higher densities of states in these polyatomics. A substantially larger rate coefficient for quenching of NO$_3^*$ by H$_2$O of $k_{12} = (6.9 \pm 0.5) \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$) was reported by Fenter and Rossi (1997). The quenching rate constants are sufficiently large that, at the pressures of N$_2$, O$_2$ and H$_2$O available in the troposphere, relaxation of NO$_3^*$ via fluorescence can be neglected.

The fraction, $f_{H_2O}$, of tropospheric NO$_3^*$ that will be quenched by collision with H$_2$O rather than N$_2$ or O$_2$ is given by expression (1):

$$f_{H_2O} = k_{12}[H_2O] / (k_{12}[H_2O] + k_{10}[N_2] + k_{11}[O_2])$$  \hspace{1cm} (1)
Using this expression we calculate that, at the Earth’s surface (1 bar pressure) and a temperature of 25 °C, \( f_{\text{H}_2\text{O}} \) can vary between 0.2 and 0.5 for relative humidities between 20 and 80%. As mentioned above, daytime concentrations of NO\(_3\) are generally low due to rapid photolysis (and reaction with NO) though measurements in polluted environments indicate maximum daytime concentrations of [NO\(_3\)] \( \approx 1 \times 10^8 \) molecule cm\(^{-3}\) (Geyer et al., 2003). The atmospheric production rate of OH via NO\(_3\) excitation may be written:

\[
P_{\text{OH}}(\text{NO}_3^*) = J_{\text{exci}}[\text{NO}_3] f_{\text{H}_2\text{O}}
\]

Using an NO\(_3\) concentration \( 1 \times 10^8 \) molecule cm\(^{-3}\) and \( J_{\text{exci}} = 0.15 \) s\(^{-1}\) (Figure 1) enables us to calculate an OH production rate (at 80% relative humidity) of \( 7.5 \times 10^6 \) molecule cm\(^{-3}\) s\(^{-1}\) if all quenching of NO\(_3^*\) by H\(_2\)O is reactive and forms OH. To put this value in context, we note that typical OH production rates from photolysis of O\(_3\) are around \( 2 \times 10^5 \) molecule cm\(^{-3}\) s\(^{-1}\), a factor of \( \approx 40 \) lower. The principal objective of this work was therefore to determine the OH production rate via NO\(_3\) photoexcitation and subsequent reaction of NO\(_3^*\) with H\(_2\)O (R12b). To best constrain these measurements, rate coefficients for total removal (quenching and chemical reaction) of NO\(_3^*\) by H\(_2\)O (\( k_{12} \)) and N\(_2\) (\( k_{10} \)) were determined.

### 2 Experimental

All experiments were conducted in a 500 cm\(^3\) jacketed photolysis cell as described previously (Wollenhaupt et al., 2000; Dillon et al., 2006) Laser light entered and exited the reaction vessel via Brewster-angle quartz-windows; laser-fluence at each wavelength being recorded using a Joule-meter located behind the exit-window. An excimer laser was used to generated \( \approx 20 \) ns pulses of light at 193 nm (ArF) or 248 nm (KrF). Dye-lasers pumped by Nd-YAG lasers were used to generate pulsed (\( \approx 6 \) ns) tuneable radiation at visible wavelengths. The pressure and the gas flow rate (300 - 2000 cm\(^3\) (STP) min\(^{-1}\)) were regulated to ensure that a fresh gas sample was available for each laser pulse for operation at 10 Hz. The pulsed laser-based schemes for generation of excited NO\(_2\) and NO\(_3\) are described below, as are the schemes for calibration of the OH signal.

Concentrations of the key reactants and precursors (NO\(_2\), HNO\(_3\) and H\(_2\)O) were monitored by UV-vis. absorption spectroscopy, reducing potential uncertainties in each of these parameters to \( \leq 10\% \). NO\(_2\) was measured in-situ using a multi-pass absorption cell positioned upstream of the reactor. Light from a halogen lamp passing through the cell was focused onto the entrance slit of a 0.5 m monochromator. A diode-array detector was used to record NO\(_2\) absorption in the visible range of light between 398 \( \leq \lambda \leq 480 \) nm at an instrumental resolution of 0.32 nm, determined from the full width at half maximum (FWHM) of the 436.8 nm Hg emission line. Optical absorption by HNO\(_3\) and H\(_2\)O was determined using a “dual beam” absorption cell (Hg-line at 184.95 nm, \( l = 43.8 \) nm) located downstream of the photolysis reactor. In experiments where both HNO\(_3\) and H\(_2\)O were present, they were added sequentially so first the optical density due to a single component was measured before the second was added and the resultant total optical density monitored.
NO₂ concentrations were calculated using a literature reference spectrum (Vandaele et al., 1998). Concentrations of HNO₃ and H₂O were calculated using cross-sections of 1.61 × 10⁻¹⁷ cm² molecule⁻¹ (Dulitz et al., 2018) and 7.22 × 10⁻²⁰ cm² molecule⁻¹ (Creasey et al., 2000).

The output from a Nd-YAG pumped dye-laser operating with Rhodamine 6G dye was frequency doubled to 282 nm and used to detect OH via excitation the A²Σ (v = 1) ← X²Π (v = 0) transition close to 282 nm. Laser-induced fluorescence was detected by a photomultiplier tube shielded by a combination of a 309 nm (± 5 nm) interference filter and BG 26 (glass) filter. Directly following experiments to measure formation of OH in the title reactions, known amounts of OH were generated via pulsed laser photolysis of HNO₃ (at 248 nm) or H₂O (at 193 nm).

$$\text{HNO}_3 + h\nu \ (248 \text{ nm}) \rightarrow \text{OH} + \text{NO}_2 \quad \text{(R13)}$$

$$\text{H}_2\text{O} + h\nu \ (193 \text{ nm}) \rightarrow \text{OH} + \text{H} \quad \text{(R14)}$$

For the experiments on NO₂⁺, a small flow of HNO₃ diluted in N₂ was added to the N₂ bath and a series of experiments was conducted that covered a range of laser fluences at 248 nm. The additional flow was compensated by reducing the main N₂ flow so that different concentrations of OH were generated in essentially unchanged conditions of pressure, temperature [NO₂] and [H₂O]. When using (R14) to calibrate the OH signal, the NO₂ supply to the experiment was replaced with N₂, and 193 nm light used to dissociate OH from the H₂O already present (in unchanged conditions of pressure, temperature and [H₂O]).

The uncertainty associated with conversion of LIF signals into OH concentrations stemmed partially from uncertainties in (measured) [HNO₃] and [H₂O] but was dominated by uncertainty in the measurement of the laser fluence at the centre of the reactor. Such measurements depended on both the accuracy of the Joule-meter and corrections for beam divergence and the assumption of a homogeneous light intensity over the cross-section of the laser beam. An overall uncertainty of 40% was estimated for the conversion of LIF-signals to absolute [OH] required for \((k_{3b} / k_3)\) determinations. For determination of \(k_{12b} / k_{12}\), the self-calibrating chemistry (R13, R15) results in a smaller contribution of laser fluence uncertainty to the overall uncertainty which is dominated by assumptions regarding the NO₃ profile (see later).

Chemicals: NO₂ (ABCR 99.99%) was subject to repeated freeze-pump-thaw cycles at 77 K prior to dilution in N₂ and storage in blackened glass bulbs; H₂O (“milli-Q” de-ionised water) and HNO₃ (prepared in house from H₂SO₄ + KNO₃) were added to the reactor via bubblers; O₃ was generated via electric discharge through O₂ in a commercial ozoniser (Anseros); N₂O₅ was prepared by mixing O₃ with NO₂ and trapping the resulting N₂O₅ at 195 K (Wagner et al., 2008); N₂ and O₂ (Westfalen, 99.999%) were used as supplied.
3 Results and Discussion

3.1 NO$_2^*$ + H$_2$O (R5)

NO$_2$ was excited at a number of different wavelengths, 532 and 567 - 647 nm; reagent concentrations and conditions for these experiments are given in Table 1. In general, large concentrations of H$_2$O were used to promote reaction of NO$_2^*$ over deactivation by other colliders, notably N$_2$, and to ensure that changes in other reagent concentrations (e.g. for calibration, see above) had a minimal effect on fluorescence quenching or other processes that impact on OH-LIF detection sensitivity.

Figure 3 displays the results of an experiment in which NO$_2$ was excited at 532 nm (at $t = 280 \mu$s) to generate $10^{13}$ to $10^{14}$ molecule cm$^{-3}$ of NO$_2^*$. The delay of 280 $\mu$s is the time between the triggering of the flash-lamps (at $t = 0$) and the Q-switch of the YAG-laser. The solid black triangles were obtained with the OH-excitation laser tuned to 282 nm (on resonance) and indicate a change in signal $\approx 200$ to 350 $\mu$s. This signal does not display the kinetic behaviour of OH in this chemical environment and remains when the OH-excitation laser is tuned off resonance (red triangles). It is also present when the 532 nm light is blocked and we conclude that this weak signal, having neither kinetics or spectroscopy characteristic of OH, is an artefact with electronic origin, possibly related to the output of the pulse generator used to trigger the laser Q-switch.

The data represented by open circles (roughly independent of reaction time) are the results of OH-calibration experiments using the 193 nm photolysis of H$_2$O ($1.5 \times 10^{17}$ molecule cm$^{-3}$) at four laser fluences between 0.3 and 6.8 mJ cm$^{-2}$ in the absence of NO$_2$. The OH concentration was calculated using a 193 nm cross section for H$_2$O of $2.1 \times 10^{-21}$ cm$^2$ molecule$^{-1}$ (Sander et al., 2011).

The roughly constant OH level over 1000 $\mu$s is consistent with the fact that OH does not react with any components of the gas-mixture. An experiment at 193 nm using the same OH-generation scheme but in the presence of NO$_2$ is displayed as solid stars. OH now decays exponentially at a rate which is consistent with its loss via reaction with NO$_2$. In this experiment, some OH was also generated by the reaction of O(1D) (formed by the 193 nm photolysis of NO$_2$) with H$_2$O and it was not used for calibration purposes. The signals obtained in the absence of NO$_2$ were converted to OH concentrations (right y-axis) using Joule meter readings as described in section 2.1.

The solid black line in Figure 3 represents the OH-signal and concentration expected from our experimental conditions (NO$_2$ concentration, H$_2$O concentration, total pressure and 532 nm laser fluence) and literature data for NO$_2$ absorption cross-sections, NO$_2^*$ deactivation rate constants and the yield of OH from NO$_2^*$ + H$_2$O reported by Li et al. (2008). Within experimental uncertainty (see below) our data are clearly not consistent with the large yield of OH reported by Li et al. (2008). In order to rule out the possibility that this is a result of using different excitation wavelengths, similar experiments were carried out in which we explored different regions of the NO$_2$ absorption spectrum. OH signals were not observed at any wavelength, enabling us to set upper limits to $k_{5b} / k_5$. The upper limits were calculated from the minimum observable OH-signal (assumed to be twice the RMS noise levels on the OH-signal) and accounting for uncertainty in parameters such as laser fluence (30%), NO$_2$ concentration (10%) and concentration of H$_2$O (10%).

7
The results are summarised in Table 1 which lists the experimental conditions in detail and in Figure 4 where we also compare to literature determinations of $k_{5b} / k_5$. The present dataset and those reported by Crowley and Carl (1997), Amedro et al. (2011) and Carr et al. (2009) found OH formation in the reaction between NO$_2^*$ and H$_2$O to be inefficient, with upper limits to $k_{5b} / k_5$ of between $6 \times 10^{-6}$ and $1.4 \times 10^{-4}$ at all wavelengths investigated. Together, these datasets contradict the yield of $1 \times 10^{-3}$ reported by Li et al (2008) for excitation across the wavelength range 560 to 630 nm. Our dataset, covering three absorption features of the NO$_2$ absorption spectrum within the range reported by Li et al. (2008) also rules out that the poor agreement is due to use of different excitation wavelengths. As discussed by Amedro et al. (2011) the use of focussed laser beams and resulting multi-photon processes are the most likely explanation for OH formation in the work of Li et al (2008). The results from this work reduce the maximum yield of OH from the reaction of NO$_2^*$ with H$_2$O to $6 \times 10^{-6}$ at 532 nm as opposed to $7 \times 10^{-5}$ measured by Crowley and Carl (1997). The assumption that this value is valid across the non-dissociative part of the absorption spectrum of NO$_2$, enables us to conclude that formation of atmospheric OH (and HONO) via R5b is insignificant.

### 3.2 NO$_3^*$ + H$_2$O (R12)

#### 3.2.1 Generation of NO$_3$

For the experiments to investigate the reaction of NO$_3^*$ with H$_2$O (R12), NO$_3$ was generated via the reaction of OH with known amounts of HNO$_3$ (R15).

$$\text{OH} + \text{HNO}_3 \rightarrow \text{NO}_3 + \text{H}_2\text{O} \quad (\text{R15})$$

The rate constant and the yield of NO$_3$ (unity) for (R15) are well known (Brown et al., 1999; Brown et al., 2001; Carl et al., 2001; Dulitz et al., 2018) enabling the time-dependent NO$_3$ concentration profile to be calculated if the initial amount of OH is known. This initial concentration of OH depends on the 248 nm laser-fluence (measured by Joule-meter, uncertainty 30%) and the HNO$_3$ concentration (measured by optical absorption at 185 nm, uncertainty 10%). As OH was formed from HNO$_3$ photolysis (R13), and the OH decay monitored, these experiments were self-calibrating as long as a sufficient excess of $[\text{HNO}_3] >> [\text{OH}] \approx [\text{NO}_3]$ was maintained. In the conditions employed in this work (see Table 2), radical loses via unwanted self- and cross-reactions of OH and NO$_3$ were < 5% of the total OH loss rate which was dominated by (R15). In experiments to measure the rate constant for NO$_3^*$ quenching by N$_2$ (R10) and H$_2$O (R12), NO$_3$ was generated via the 248 nm photolysis of N$_2$O$_5$:

$$\text{N}_2\text{O}_5 + hv \ (248 \text{ nm}) \rightarrow \text{NO}_3 + \text{NO}_2 \quad (\text{R16})$$

In this scheme of NO$_3$ generation NO$_3$ is formed instantaneously (in contrast to the reactions (R13 and R14).

#### 3.2.2 Quenching of NO$_3^*$ by N$_2$ and H$_2$O ($k_{10}$ and $k_{12}$)

The fate of electronically excited NO$_3$ radicals in the atmosphere is controlled by the relative rate of quenching by H$_2$O and the predominant bath-gases N$_2$ and O$_2$, which depends both on the concentration of H$_2$O and on the quenching rate
coefficients $k_{10}$, $k_{11}$ and $k_{12}$. As the rate constant for quenching of NO$_3^*$ by H$_2$O ($k_{12}$) has been addressed only briefly in a single study (Fenter and Rossi, 1997) and the value derived ($6.9 \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$) is unexpectedly large, we chose to re-measure $k_{12}$. In these experiments, NO$_3$ was generated in (R16) and He was used as the main bath gas, with traces of N$_2$ and H$_2$O added.  

An excitation laser-pulse at 662 nm was triggered when the NO$_3$ concentration was close to its maximum value (i.e. when > 95% of the primary-OH had been consumed by reaction with HNO$_3$) to generate NO$_3^*$. Time dependent fluorescence from NO$_3^*$ ($\lambda > 690$ nm) was detected using a red-sensitive photo-multiplier and recorded on a 100 MHz, digital oscilloscope. Fluorescence decay constants in the presence of various concentrations of H$_2$O were then used to derive $k_{12}$. We also conducted a set of experiments using N$_2$ as quenching molecule to test our experimental methodology by comparison with literature measurements of $k_{10}$.

NO$_3$ fluorescence profiles from these experiments are displayed in Figure 5, where datasets are depicted in which various amounts of N$_2$ (Fig. 5a) and H$_2$O (Fig. 5b) were added to the He bath gas. The fluorescence decay rate constant ($k'_{f}$) derives from the sum of processes that depopulate the excited state and includes fluorescence, inter-system crossing as well as quenching by N$_2$, H$_2$O and N$_2$O$_5$ with rate constants ($k_{f}$, $k_{isc}$, $k_{f}(N_2)$, $k_{f}(H_2O)$ and $k_{f}(N_2O_5)$, respectively.

$$k'_{f} = k_{f} + k_{isc} + k_{q}(N_2)[N_2] + k_{q}(H_2O)[H_2O] + k_{q}(N_2O_5)[N_2O_5]$$

In line with previous studies Nelson et al. (1983), the slow component of the NO$_3$ fluorescence was found to decay mono-exponentially (black and red lines in Figs. 5a and 5b) and depended on the pressure of N$_2$ or H$_2$O.

The decay constant ($k'_{f}$) was derived from exponential fits to the data and plotted against the concentration of N$_2$ or H$_2$O (Figure 6) to obtain (from the slopes) the rate constants $k_{10}$ and $k_{12}$ for quenching by N$_2$ and H$_2$O, respectively. Assuming negligible contribution from OH, NO$_2$ and NO$_3$, due to their low concentrations, the y-axis intercepts in Fig. 6 ($\approx 0.5-0.8 \times 10^6$ s$^{-1}$) are the combined terms $k_{f} + k_{isc} + k_{q}(N_2O_3)[N_2O_3]$, where $k_{q}(N_2O_5)$ is the unknown rate constant for quenching of NO$_3^*$ by N$_2$O$_5$. As the collision-free lifetime of excited NO$_3$ is several hundred µs, the terms $k_{f}$ and $k_{isc}$ contribute insignificantly to the fluorescence decay. The intercept ($\approx 5-8 \times 10^5$ s$^{-1}$) is consistent with N$_2$O$_5$ concentrations in the range 10$^{15}$ molecule cm$^{-2}$ and a value of $k_{q}(N_2O_3)$ of the order of 10$^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$.

Our result obtained in N$_2$, $k_{10} = (2.1 \pm 0.2) \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, is in reasonable agreement with the value of $(1.7 \pm 0.2) \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ reported by Nelson et al. (1983). In contrast, our result for quenching by water vapour, $k_{12} = (1.6 \pm 0.3) \times 10^{-10}$ cm$^3$ molecule$^{-1}$ s$^{-1}$, more than a factor of four lower than that reported by Fenter and Rossi (1997). As both studies used 662 nm excitation of NO$_3$ and similar methods to derive $k_{12}$, the differences are likely to be related to the measurement of the H$_2$O concentration. As we measured the H$_2$O concentration in situ (optical absorption at 185 nm) the uncertainty of our result is expected to be determined by uncertainty in the absorption cross section of H$_2$O at this wavelength, which, based on good agreement across several measurements (Cantrell et al., 1997; Hofzumahaus et al., 1997; Creasey et al., 2000) we estimate to be < 10%. Fenter and Rossi (1997) relied on flow measurements to derive the
concentration of H$_2$O in their experiments. Because of this, we consider our measurement of $k_{10}$ the more accurate and use this value for further evaluation of our experiments to derive $k_{12b}/k_{12}$.

### 3.2.3 Yield of OH from NO$_3^*$ + H$_2$O

Figure 7 displays the results of an experiment using three pulsed lasers. The first (excimer laser at time zero) generated OH from the 248 nm photolysis of HNO$_3$. In this particular experiment the HNO$_3$ concentration (monitored at 185 nm) was $6.3 \times 10^{15}$ molecule cm$^{-3}$ and a laser-fluence of 13 mJ cm$^{-2}$ was used to generate $2.0 \times 10^{12}$ OH cm$^{-3}$. This OH monitored by the 282 nm LIF-laser out to a reaction time of 10ms (open circles in Fig. 7), was observed to decay at a rate consistent with its well-characterised reaction with HNO$_3$ $k_{15}(298$ K, 22.5 Torr) = $1.3 \times 10^{-13}$ cm$^3$ molecule$^{-1}$ s$^{-1}$ (Dulitz et al., 2018). NO$_3$ is the unique product of reaction (R15) (Brown et al., 2001; Carl et al., 2001). The NO$_3$ profile (dashed line), calculated from initial OH and HNO$_3$ concentrations, is also displayed in Fig. 7. Here we calculate that 97% of the initial OH will react with HNO$_3$, the balance resulting from diffusion and reaction with NO$_2$. The decay of NO$_3$ at long reaction times is due to NO$_3$ diffusion from the reaction volume so that its concentration at 8.28 ms (when the 662 nm laser is triggered) was reduced by $\approx$ a factor of two compared to the stoichiometric yield of $2 \times 10^{12}$ molecule cm$^{-3}$ (i.e. when all OH is converted to NO$_3$). The decay of NO$_3$ was calculated from the known diffusion loss constant for OH at this pressure and the relative reduced-masses of OH and NO$_3$. A delay of 8.28 ms allowed the primary-OH to decay to very low values (i.e. $\approx 10^9$ molecule cm$^{-3}$) before triggering the 662 nm excitation laser. The measured laser fluence at 662 nm was then combined with the NO$_3$ concentration at 8.28 ms to calculate the fractional excitation of NO$_3$ (generally about 10%) and thus the concentration of NO$_3^*$ formed. When using very large laser fluences at 662 nm we calculate that the transition was saturated and then assume equal concentrations of ground and excited state NO$_3$ directly after the excitation pulse.

The solid lines starting at $t = 8.28$ ms represent the expected OH-signal if the value of $k_{12b}/k_{12}$ were 0.0, 0.01, 0.05 and 0.1 and were calculated using the rate constants for quenching of NO$_3^*$ by N$_2$ and H$_2$O as derived in this study as well as the concentrations of N$_2$ and H$_2$O.

Clearly, the data from the experiment illustrated in Fig. 7 are consistent with a value of $k_{12b}/k_{12}$ that lies between 0 and 1%. Similar experiments were repeated for different starting conditions and photoexcitation wavelengths (623, 629 and 662 nm) corresponding to strong absorption features of NO$_3$. No evidence for OH production in (R12) was observed in any experiment and an upper limit to the yield of OH was obtained from the random noise on the experimental OH-trace data and the expected OH signal. These values are tabulated in Table 2. The major sources of uncertainty in the calculated OH-yield are uncertainty in the measurement of laser fluences (30%) required to calculate the initial OH and NO$_3^*$ concentrations and assumptions related to the (unmeasured) NO$_3$ time profile. NO$_3$ is relatively unreactive in this system as it does not react with HNO$_3$ and only slowly with NO$_2$ (formed in R13) at these pressures. We calculate that $\approx$ 5% of the NO$_3$ formed is lost via reaction with OH ($k$(OH+NO$_3$) = $2 \times 10^{-11}$ cm$^3$ molecule$^{-1}$ s$^{-1}$) (Atkinson et al., 2004), its major removal processes being diffusive transport. The diffusive loss rate constant for NO$_3$ in this system was calculated from the known diffusive loss rate
constant of OH under the same conditions of pressure and temperature. In the absence of corroborative measurements of the NO3 profile in these experiments, we conservatively assume a factor two uncertainty in the NO3 concentration at the time of the excitation pulse. We thus derive an upper-limit of \( k_{12b} / k_{12} < 0.03 \) following photoexcitation at 623, 629 and 662 nm. This indicates either that the rapid quenching of NO3* by H2O predominantly involves energy transfer rather than reaction, or that the products formed in reactive quenching to not include OH.

4 Atmospheric implications and conclusions

The results obtained in this work and elsewhere (Crowley and Carl, 1997; Carr et al., 2009; Amedro et al., 2011) clearly demonstrate that the large values of \( k_{5b} / k_5 \) reported by Li et al. (2008) were erroneous. In this work we were able to reproduce, extend and improve upon previous results (i.e. obtain smaller upper-limits for \( k_{5b} / k_5 \)). The extension of the database to a wider range of photoexcitation wavelengths was important, since the majority of the data from Li et al. (2008) were obtained at wavelengths red-shifted from those of the other groups. In the modelling study by Wennberg and Dabdub (2008) the largest impacts of (R5b) on air quality (enhancements in O3 of \( \approx 40\% \)) were found when using \( k_{5b} / k_5 = 10^{-3} \) from Li et al. (2008). Small but still significant impacts changes in O3 and particle mixing ratios were calculated when using the upper-limit of \( k_{5b} / k_5 = 7 \times 10^{-5} \) provided by (Crowley and Carl, 1997). Results from this work, with upper-limits to \( k_{5b} / k_5 \) an order of magnitude smaller than those available previously, enable us to conclude that the formation of OH in NO2* + H2O is not an important atmospheric process.

Our upper limits of 3% to OH formation from the reactive quenching of NO3* by H2O can be put in context using equations (1) and (2). We combine our measurements of \( k_{10} = 2.1 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \) and \( k_{12} = 1.6 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \) with the literature value for \( k_{11} (2.1 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}, \text{ Nelson et al. (1983)} \) to derive \( f_{\text{H2O}} = 0.16 \) at 25°C and a relative humidity of 80%. Using the same excitation rates and concentrations of NO3 described in section 1.1 and our upper limit of \( k_{12b} / k_{12} = 0.03 \), we derive an OH production rate of \( \approx 7 \times 10^4 \text{ OH cm}^{-3} \text{ s}^{-1} \). Whilst this value is \( \approx \) two orders of magnitude lower than that calculated in section 1.1 where we assumed that all NO3* + H2O interactions form OH and used the high value of \( k_{12} \) from the literature (Fenter and Rossi, 1997), it is non-negligible compared to OH production rates from photolysis of O3 (see section 1.2) and may still represent an important contribution to OH formation in environments where OH generation via traditional routes involving absorption of UV radiation is suppressed e.g. at high-latitudes in winter.

The low yield of OH most likely results from the dominance of collisional energy transfer over reactive quenching of NO3 by H2O \( (k_{12b} << k_{12}) \). However, we also consider the possibility that the non-observation of OH in our experiments reflects the fact that the preferred products are HONO + HO2 (i.e. \( k_{12c} > k_{12b} \)) even though the molecular rearrangements required to form these products are less straightforward than for formation of OH and HNO3 if excited state NO3 has the same (approximate) D3h symmetry as the ground state and formally contains no O-O bonds. The conversion of HO2 to (detectable) OH via addition of
NO was not feasible owing to the rapid reaction of NO with NO\textsubscript{3} ($k(\text{HO}_2 + \text{NO}) \approx 8 \times 10^{-12}$ cm\textsuperscript{3} molecule\textsuperscript{-1} s\textsuperscript{-1}, $k(\text{HO}_2 + \text{NO}) \approx 3 \times 10^{-11}$ cm\textsuperscript{3} molecule\textsuperscript{-1} s\textsuperscript{-1} (Atkinson et al., 2004; IUPAC, 2018).

Given that our experiments were blind to formation of HO\textsubscript{2} or HONO a detailed discussion of the atmospheric role of reaction (R12c) is not warranted. However, the potential importance of reaction R12c can be illustrated by assuming a 10% yield of HONO and HO\textsubscript{2} ($k_{12c} / k_{12} = 0.1$) and the same temperature, NO\textsubscript{3} concentration and relative humidity outlined above. With this scenario, we calculate production rates of HO\textsubscript{2} and HONO of $\approx 2 \times 10^5$ molecule cm\textsuperscript{-3} s\textsuperscript{-1}. For HONO, this production rate is comparable to its formation in the gas-phase reaction between OH and NO under low NO\textsubscript{X} conditions but lower than the missing production rate of $\approx 1-5 \times 10^6$ molecule cm\textsuperscript{-3} s\textsuperscript{-1} that has been observed in several environments as summarised by Meusel et al. (2016). In terms of HO\textsubscript{2} formation, a rate of $2 \times 10^5$ molecule cm\textsuperscript{-3} s\textsuperscript{-1} would be comparable to that obtained by the photolysis of $\approx 0.5$ ppbv of HCHO (assuming a J-value for HCHO of $\approx 2 \times 10^{-5}$ s\textsuperscript{-1}). In conclusion, whilst our experiments indicate that the reactive quenching of excited NO\textsubscript{3} by water vapour is inefficient compared to collisional deactivation, we cannot rule out that this reaction plays a role in HO\textsubscript{X} or HONO production. Experiments sensitive to HO\textsubscript{2} or HONO formation and theoretical calculations could help shed light on this.

Acknowledgements

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References


Table 1  Experimental conditions and results for NO$_2^*$ + H$_2$O (R5)

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<th>$E_\lambda$</th>
<th>n</th>
<th>$P$</th>
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Notes: $\lambda$ = excitation wavelength (nm). $E_\lambda$ = excitation laser fluence (in $10^{16}$ photons pulse$^{-1}$ cm$^{-2}$); $n$ = number of repeat experiments; $P$ = bath-gas (N$_2$) pressure (mbar); units of concentration were $10^{15}$ molecule cm$^{-3}$.

Table 2 – Experimental conditions and results for NO$_3^*$ + H$_2$O (R12)

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<td>47</td>
<td>6.3</td>
<td>&lt; 0.003</td>
</tr>
<tr>
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<td>16</td>
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<td>16</td>
<td>49</td>
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Notes: $\lambda$ = excitation wavelength (nm). $E_\lambda$ = excitation laser fluence (in $10^{16}$ photons pulse$^{-1}$ cm$^{-2}$); $P$ = bath-gas (N$_2$) pressure (mbar); units of concentration were $10^{15}$ molecule cm$^{-3}$.
Figure 1: Rate constants for dissociative (black line, J-diss) and non-dissociative (blue line, J-exci) excitation of NO$_3$. The data use solar radiation actinic flux at the surface at 50°N and a solar zenith angle (SZA) of 27° (red line) as well as the NO$_3$ absorption cross sections and quantum yields. J-values (and fraction of NO$_3$ dissociated) were obtained by integration of the excitation rate (quanta s$^{-1}$ nm$^{-1}$) over the wavelength range of absorption.
Figure 2: Net effects of reactive removal of NO$_2^*$ and NO$_3^*$ by H$_2$O.
Figure 3: Photo-excitation of NO$_2$ at 532 nm. The open circles are OH-calibrations obtained by the 193 nm photolysis of $1.5 \times 10^{17}$ molecule cm$^{-3}$ H$_2$O (in the absence of NO$_2$) at different laser fluences (mJ cm$^{-2}$). The solid stars are data points from an OH calibration in the presence of NO$_2$. The black triangles are data obtained by photoexcitation of [NO$_2$] = $4.0 \times 10^{15}$ molecule cm$^{-3}$ using 532 nm (50 mJ cm$^{-2}$) in the presence of [H$_2$O] = $1.5 \times 10^{17}$ molecule cm$^{-3}$. The red triangles are the results of an identical experiment, but with the OH-excitation laser tuned away from the OH-feature at 282 nm. The solid black line represents the OH signal and concentration expected from the yield of OH from NO$_2^*$ + H$_2$O reported by Li et al. (2008).
Figure 4: Summary of data obtained following photoexcitation of NO$_2$ at various wavelengths. The data from this study, Crowley and Carl (1997), Carr et al. (2009) and Amedro et al. (2011) are all upper limits, indicated by the down-arrows. The NO$_2$ absorption cross-sections were taken from Van Daele et al (1998).
**Figure 5**: Exponential decay of fluorescence from NO$_3$ following photoexcitation at 623 nm in the presence of N$_2$. 

A) NO$_3$ fluorescence signal (a.u.) vs. time (s) with varying [N$_2$] (molecule cm$^{-3}$)

- 1: $3.76 \times 10^{16}$
- 2: $5.86 \times 10^{16}$
- 3: $9.85 \times 10^{16}$
- 4: $1.37 \times 10^{17}$
- 5: $1.80 \times 10^{17}$

B) NO$_3$ fluorescence signal (a.u.) vs. time (s) with varying [H$_2$O] (molecule cm$^{-3}$)

- 1: 0
- 2: $4.92 \times 10^{16}$
- 3: $1.32 \times 10^{16}$
- 4: $2.12 \times 10^{16}$
- 5: $4.31 \times 10^{16}$
Figure 6: Plots for the determination of total quenching rate coefficients for NO₃⁺ with N₂ (R10) and
with H₂O (R12) at 296 K: $k_{10} = (2.1 \pm 0.1) \times 10^{-11}$ cm³ molecule⁻¹ s⁻¹; $k_{12} = (1.6 \pm 0.3) \times 10^{-10}$ cm³
molecule⁻¹ s⁻¹. The error bars are statistical uncertainty (2σ) from the fits to OH-decays as
exemplified in Figure 5.
Figure 7: Plot of primary OH and expected OH (solid lines after 8.28 ms) from NO$_3^*$ + H$_2$O at various values (0 to 10%) of $k_{12b} / k_{12}$. The initial OH concentration (right y-axis) was $2.0 \times 10^{12}$ molecule cm$^{-3}$. The dashed red line displays the calculated NO$_3$ concentration, which at 8.28 ms (time of 662 nm excitation pulse) was $9 \times 10^{11}$ molecule cm$^{-3}$. In these conditions 50% of available NO$_3$ was excited to NO$_3^*$ by absorption at 662 nm; 35% of this NO$_3^*$ proceeded to react with H$_2$O in (R12), with the balance quenched by N$_2$ or HNO$_3$. The solid lines ($t > 8.28$ ms) represent expected OH signals for values of $k_{12b} / k_{12}$ between 0 and 10%.