

Spatially separate production of hydrogen oxides and nitric oxide in lightning

Jena M. Jenkins¹, William H. Brune¹

¹Department of Meteorology and Atmospheric Science, Pennsylvania State University, University Park, PA, USA

Correspondence to: Jena M. Jenkins (jzj76@psu.edu)

Abstract. The atmosphere’s most important oxidizer, the hydroxyl radical (OH), is generated in abundance by lightning, but the contribution of this electrically generated OH (LOH) to global OH oxidation needs to be better quantified. Part of the uncertainty in this contribution is due to the abundant nitric oxide (NO) also generated in lightning, which would rapidly remove the LOH before it can oxidize other pollutants in the atmosphere. However, atmospheric observations and a previous laboratory study show extreme LOH coexists with extreme NO. The only way this LHO_x can possibly survive is if LOH production is spatially separated from the NO production in lightning flashes and laboratory sparks. This hypothesis of spatially separate OH and NO production is further tested here in a series of laboratory experiments, where the OH decays were measured from spark discharges in air which had increasing amounts of NO added to it. The LOH decayed faster as more NO was added to the air, indicating that the LOH was reacting with the added NO, and not the spark NO. Thus, LOH from lightning flashes is not immediately consumed by the electrically generated NO but is available to oxidize other pollutants in the atmosphere and contribute to global OH oxidation. Subsequent modelling of the laboratory data also supports the spatially separate production of LOH and NO_x and further suggests that substantial HONO may also be produced by sparks and lightning in the atmosphere.

1 Introduction

Lightning and other electrical discharges have been shown to directly generate extreme amounts of the atmosphere’s primary oxidant, the hydroxyl radical (OH), and the closely related hydroperoxyl radical (HO₂) in field studies (Brune et al. 2021; Brune et al., 2022), laboratory studies (Jenkins et al. 2021; Ono and Oda, 2002), and modelling studies (Bhettanabhotla et al., 1985; Ripoll et al. 2014). The first reported field measurements of electrically generated OH and HO_x (together called the hydrogen oxides or HO_x) were from the Deep Convective Clouds and Chemistry campaign in 2012, where as much as ~2 ppbv of electrically generated HO_x (LHO_x) was measured (Brune et al., 2021). Subsequent laboratory studies showed that both lightning and weaker cloud electrical discharges, called corona discharges, were generating the extreme amounts of LHO_x and that LHO_x was initially generated as equal amounts of LOH and LHO₂ (Jenkins et al., 2021). Based on these studies, lightning and corona discharges in thunderstorms are together estimated to account for as much as 2-16% of global OH. However,

Deleted: 16802,

Deleted: remains highly uncertain

Deleted: is

Deleted: evidence from

Deleted: indicated

Formatted: Subscript

Deleted: is not immediately consumed by NO, possibly because LOH’s production

Deleted: ,

Deleted: is

Formatted: Subscript

Formatted: Subscript

Formatted: Subscript

Formatted: Subscript

Formatted: Subscript

Formatted: Subscript

40 narrowing down the uncertainty of this range will require more work. The frequency, duration, and location of corona discharges is not well known, complicating attempts to estimate global OH production from these discharges.

In comparison, it is accepted that lightning flashes occur at a rate of 44 s^{-1} globally (Christian et al., 2003), last <1 second (Rakov and Uman, 2006), and are mostly detected by satellites and lightning networks. The extreme amount of nitrogen oxide (NO) also generated in lightning makes estimating the impact of LHO_x difficult, as theoretically this NO will rapidly remove the extreme OH before it oxidizes other chemical species in the atmosphere, such as methane, carbon monoxide, sulfur dioxide, or other pollutants. However, evidence from a previous laboratory study shows that LHO_x is not immediately destroyed by electrically generated NO (LNO). In Jenkins et al. (2021), laboratory sparks were generated inside a flow tube, and the subsequent LNO and LHO_x formed from these discharges was measured. Hundreds of pptv of LHO_x was observed to decay over hundreds of milliseconds, while simultaneously 1-2 ppmv of LNO was also measured. When these same measurements of LHO_x and LNO from the laboratory experiments were input into a photochemical box model, the Framework for 0-D Atmospheric Modelling (F0AM) (Wolfe et al., 2016) with the Master Chemical Mechanism v3.3.1 (Jenkin et al., 2015), the model predicted that LNO should have titrated all LHO_x away in less than 10 ms, a small percent of the hundreds of milliseconds over which the LHO_x decay was actually observed. It is unlikely that this discrepancy is due to some unimagined chemistry considering how well studied this chemistry is. Therefore, the only logical conclusion is that LHO_x and LNO generation are spatially separated for the spark, preventing their immediate reaction.

55 Spatially separate production is possible due to the structure of and different types of energy present in lightning flashes and sparks. At the center of a lightning flash is a $\sim 1\text{-}2$ cm diameter core (Rakov and Uman, 2006) with air temperatures exceeding 30,000K (Orville, 1968a). Surrounding this hot core is a weaker area of electrical discharge, called the corona sheath. The air temperature in the corona sheath is near ambient, and the electrical discharges from the sheath extend radially several meters from the hot core (Rakov and Uman, 2006), so that the ratio of the volume of the corona sheath to the volume of the core is at least 10^4 . Some of the radiation emitted by lightning flashes is in the ultraviolet (UV) range, composed of both broad spectrum and line emissions (Orville, 1968b), and including wavelengths <300 nm that are emitted from the sun but normally not present in the troposphere due to their absorption in the higher levels of the atmosphere by ozone. This UV radiation is generated by both the hot core and the corona sheath. The reach of the UV radiation depends on the wavelength and scattering the radiation encounters but can be as much as tens of meters. Sparks are essentially a smaller scale version of lightning flashes, still composed of a hot core (though not as hot as lightning) surrounded by a weaker and cooler corona sheath and emitting UV radiation (though not as much as lightning).

70 The differences between the core and corona sheath lead to different chemistry occurring in each area. For example, the extremely high temperatures of the lightning flash or spark core are required to dissociate stable N_2 and make the extreme amounts of NO present in lightning flashes via the Zel'dovich Mechanism (Chameides et al., 1977):

Deleted: the exact contribution of electrical discharges to global atmospheric OH is highly uncertain.

Deleted: weaker electrical

Deleted: , like streamers or corona, are

Deleted: types

Deleted: we know

Deleted: can

Deleted: when and where they occur with

Deleted: mapping arrays

Deleted: ,

Deleted: but t

Deleted: lightning generated OH and HO_2 (together called the hydrogen oxides or

Deleted:)

Deleted: could

Deleted: suggest

Deleted: electrically generated HO_x (

Deleted:)

Formatted: Subscript

Formatted: Subscript

Deleted: LHO_x generated from a laboratory spark was measured decaying over the course of hundreds of milliseconds, while a modelled decay generated with a

Formatted: Subscript

Deleted: found that

Deleted: should decay

Deleted: with the extreme LNO simultaneously generated.

Formatted: Subscript

Deleted: This discrepancy between the laboratory and model decays suggests that either

Deleted: in

Deleted: or some chemistry is missing or incorrect in the model

Deleted: and cooler

Deleted: ,

Deleted: ing

Deleted:)

Formatted: Superscript

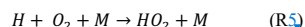
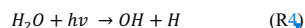
Deleted: The

Deleted: (Chameides et al., 1977),



The air cools down rapidly after the lightning flash, removing the energy required for the reverse reactions to convert NO back to N_2 and O_2 faster than these reactions can occur. As a result, elevated NO remains after the lightning flash is completed.

Conversely, without the high temperatures, the corona sheath makes several orders of magnitude less LNO (Rehbein and Cooray, 2001; Bhetanabhotla et al., 1985), so less than 1% of the spark NO is made outside the core. However, large amounts of OH, though not HO_2 , are also made by combustion at the high temperatures of the core (Dyer and Crosley, 1982; Bhetanabhotla et al. 1985; Ripoll et al., 2014), while both OH and HO_2 are made through multiple pathways in the corona sheath. These pathways include, for example, OH-forming reactions like $electron + H_2O \rightarrow OH + H$, or $O^1D + H_2O \rightarrow 2OH$, reactions that form HO_2 like $H + O_2 + M \rightarrow HO_2 + M$, or UV radiation directly dissociating water vapor at wavelengths <200 nm, directly producing equal amounts of OH and HO_2 :



In short, LNO production is contained in the very narrow hot core, while HO_x production occurs in both the hot core and in a volume extending several meters the outside the hot core in the corona sheath. Thus, spatially separate LNO and LNO production is possible.

To further test the hypothesis that LNO and LNO production are spatially separated in spark discharges, we conducted a series of laboratory experiments in which the LOH and LHO₂ decays from spark discharges in air were measured with different amounts of background NO added into the air flow, from 0 ppbv up to 1000 ppbv of added NO. The decays from the laboratory experiments are also compared to decays calculated by F0AM with MCM to see if the model can successfully reproduce these decays. If LHO_x decays faster as the background NO mixing ratio is increased, then LHO_x is mostly or entirely reacting with background NO instead of spark LNO, confirming that LHO_x and LNO generation is spatially separated in the spark. Otherwise, if the LHO_x decays are unaffected by the amount of added NO, then LHO_x is mostly or entirely reacting with spark LNO, LHO_x and LNO are likely generated in the same location, and some unimagined chemistry is causing the discrepancy between model and measurement.

Formatted: Centered

Formatted: Subscript

Formatted: Subscript

Deleted: but I

Deleted: HO_x

Deleted: can

Deleted: be

Formatted: Subscript

Formatted: Subscript

Deleted: . Corona can make extreme HO_x through multiple pathways (Bruggeman and Schram, 2010) while making orders of magnitude less LNO than lightning or sparks (Rehbein and Cooray, 2001). ...

Deleted: an electron dissociating a water molecule (

Deleted:

Deleted:)

Deleted: an excited state oxygen atom

Deleted: dissociating a water molecule (

Deleted:)

Deleted: among several others

Deleted: .

Formatted: Subscript

Deleted: can also make extreme OH while making little to (... [1]

Deleted: producing

Deleted: HO_x in two steps

Formatted: Subscript

Deleted: 1

Deleted: 2

Deleted: Thus, with the

Deleted: mostly

Deleted: to the

Deleted: but

Deleted: the HO_x

Deleted: ring

Deleted: both in

Formatted: Subscript

Deleted: outside

Deleted: and the UV radiation

Deleted: ,

Deleted: were conducted where

Deleted: unaccounted-for

170 2 Methods

2.1 Laboratory Experimental Setup

The laboratory setup was nearly identical to the setup used in our previous LHO_x studies (Jenkins et al. 2021; Jenkins and Brune, 2023). Purified and dried air, with an OH reactivity of $\sim 0.35 \text{ s}^{-1}$ (Brune & Jenkins, 2024), was flowed through a bubbler to add a controlled amount of water vapor, then mixed with dry air that flowed down a quartz (previously Pyrex®) tube (50 mm OD x 46 mm ID x 105 cm) at 50 standard liters per minute (slpm), through spark discharges, and over to instruments for measuring OH and HO₂ (Ground-based Tropospheric Hydrogen Oxides Sensor [GTHOS; Faloona et al., 2004]), NO-NO₂-NO_x (ECO PHYSICS nCLD 855Y), and O₃ (Kalnajs & Avallone, 2010). A solid-state Tesla coil (Eastern Voltage Research, Plasmasonic® 1.3) was used to generate the sparks across a 0.7 cm gap between tungsten wire electrodes (0.10 cm diameter) inside the flow tube. The sparks were generated in packets of 10 sparks, with $\sim 75 \text{ ms}$ between each spark in the packet, as signals from individual sparks were too narrow to consistently measure even at the 5 Hz sampling rate of GTHOS. The NO_x analyzer collected data at a rate of 2 Hz and the O₃ analyzer collected data at a rate of 1 Hz. Each electrode was attached to a copper rod; one copper rod was attached via a copper wire cable to the output toroid of the Tesla coil, while the other was attached to an electrical ground. All discharges were generated using the same Tesla coil settings. Pressure (MKS Baratron® Type 222) was monitored ahead of the inlet for GTHOS and the Teflon tubing leading to the NO_x and O₃ analyzers, temperature was measured both before air entered the flow tube (Vaisala HMT310) and as the air exited (thermistor), and the water vapor mixing ratio (Vaisala HMT310) was also measured before the air entered the flow tube. The air velocity was measured with an anemometer (TSI Inc., 8455-09) before running experiments, and the flow in the tube was previously determined to be laminar that is not fully developed (Jenkins et al., 2021). A short piece of Teflon tubing (1.3 cm diameter x 2.5 cm long) was placed on the GTHOS inlet, and the opening of the Teflon tube leading to the NO_x and O₃ analyzers was positioned $\sim 2 \text{ mm}$ downstream of the GTHOS opening and facing into the short piece of Teflon tubing. This arrangement ensured that GTHOS and the NO_x and O₃ analyzers all sampled from the same volume. The absolute uncertainty and limit of detection at the 68% confidence level was $\pm 20\%$ and $\sim 1 \text{ pptv}$ for the HO_x measurements from GTHOS, $\pm 10\%$ and $\sim 1\text{--}3 \text{ ppbv}$ for the NO_x measurements, and $\pm 5\%$ and $\sim 20 \text{ ppbv}$ for the O₃ measurements. A diagram of the laboratory setup is shown in Figure 1.

Deleted: S

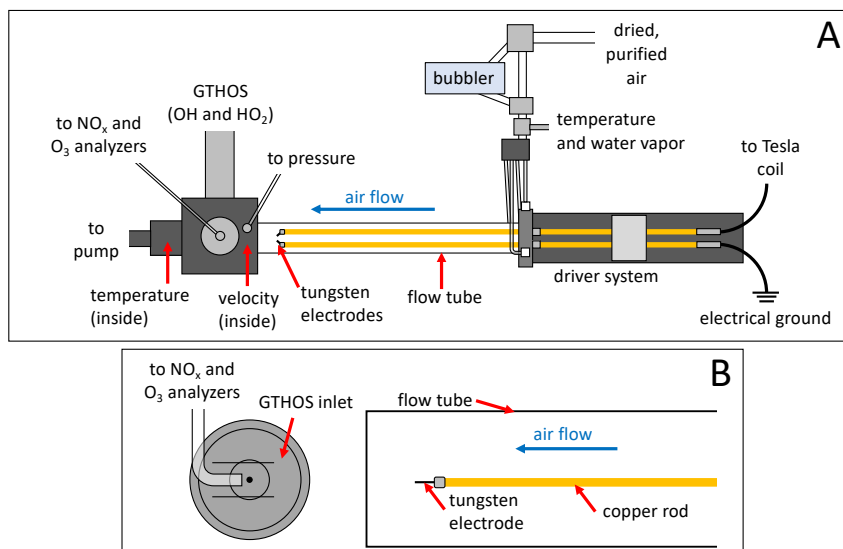


Figure 1: (A) Top-down diagram of the laboratory experimental setup showing the key components. (B) Side-view showing a close-up of the GTHOS inlet and Teflon line leading to the NO_x and O₃ analyzers, which sample from the same volume as GTHOS in a 1.3-cm diameter tube placed over the GTHOS inlet (shown as two horizontal lines), along with the relative positions of the flow tube, copper rod, and tungsten electrode. Neither (A) nor (B) is shown to scale.

As in real lightning, the core of the sparks is small relative to volume available for the corona sheath and ultraviolet radiation to occupy. Based on the visible light, the spark core is estimated to be ~1 mm in diameter across the 0.7 cm spark gap, so the core volume occupies ~0.006 cm³. Assuming a similar ratio of corona sheath to core in the sparks that is present in lightning, then the spark corona sheath can occupy a volume as large as 55 cm³ with a radius of 5 cm, although the 4.6 cm inner diameter of the flow tube will be the actual cut off point for the corona sheath. We can detect the UV radiation from the spark discharges with a spectrometer placed outside of the flow tube, so the UV radiation also travels well beyond the spark core. The air that the NO instrument samples through a 4 m long 1/4" Teflon line is well mixed, indicating that the actual core NO is much higher than measured. However, GTHOS pulls 6 slpm and thus each 0.2 second measurement contains a volume of surrounding air hundreds of times larger than the ~0.006 cm³ volume of core air. Because GTHOS is right at the exit of the flow tube and the flow is laminar and 50 slpm, molecular diffusion mixes the core air into a volume less than 1 cm³ at the longest reaction time of ~0.5 s. Thus, GTHOS samples both core and sheath air, but they are spatially separated in the flow tube when sampled.

Deleted: ¶

Formatted: Superscript

Formatted: Superscript

215 The experiments were conducted as follows. To capture the LHO_x decay, the copper rods were moved by a driver system so
discharges were generated in 5 different positions in the flow tube, over a total distance of 27.5 cm. In each position, four spark
packets were generated, with 5 second spacing between each packet. For one of the four spark packets, the laser on GTHOS
was switched to a wavelength slightly off the OH absorption wavelength to confirm the absence of electrical interference in
the OH and HO₂ signals. By moving the discharge, the distance between the discharge and instrument inlets was changed,
which also changed the time between the LHO_x generation and measurement, producing the LHO_x decay over time. The
220 different amounts of added NO in the system were created by adding NO (Linde, 4.83 ppm) to the air flow before it entered
the flow tube to create mixing ratios of 0, 50, 100, 250, 500, or 1000 ppbv (all within ± 6%). Because lightning can occur at
any pressure in the troposphere, data were collected at pressures of 970 hPa, 770 hPa, 570 hPa, and 360 hPa (all within ± 2%)
to cover most of the tropospheric pressures. Data were also collected at water vapor mixing ratios between 2000-2400 ppmv
and temperatures between 289-294K.

225 Normally GTHOS uses two detection axes to simultaneously measure OH and HO₂, but only one detection axis was available
when these experiments were conducted. To obtain both OH and HO₂ measurements for these experiments, OH was measured
in a set of experiments, and total HO_x was measured in another set of experiments conducted under the same conditions. The
average OH measured at each position was subtracted from the total HO_x generated at the same position and collected under
230 the same conditions to determine the HO₂ generated.

2.2 Laboratory Data Processing

Each 10-spark discharge packet created a single spike in the OH, HO₂, NO, and NO_x signals. Figure 2 shows the OH and NO
signals from the spark packets over time for one experiment. No O₃ was detected in these experiments. These spikes were
integrated over time to determine the total amount of chemical generated by the spark discharge. For the OH and HO₂
235 measurements, the peaks were about ~1.2 seconds wide and were integrated over 2.2 seconds, while the NO and NO_x peaks
were ~4.8 seconds wide and were also integrated over 4.8 seconds. From previous tests, only about 85% of the generated LNO_x
is sampled (Jenkins et al., 2021), so the LNO and LNO₂ results were corrected up 15% to account for the LNO_x that is not
sampled. OH and HO₂ have similar diffusion coefficients to NO_x (Tang et al., 2014), so OH and HO₂ were also corrected up
15% to account for sampling. Additionally, the lifetime of NO_x is long relative to the time it spends in the flow tube (hours vs
240 <0.5 seconds, respectively), so any change in the NO_x mixing ratio across the different positions was assumed to come from
diffusion and not chemical loss. The average change in NO_x over the different discharge positions in the flow tube is shown
in Figure S1₄ for all four pressures tested. The LOH and LHO₂ measurements were also corrected up based on the NO_x diffusion
to account for diffusion losses.

Deleted: s

Deleted: 3

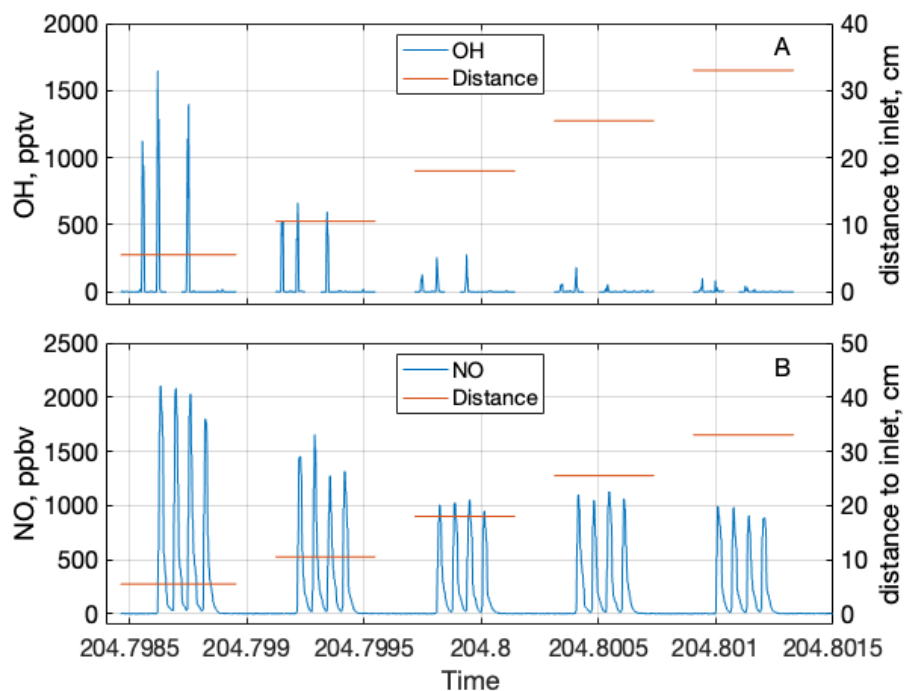


Figure 2: Change in OH (A) and NO (B) mixing ratios due to the spark discharges at each of the five discharge positions at 770 hPa and 0 ppbv of added NO. Each peak is from one spark packet containing 10 sparks. OH and NO mixing ratios are indicated by the blue lines and use the y-axes on the left side of their respective subplots, while the distances from the discharge to the GTHOS inlet and Teflon line leading to the NO_x analyzer are indicated by the orange lines and use the y-axes on the right side.

Both the LOH and LHO₂ decays were fitted with equations assuming constant, first-order losses. These equations were extrapolated back to time-zero to determine the initial amount of these species generated in the discharge. In some experiments, the HO_x decay was fast enough that the HO_x data became too small and imprecise to use at farther discharge positions in the flow tube. If at least 3 positions had clear OH and HO₂ signals, the decay was included in the results; if only 2 positions or less

were available, the data were not used in the results, as there was not enough confidence in the extrapolated fit. Consequently, not all pressures have results for all the different amounts of added NO.

The initial LNO_x formed in the discharges was taken as the LNO_x in the position closest to the instrument inlets as it was least affected by diffusion. NO₂ made up <10% of total NO_x.

2.3 Model Setup

The modelling experiments were conducted using F0AM v3 with MCM 3.3.1 chemistry. The laboratory data were collected in 10 spark packets, but the chemical measurements were scaled down to single spark equivalents before inputting them into the model. The reason for scaling down is two-fold. First, even at the slowest speed in the flow tube, one spark will travel ~7 cm before the next one occurs, and previous work has shown that the HO_x and NO_x measurements scale proportionally to the number of sparks in the packet (Jenkins et al., 2021), indicating that the chemicals generated by sparks within a packet are likely not overlapping. Second, due to the nonlinear chemistry between HO_x and NO_x, we cannot assume that any modelling done with 10 sparks will scale simply to a single spark. Therefore, because each spark within a packet can be treated as an independent event, the modelling was done using HO_x and NO_x values scaled down to a single spark.

The initial OH and HO₂ determined from the extrapolation of the laboratory decays, scaled down 10-fold, were chosen as the initial OH and HO₂ (respectively) for the model runs. Using this same initial HO_x, three cases using different amounts of initial NO_x were tested. In the first case, only the added NO was included in the model, and no spark NO_x was included. In the second case, the added NO plus all the spark NO_x was included, and in the third case, the added NO plus only a small percentage of the spark NO_x was included. The purified air used in the laboratory experiments was found to contain ~20 ppbv of CO (Thermo Scientific, 48i-TLE) which was also included in all the model experiments, along with wall loss at a rate of 0.9 s⁻¹ for OH (no wall loss was observed for HO₂). Model tests confirmed that even if up to 20 ppbv of O₃ (our limit of detection) had been generated in the laboratory experiments, it would not have significantly affected the HO_x decays, so O₃ was not included in any of the model runs shown here. The model experiments were set to simulate 0.5 seconds of reaction time, enough to cover the longest reaction timescale of the laboratory experiments, using the same pressure, temperature, and water vapor as the laboratory experiments, and included no dilution.

3 Results

3.1 Laboratory Results

As an increasing amount of NO was added to the air flow in the laboratory experiments, the OH and HO₂ decays became progressively steeper, as shown Figure 3 (970 hPa and 360 hPa), Figure S2 (770 hPa and 570 hPa), and Figure S3 (average

Deleted: 1

Deleted: 4

Deleted: 5

slopes for all experiments). In other words, both OH and HO₂ decayed faster as more NO was added to the air flow. This dependence of the OH and HO₂ decays on the added NO indicates that LHO_x is reacting mostly with the added NO, and little or not at all with the spark NO_x, supporting the hypothesis that the HO_x we measure from spark and lightning discharges is produced separately from the spark NO_x. The average LNO_x generated in the laboratory experiments is shown in Figure S4

Deleted: 6

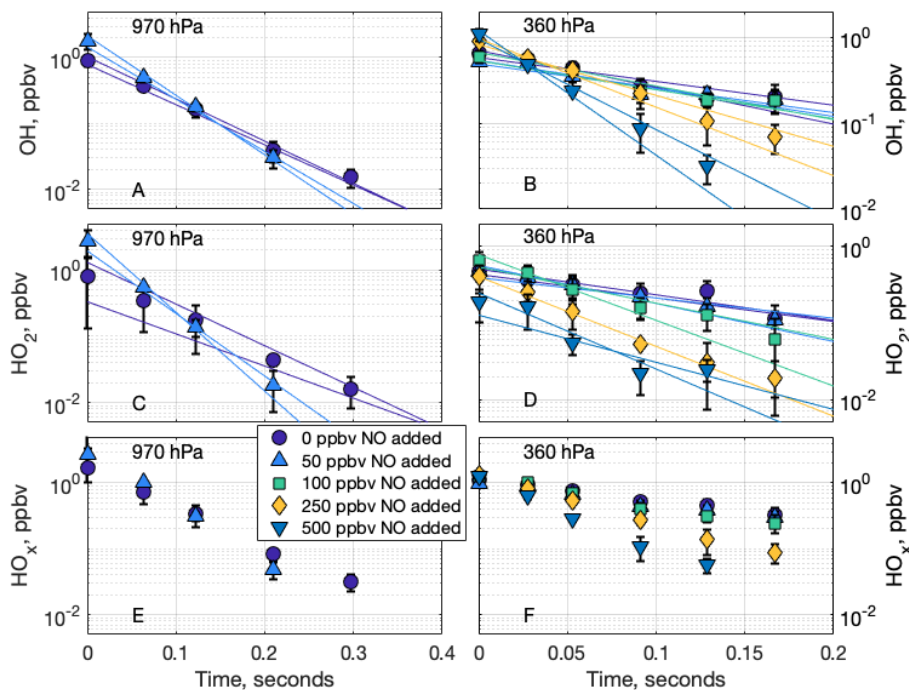


Figure 3: Laboratory decays of OH (A,B), HO₂ (C,D), and net HO_x (E,F) at 970 hPa (A,C,E) and 360 hPa (B,D,F). The markers are the averaged data points containing 3 or 6 measurements from 1 or 2 laboratory experiments, respectively. The markers at time zero are the averaged extrapolated values from the decays. The lines on A, B, C, D are the linear fits to the individual decays. Error bars are the standard deviation from averaging the multiple laboratory measurements.

Deleted: 1

3.2 Laboratory versus model decays

Comparing the laboratory OH decays to the model decays from F0AM further supports the separate production of LHO_x and LNO, but also indicates that LHO_x and LNO or other chemical products from the spark discharges are likely interacting. For example, at 770 hPa and 0 ppbv of added NO, the laboratory LHO_x measurements decay neither as fast as when 100% of the spark NO_x is added to the model nor as slowly as when no spark NO_x is added to the model (Figure 4A,B). If LHO_x and LNO_x were generated in the same place, the laboratory LHO_x decays would match the model decay with 100% LNO_x included, and if LHO_x and LNO_x did not interact at all, the laboratory decays would match the 0% LNO_x model case. The laboratory decays falling in between the two model runs indicates that LHO_x is either partially interacting with LNO_x, or it is interacting with some other product(s) from the sparks.

As the background NO was increased, the gap between the laboratory decay and 0% LNO_x model case decreases (Figure 4C,D), and this gap decreases further as more background NO was added (Figure 4E,F). This decrease in the difference between the laboratory and model decays is likely because as the background NO was increased, it accounted for an increasing amount of the HO_x reactivity compared to the spark products. This increasing agreement between the model and laboratory decays as the added NO increased can be seen at 970 hPa, 570 hPa, and 360 hPa as well (Figures S5, S6, S7, respectively), and is another indicator that LHO_x is mostly made separate from the LNO_x made in the spark hot channel.

Deleted: 2

Deleted: 2

Deleted: 2

Deleted: 7

Deleted: 8

Deleted: 9

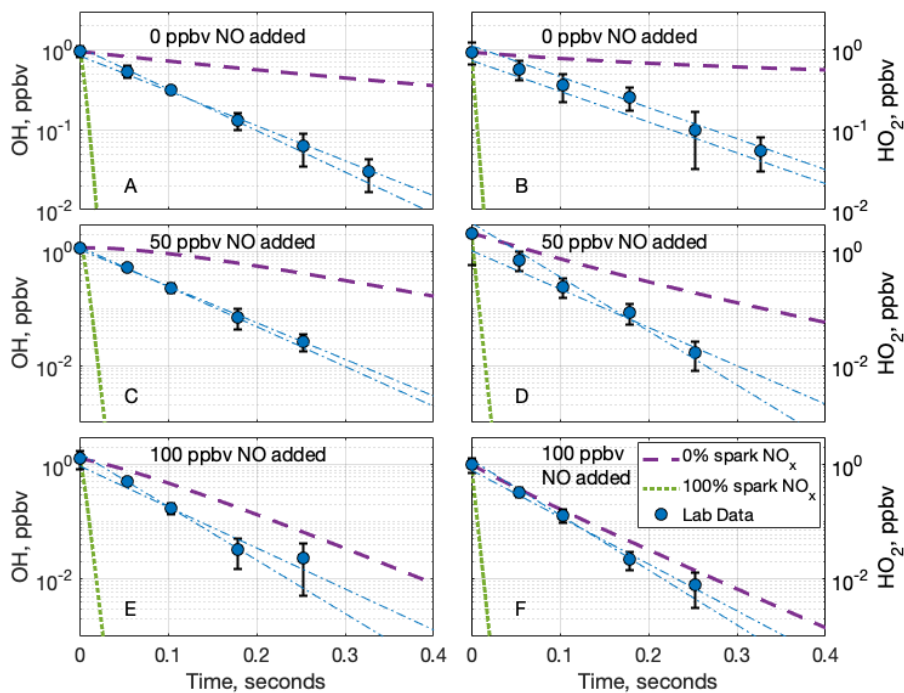


Figure 4: Comparison of measured OH (A,C,E) and HO₂ (B,D,F) laboratory decays and two model decays at 770 hPa and (A,B) 0 ppbv of added NO, (C,D) 50 ppbv of added NO, and (E,F) 100 ppbv of added NO. The dashed purple lines are the model decay with only the added NO, and includes no NO_x from the spark, and the dotted green lines are the model decay with the added NO and all of the spark NO_x. The blue circles are the average laboratory measurements and average extrapolated value at time zero, while the dashed-dotted blue lines are the individual extrapolated linear fits to the laboratory data. Error bars are the standard deviation from averaging multiple measurements.

3.3 Improving the measurement-model agreement

The agreement between the laboratory and model decays is at its worst when 0 ppbv of NO was added in the laboratory experiments. As these cases are also the most relevant to the atmosphere, trying to resolve this disagreement can also give insight into lightning chemistry in the atmosphere.

Deleted: 2

When we first observed this measured-modelled discrepancy in Jenkins et al., 2021, we were able to resolve the discrepancy for both OH and HO₂ by including just 0.5% of the spark NO_x in a model run. However, the model in the previous study was initialized using the full 10 spark packet data and also did not include the OH wall loss. Here, adding 3% of the spark NO_x to the model (amounting to 61.6, 62.8, 69.9, and 90.7 ppbv of NO_x at 970, 770, 570, and 360 hPa, respectively) brings agreement within uncertainty to the laboratory HO₂ data, but the OH data is still overestimated by the model (Figure S8). Adding 5% (104, 105, and 117 ppbv at 970, 770, and 570 hPa, respectively) or 10% (303 ppbv at 360 hPa) of LNO_x instead brings measured-modelled agreement for OH, but the HO₂ data is then consistently underpredicted by the model (Figure S9). There is no amount of LNO_x that can match the OH and HO₂ measurements simultaneously, leaving some chemistry still unaccounted for in the model.

Adding ~10 s⁻¹ of OH reactivity into the model along with 3% LNO_x can resolve the discrepancy (Figure S10) within uncertainty. What chemical species could be responsible for this reactivity? In addition to the HO_x, NO_x and O₃ we measure, many other species are generated in sparks as well, including atoms, ions, and excited states such as O, N, H, N₂⁺, O(¹D), O⁻, and others; other molecules that are primary products of the discharge, like N₂O and CO; and secondary products formed from reaction between or within the first two categories, like H₂O₂, HONO, and NO₂ (Bhetanabhotla et al., 1985; Boldi, 1992; Ripoll et al., 2014). For one (or more) of these species to account for the missing reactivity, it must fulfill a few criteria. First, its lifetime needs to be long enough so it is still present over the time frame we measure the HO_x decays, at least 0.2-0.5 seconds post-discharge. Second, it needs to react with OH on the same 0.2-0.5 second time frame, so it must either react with OH quickly or be present in large enough quantities to compensate for a slow reaction rate. Third, it must spatially overlap with the LHO_x we measure, so either it is produced in the corona sheath and/or UV radiation, or it is produced in large amounts in the hot core, with ~3% mixing out as we think LNO_x is doing. Lastly, the reaction between OH and this species must not produce HO₂. The mismatch between the model and measurements is because OH is overpredicted by the model relative to HO₂. If the reaction between OH and the missing species yields HO₂, then instead of increasing the OH loss rate, OH will be quickly recycled through the reaction $HO_2 + NO \rightarrow OH + NO_2$.

Neither of the first two categories of species, the atoms, ions, and excited states or the other primary molecules, can account for the missing reactivity in the model. The lifetime of the atoms, ions, and excited states species will be too short to affect the HO_x decays over 0.2-0.5 seconds, failing the first criterion. On the other hand, the primary products CO and N₂O fail the second criterion. Both species are longer lived than the first category, but their reactions with OH are relatively slow, and not enough of these species will be produced to compensate. For example, only about ~340 ppbv of N₂O is expected to be made in the combined hot core and corona sheath of a lightning flash (Brandvold et al., 1989; Brandvold et al., 1996; Donohoe et al., 1977; Hill et al., 1984; Levine et al., 1979), but ~11,000 ppmv would need to be produced in the laboratory sparks to compensate for a reaction rate of $k_{N_2O+OH} = 3.8 \times 10^{-17} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1}$ (Biermann et al., 1976). The reaction between CO

Deleted: 10

Deleted: 11

Deleted: 2

and OH is faster, with $k_{CO+OH} = 2.3 \times 10^{-13} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1}$ at 970 hPa in F0AM, and only ~1.8 ppmv of CO is needed to satisfy the missing reactivity in the model. But this 1.8 ppmv is ~12% of the 14.6 ppmv of CO expected to be made in the lightning hot core (Bhettanabhotla et al., 1985; Levine et al., 1979), and it is unlikely that the laboratory sparks are making as much CO as a lightning flash. The reaction of CO and OH also produces HO₂, leading to OH recycling.

The secondary discharge products are long-lived enough to still exist 0.2-0.5 seconds after the discharge, and their reaction rates with OH are faster than the rates with the primary products, so less of them are required to satisfy the missing reactivity compared to the primary products. Still, modelling results indicate that at most ~400 ppbv of H₂O₂ is generated in the lightning hot channel, and if only 3% of the hot channel mixes out, then this will not be enough to satisfy the ~250 ppbv of H₂O₂ needed to account for the missing OH reactivity in the sparks based on the reaction rate of $k_{H_2O_2+OH} = 1.7 \times 10^{-12} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1}$ from F0AM. Additionally, the reaction of OH and H₂O₂ produces HO₂. For NO₂, we have already included 3% of what we measure in the laboratory experiments in the model runs, which amounts to <10 ppbv of NO₂.

HONO, however, could account for the missing reactivity. It meets all four of the criteria: it lasts long enough to affect the HO_x decays; its reaction with OH does not recycle HO_x; it can react with OH over the 0.2-0.5 second time frame; and production of HONO in the core is expected to be high enough that only ~3% overlapping from the core could account for the OH reactivity. A model study including HONO production in the hot lightning core suggests as much as 12.6 ppmv of HONO can be generated within 10 ms of the discharge (Bhettanabhotla et al., 1985), and we only need ~70 ppbv of HONO to fulfill the missing reactivity, using the F0AM reaction rate of $k_{OH+HONO} = 6.1 \times 10^{-12} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1}$. Even considering that the laboratory sparks are smaller and cooler than a real lightning flash, substantial HONO production in the range of 1-2 ppmv is possible for the laboratory sparks as well.

Deleted: very much

Chemical models of the hot lightning channel show that both LNO and LOH production is extreme inside the lightning hot channel. For example, the model from Bhettanabhotla et al. (1985) has as much as 4300 ppmv of LNO and 860 ppmv of LOH initially produced, while the simulations of Ripoll et al. (2014) has as much as 42000 ppmv of LNO and 8400 ppmv LOH, with LNO and LOH within an order of magnitude of each other in the shock front. Little to no HO₂ is expected to be generated in the hot channel (Bhettanabhotla et al., 1985; Ripoll et al., 2014). As a test, a model experiment was run assuming 4 ppmv of LNO is initially produced in the laboratory sparks, which is only ~1.4-2 times our laboratory measurements for LNO, along with 2.8 ppmv of hot core LOH and no other chemicals added. The result of this experiment is HONO production in the range of 1-2 ppbv across all pressures (Table 1). Additionally, this HONO is generated fast, before we make our first measurement of HO_x in the laboratory flow tube. All the core LOH is also titrated to <1 pptv (our limit of detection in these experiments) over the same time frame the HONO is generated, so it would not be detected by GTHOS in the laboratory experiments. This model result is consistent with our laboratory observations because if substantial core LOH remains beyond the time the first measurement is made in the laboratory, then we would expect to detect significantly more LOH than LHO₂ during the

experiments instead of the relatively equal amounts of LOH and LHO₂ that are actually detected. This result is also in line with the Bhetanabhotla et al. (1985) model prediction that all the core LOH should decay away very rapidly. The only model case where the core LOH is not titrated to less <1 pptv before the first laboratory measurement is made is at 360 hPa, but even at this pressure, the model predicts that HONO, NO, and NO₂ are all within 1% of their final values when that first measurement is made.

Table 1. Comparison of the averaged NO and NO₂ measured in the laboratory experiments and the predicted NO, NO₂, and HONO from a model run starting with 4 ppmv of LNO and 2.8 ppmv of LOH.

	970 hPa		770 hPa		570 hPa		360 hPa	
	Lab	Model	Lab	Model	Lab	Model	Lab	Model
NO (ppbv)	1850	1820	1950	1870	2200	1930	2900	2040
NO ₂ (ppbv)	220	380	140	410	140	440	110	490
HONO (ppbv)	-	1670	-	1590	-	1490	-	1300
Time* (s)	0.064	0.019	0.055	0.027	0.042	0.0384	0.028	0.074

*For the laboratory data, time is when the first HO_x measurement is made post-spark. For the model data, time is when OH has been titrated to <1 pptv, our limit of detection in these experiments.

This model run demonstrates that HONO can be formed fast and in large amounts in the spark discharges. The initial chemistry in the sparks is occurring at thousands of degrees Celsius with electrons and many other chemical species besides NO and OH present, and the production of these species may have spatial dependencies that we cannot incorporate or account for in F0AM. These limitations may explain why the model does not entirely reproduce the NO and NO₂ laboratory measurements. Still, the model results are within an order of magnitude of the laboratory results while simultaneously producing substantial HONO. Adding 3% of the modelled HONO from Table 1 into the model of the laboratory decays drastically improves the agreement between the modelled and measured OH, and in some cases brings the modelled and measured decays into agreement within the laboratory uncertainty (Figure 5). A diagram of the simplified HO_x and NO_x spark chemistry discussed in the preceding paragraphs is shown in Figure 6.

Deleted: 3

Deleted: 4

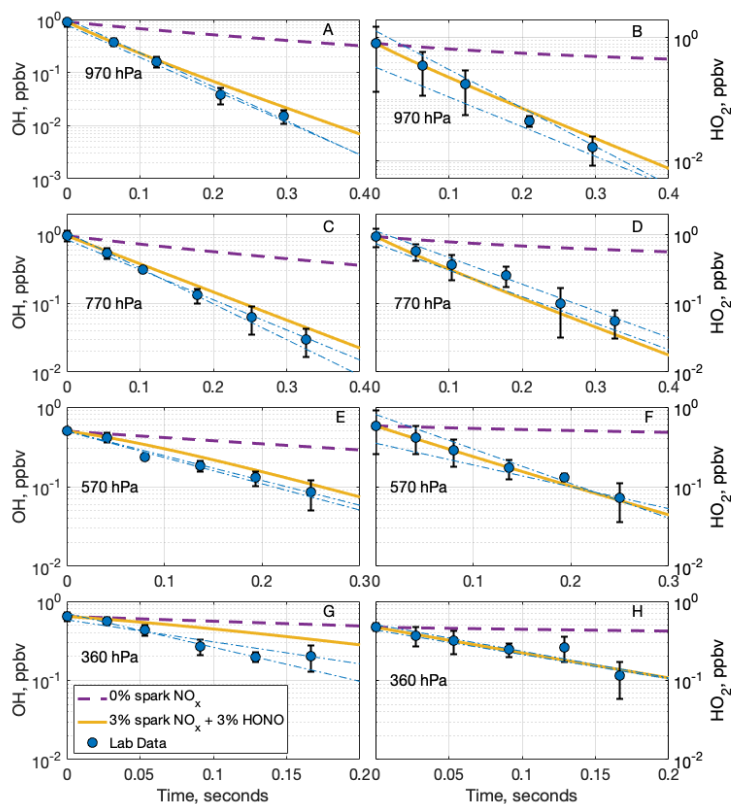


Figure 5: Comparison of measured OH (A,C,E,G) and HO₂ (B,D,F,H) laboratory decays and two model decays at (A,B) 970hPa, (C,D) 770 hPa, (E,F) 570 hPa, and (G,H) 360 hPa. The dashed purple lines are the model decay including no NO_x from the spark, and the solid yellow lines are the model decay including 3% the spark NO_x and 3% of the HONO predicted to be generated in a model run. The blue circles are the average laboratory measurements and average extrapolated value at time zero, while the dashed-dotted blue lines are the individual extrapolated linear fits to the laboratory data. Error bars are the standard deviation from averaging multiple measurements.

Deleted: 3

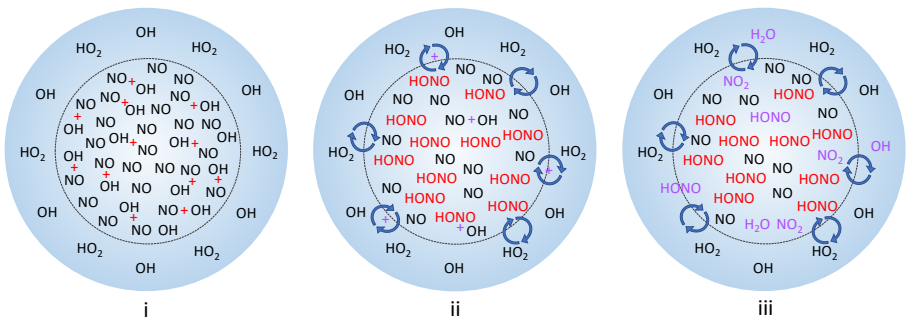


Figure 6: Simplified progression of the proposed HO_x and NO_x chemistry in spark and lightning discharges. (i) Initially, extreme amounts of NO and OH are made inside the lightning hot channel, indicated by the dashed inner circle, while OH and HO_2 are produced outside the hot channel in the corona sheath and UV radiation. (ii) The NO and OH in the hot channel react and form HONO, while the species in the hot channel and corona sheath start to mix together. (iii) Inside the hot channel, any remaining OH reacts with NO and HONO, forming either more HONO or H_2O and NO_2 , respectively. Where the hot channel and corona sheath have started mixing, OH and HO_2 from the corona sheath react with NO from the core, forming HONO or OH and NO_2 , respectively, while OH from the corona sheath and HONO from the core can also react to form H_2O and NO_2 .

450 **4 Conclusions**

Both the laboratory and model results across all the tested pressures confirm that the OH and HO_2 we measure from sparks are generated outside the lightning hot channel, separate from the core where the LNO is generated. It took 3% NO and 3% HONO to resolve the measured-modelled discrepancy in these laboratory experiments, where the sparks occurred in a flow tube with laminar flow and a fast air velocity. In the atmosphere, the percentage of NO or HONO reacting with LHO_x could be lower or higher than 3%, depending on the turbulence and air velocity where the lightning flash occurs, and likely varies from one lightning flash to the next. But the overall conclusion, that the HO_x generated outside the hot channel only partially interacts with the hot channel products, will still be true in the atmosphere.

Additionally, these results indicate only that the substantial LHO_x we measure is generated outside the hot channel; they do not imply that no LHO_x is generated in the hot channel. As stated previously, modelling studies of the lightning hot channel indicate that substantial LHO_x is also generated in the hot channel, likely even more than we measure outside the hot channel.

Deleted: 4

Deleted: likely

Deleted: ¶

465 But this hot channel HO_x will be rapidly titrated away in the presence of the large NO also generated in the core, becoming
substantial HONO. As for the LHO_x we measure outside the hot channel, LHO_x production has been found to be proportional
to ultraviolet radiation (UV) production in corona discharge (Jenkins et al., 2022), and UV may also be responsible for the
LHO_x we measure in sparks and lightning. The consequence of this spatially separate production of LHO_x and LNO is that
470 LHO_x is not immediately consumed by LNO in lightning flashes, but instead is available to oxidize other pollutants in the
atmosphere and contribute to global OH oxidation.

While we did not test the full range of possible tropospheric pressures and temperatures in this study, we still expect that these
results apply for the lower pressures and lower temperatures found in the upper troposphere where most lightning occurs.
Regardless of where it occurs in the troposphere, a lightning flash is composed of a hot core surrounded by a corona sheath
475 and UV radiation, so HO_x and NO_x production is also expected to be spatially separate in the upper troposphere. Our previous
study showed that the initial LNO_x mixing ratio is independent of temperature and only slightly dependent on pressure, with
less than a factor-of-2 difference in production between 970 hPa and 250 hPa, while the initial LHO_x mixing ratio is
independent of pressure and decreases with decreasing temperature, depending on the available water vapor (Jenkins and
Brune, 2023). Therefore, we expect roughly the same LNO_x production in the upper troposphere as was observed in the
480 experiments here, with likely ~200-300 pptv of LHO_x produced. The modelling results showed that for all the pressures tested
in this study, the reaction $OH + NO + M \rightarrow HONO + M$ accounts for over half of the OH loss, while the reaction $HO_2 +$
 $NO \rightarrow OH + NO_2$ accounts for 80% of the HO₂ loss. The rates of these two reactions increases with decreasing temperature,
although the rate of $OH + NO + M$ is also pressure dependent. However, further modelling tests using the lowered LHO_x
production with the same LNO_x as was measured at 360 hPa demonstrate even at 200 hPa and 220 K, the reactions $OH +$
485 $NO + M$ and $HO_2 + NO$ still accounts for more than 50% of the OH loss and 80% of the HO₂ loss, respectively. Thus, based
on this information, we also expect the same subsequent HO_x-NO_x chemistry to occur in the upper troposphere as shown for
the pressures and temperatures here.”

490 Differences in the model and laboratory HO_x decays are resolved if substantial HONO is produced in the spark discharges and
therefore HONO would also be a substantial product of lightning in the atmosphere. Aside from the substantial HONO
production predicted in two lightning chemistry models (Bhettanabhotla et al., 1985; Hill & Rinker, 1981), enhanced HONO
has been measured inside two different electrified convective clouds (Dix et al., 2009; Heue et al., 2014). The field studies
estimate HONO mixing ratios from 37-160 pptv inside the clouds, a range much lower than what the modelling studies predict
and the 1-2 ppbv we expect is made in the laboratory sparks. However, both field cases measured the HONO using differential
495 optical absorption spectroscopy, generating long path measurements that are averaged over the entire length of the cloud, while
higher resolution measurements would likely show very high HONO mixing ratios in lightning affected air and lower mixing
ratios in air lightning did not pass through. We are not aware of any laboratory measurements of electrically produced HONO.

Deleted: ,

Deleted: Comparing the model and laboratory HO_x decays revealed that substantial HONO is a likely product of our spark discharges, and therefore is also likely to be a substantial product of lightning in the atmosphere.

Formatted: Subscript

Measurements of electrically generated HONO, either in the laboratory or at higher spatial resolution in the field, would thus be a good target for future work.

505

Data Availability All data shown in the figures is publicly available at Jenkins and Brunc (2024).

Author Contribution Investigation, Methodology, Visualization, Original Manuscript Draft were by MJJ. Funding Acquisition was by WHB. Conceptualization and Reviewing and Editing of the Manuscript were by WHB and MJJ.

510

Competing Interests The authors declare that they have no conflict of interest.

Acknowledgements We thank P. Stevens for lending us a microchannel plate detector after ours failed.

515

References

Bhetanabhotla, M. N., Crowell, B. A., Coucouvinos, A., Hill, R. D., and Rinker, R. G.: Simulation of trace species production by lightning and corona discharge in moist air, Atmos. Environ., 19, 1391-1397, doi:10.1016/0004-6981(85)90276-8, 1985.

520 Biermann, H. W., Zetzsch, C., and Stuhl, F.: Rate Constant for the reaction of OH with N₂O at 298 K, Berich. Bunsen. Gesell., 80, 909-911, doi:10.1021/i160062a006, 1976.

Boldi, R. A.: A model of the ion chemistry of electrified convection, Ph.D. dissertation, Massachusetts Institute of Technology, 1992.

525 Brandvold, D. K., Martinez, P., and Dogruel, D.: Polarity Dependence of N₂O Formation From Corona Discharge, Atmos. Environ., 23, 1881-1883, doi:10.1016/0004-6981(89)90513-1, 1989.

Brandvold, D. K., Martinez, P., and Hipsh, R. Field measurements of O₃ and N₂O produced from corona discharge, Atmos. Environ., 30, 973-976, doi:10.1016/1352-2310(95)00234-0, 1996.

530 Bruggeman, P., and Schram, D. C.: On OH production in water containing atmospheric pressure plasmas, Plasma Sources Sci. T., 19, 045025, doi:10.1088/0963-0252/19/4/045025, 2010.

Deleted: Unfortunately, there is presently little data on electrically generated HONO. Only one modelling study confirms that fast, substantial HONO formation is possible in the aftermath of a lightning flash (Bhetanabhotla et al., 1985); other modelling studies of lightning flash chemistry do not mention HONO, nor are we aware of any laboratory or field studies measuring electrically produced HONO. The Deep Convective Clouds and Chemistry campaign, where the first LHO₂ measurements were made in the field, also did not deploy any HONO measuring instruments.

Deleted: field or

545 Brune, W. H., and Jenkins, J. M.: Is the reaction rate coefficient for $\text{OH} + \text{HO}_2 \rightarrow$
 $\text{H}_2\text{O} + \text{O}_2$ dependent on water vapor?, *JACS Au*, 4, 4921–4926, doi: 10.1021/jacsau.4c00905, 2024.

Brune, W. H., Jenkins, J. M., Olson, G. A., McFarland, P. J., Miller, D. O., Mao, J., and Ren, X.: Extreme hydroxyl amounts
generated by thunderstorm-induced corona on grounded metal objects, *P. Natl. Acad. Sci. USA*, 119, e2201213119,
550 doi:10.1073/pnas.2201213119, 2022.

Brune, W. H., McFarland, P. J., Bruning, E., Waugh, S., MacGorman, D., Miller, D. O., Jenkins, J. M., Ren, X., Mao, J., and
Peischl, J.: Extreme oxidant amounts produced by lightning in storm clouds, *Science*, 372, 711–715,
doi:10.1126/science.abg0492, 2021.

555 Christian, H. J., Blakeslee, R. J., Boccippio, D. J., Boeck, W. L., Buechler, D. E., Driscoll, K. T., Goodman, S. J., Hall, J. M.,
Koshak, W. J., Mach, D. M., Stewart, M. F.: Global frequency and distribution of lightning as observed from space by the
Optical Transient Detector, *J. Geophys. Res.-Atmos.*, 108, ACL 4-1-ACL 4-15, doi:10.1029/2002JD002347, 2003.

560 [Dix, B., Brenninkmeijer, C. A. M., Frieß, U., Wagner, T., & Platt, U.: Airborne multi-axis DOAS measurements of atmospheric
trace gases on CARIBIC long-distance flights, *Atmos. Meas. Tech.*, 2, 639–652. doi: 10.5194/amt-2-639-2009, 2009.](#)

Donohoe, K. G., Shair, F. H., and Wulf, O. R.: Production of O_3 , NO, and N_2O , in a pulsed discharge at 1 atm. *Ind. Eng. Chem.*
Fund., 16, 208–215, doi:10.1021/i160062a006, 1977.

565 Dyer, M. J., and Crosley, D. R.: Two-dimensional imaging of OH laser-induced fluorescence in a flame, *Opt. Lett.*, 7, 382–
384, doi:10.1364/OL.7.000382, 1982.

Faloona, I. C., Tan, D., Leshner, R. L., Hazen, N. L., Frame, C. L., Simpasa, J. B., Harder, H., Martinez, M., Di Carlo, P., Ren,
570 X., Brune, W. H.: A laser-induced fluorescence instrument for detecting tropospheric OH and HO_2 : Characteristics and
calibration, *J. Atmos. Chem.*, 47, 139–167, doi:10.1023/B:JOCH.0000021036.53185.0e, 2004.

[Heue, K.-P., Riede, H., Walter, D., Brenninkmeijer, C. A. M., Wagner, T., Frieß, U., Platt, U., Zahn, A., Stratmann, G., &
Ziereis, H.: CARIBIC DOAS observations of nitrous acid and formaldehyde in a large convective cloud. *Atmos. Chem. Phys.*,
575 14, 6621–6642. doi: 10.5194/acp-14-6621-2014, 2014.](#)

[Hill, R. D., & Rinker, R. G.: Production of nitrate ions and other trace species by lightning. *J. Geophys. Res.-Oceans*, 86, 3203–
3209. doi: 10.1029/JC086iC04p03203, 1981.](#)

580 Hill, R. D., Rinker, R. G., and Coucouvinos, A.: Nitrous oxide production by lightning, *J. Geophys. Res.-Atmos.*, 89, 1411-1421, doi:10.1029/JD089iD01p01411, 1984.

Jenkin, M. E., Young, J. C., Rickard, A. R.: The MCM v3.3.1 degradation scheme for isoprene, *Atmos. Chem. Phys.*, 15, 11433-11459, doi:10.5194/acp-15-11433-2015, 2015.

585 Jenkins, J. M., Brune, W. H., and Miller, D. O.: Electrical discharges produce prodigious amounts of hydroxyl and hydroperoxyl radicals, *J. Geophys. Res.-Atmos.*, 126, e2021JD034557, doi:10.1029/2021JD034557, 2021.

Jenkins, J. M., Olson, G. A., McFarland, P. J., Miller, D. O., and Brune, W. H.: Prodigious Amounts of Hydrogen Oxides Generated by Corona Discharges on Tree Leaves, *J. Geophys. Res.-Atmos.*, 127, e2022JD036761, doi:10.1029/2022JD036761, 2022.

590 Jenkins, J. M., and Brune, W.H.: Effect of Temperature and Water Droplets on Production of Prodigious Hydrogen Oxides by Electrical Discharges, *J. Geophys. Res.-Atmos.*, 128, e2023JD039362, doi:10.1029/2023JD039362, 2023.

595 Jenkins, J. M., and Brune, W. H.: Spatially separate production of hydrogen oxides and nitric oxide in lightning, *datacommons@psu [dataset]*, <https://doi.org/10.26208/0VND-TQ52>, 2024.

Kalnajs, L. E., & Avallone, L. M.: A novel lightweight low-power dual-beam ozone photometer utilizing solid-state optoelectronics. *J. Atmos. Ocean. Tech.*, 27, 869–880, doi:10.1175/2009JTECHA1362.1, 2010.

600 Levine, J. S., Hughes, R. E., Chameides, W. L., and Howell, W. E.: N₂O and CO Production by electric discharge: Atmospheric implications, *Geophys. Res. Lett.*, 6, 557-559, doi:10.1029/GL006i007p00557, 1979.

605 Ono, R., and Oda, T. Measurement of hydroxyl radicals in pulsed corona discharge. *J. Electrostat.*, 55, 333–342, doi:10.1016/S0304-3886(01)00215-7, 2002.

Orville, R. E.: A High-Speed Time-Resolved Spectroscopic Study of the Lightning Return Stroke: Part II. A Quantitative Analysis, *J. Atmos. Sci.*, 25, 839-851, doi:10.1175/1520-0469(1968)025<0839:AHSTRS>2.0.CO;2, 1968a.

610 Orville, R. E.: A High-Speed Time-Resolved Spectroscopic Study of the Lightning Return Stroke: Part I. A Qualitative Analysis, *J. Atmos. Sci.*, 25, 827-838, doi:10.1175/1520-0469(1968)025<0827:AHSTRS>2.0.CO;2, 1968b.

Rakov, V. A. and Uman, M. A.: *Lightning: Physics and Effects*, Cambridge University Press, ISBN 0-521-03541-4, 687 pp.,
615 2006.

Rehbein, N., and Cooray, V. NO production in spark and corona discharges, *J. Electrostat.*, 51–52, 333–339,
doi:10.1016/S0304-3886(01)00115-2, 2001.

620 Ripoll, J.-F., Zinn, J., Jeffrey, C. A., and Colestock, P. L.: On the dynamics of hot air plasmas related to lightning discharges:
1. Gas dynamics, *J. Geophys. Res.-Atmos.*, 119, 9218–9235, doi:10.1002/2013JD020068, 2014.

Tang, M. J., Cox, R. A., and Kalberer, M.: Compilation and evaluation of gas phase diffusion coefficients of reactive trace
gases in the atmosphere: volume 1. Inorganic compounds, *Atmos. Chem. Phys.*, 14, 9233–9247, doi:10.5194/acp-14-9233-
625 2014, 2014.

Wolfe, G. M., Marvin, M. R., Roberts, S. J., Travis, K. R., and Liao, J.: The Framework for 0-D Atmospheric Modeling
(F0AM) v3.1, *Geosci. Model Dev.*, 9, 3309–3319, doi:10.5194/gmd-9-3309-2016, 2016.

