Second round review

We thank the Referee #2 again for his review. To facilitate the review process we have copied the reviewer comments in black text. Our responses are in regular blue font. We have responded to all the referee comments and made alterations to our paper (in **bold text**). Page and line numbers refer to the *first revised* manuscript.

Referee #2

The authors have largely addressed my concerns in their rebuttal. I am happy that the paper can form the basis of ongoing discussions in the field of OFR deployment and interpretation. I am particularly satisfied with the RTD analysis and its explanation of the deviation from plug flow. I am still slightly concerned by the continued separation of the gaseous and condensed phase processes. I am in full agreement with the authors that "OFR modeling is a subfield in itself, and our group cannot be expected to address every single possible topic". However, where a process can have a substantial influence on the processes that are the subject of a manuscript, then this possibility should be acknowledged.

It can be argued that the two statements in point 2.2 of the authors response:

i) "The presence of aerosols has typically negligible impacts on the gas-phase chemistry" and

ii) ... "gas-phase species have only limited impacts on OA"

are not demonstrably correct for all conditions in OFRs.

R2.5) To rebut i), consider the typical concentrations in diesel emissions. Concentrations of NOx in raw diesel exhaust are typically between 50 and 1000 ppm depending on running conditions and technology (and can be very much higher during transients and below 17 degrees C when EGR is not mandated). Clearly this is the sort of NOx target regime of the current manuscript. Whilst PM emissions do not respond in the same way as NOx to engine technologies (e.g. EGR generally increases PM whilst decreasing NOx, and only DPF fitted vehicles have significantly reduced PM) or load-speed conditions, typical concentrations from a modern light-duty (EURO5) diesel generally range from between 1 and 30 mg/m³ in raw exhaust. Assuming 80 nm modal diameter, 1 mg/m³ will provide a mass transfer rate ("condensation sink") of about 4 s⁻¹ (using an uptake coefficient of unity); so a lifetime of 0.25 s for such a condensing gas (and 30 mg/m³ would give a lifetime of less than 0.01 s). A lower uptake coefficient would obviously lead to a longer lifetime (e.g. 1 s for 0.01 at 30 mg/m³).

Lines 88 to 92 explicitly include OFR conditions where there is a substantial likelihood of such high primary PM mass (an urban tunnel, "where NOx was high enough to be a major OH reactant"... and ... emissions of vehicles, biomass burning, and other combustion sources, "where NO can often be hundreds of ppm"). Looking at the Karjalainen et al., 2016 case presented in Figures 7 and 8, the authors are carrying out calculations under raw, 12 x and 100 x dilution

conditions for gasoline engine emissions. Figure 7 in Karjalainen reported average primary PM values of 0.45 mg/m³ for parts of the test cycle (assumed raw), rising to more than 10 mg/m³ including the SOA from a gasoline engine. Similarly, the Link et al., 2016 study of diesel emissions at 45 - 110 dilution employed no primary particle removal technology to emissions from a turbocharged, intercooled, heavy-duty, off-road diesel engine likely to emit massively more than the light-duty levels stated above (in excess of 100 mg/m³ is readily possible in raw exhaust from such engines). In both these cases the mass transfer of potentially condensing closed shell and radical species to PM could clearly provide very significant sinks of gaseous components that should be considered in a model of OFRs.

We realize that the Referee is not understanding what we intended to communicate with our statements. When we stated that the "gas-phase chemistry" was not significantly perturbed by the presence of particles, we were referring to the gas-phase chemistry that we are modeling, i.e., the radical and NO_x - NO_y chemistry, as well as the consumption of VOCs and other OH reactants. We did not intend to include the physical partitioning of semivolatile and low volatility species, which clearly is an area of strong interaction of the gas and particle phases (but for the most part does not involve chemistry). We will clarify the language, as described below, to remove this potential source of confusion.

We thank the Referee for providing detailed examples for high condensational sink in combustion exhausts. Nevertheless, even if raw exhausts are injected into the reactor, they still cannot have significant impacts on the major gas-phase oxidants, since VOCs and NO_x in raw exhausts, which are also proportionally higher, still dominate total oxidant sink. Dilution of raw exhausts simultaneously lowers condensational sinks and gas-phase reactive oxidant sinks, with their relative importance remaining the same.

We have added a few sentences to clarify this point at the end of the paragraph i) in L151. The modified paragraph now reads:

i) "The presence of aerosols has typically negligible impacts on the gas-phase chemistry of radicals, NO_x/NO_y, and OH reactants studied here. Condensational sink (CS) of ambient aerosols can rarely exceed 1 s⁻¹ even in polluted areas and is usually 1-3 orders of magnitude lower (Donahue et al., 2016; Palm et al., 2016). Thus, even under the assumption of unity uptake coefficient, CS cannot compete with OHR_{ext} (usually on the order of 10 s⁻¹ or higher) in OH loss. Uptake of NO onto aerosols only occurs through the reaction with RO₂ on particle surface (Richards-Henderson et al., 2015), which is formed very slowly (see below) compared to gas-phase HO_x and NO_x chemistry. Uptake of HO₂, O₃, NO₃ etc. is even more unlikely to be of importance due to lower uptake coefficients (Moise and Rudich, 2002; Moise et al., 2002; Hearn and Smith, 2004; Lakey et al., 2015). Combustion exhausts can have high aerosol loadings with condensational sinks on the order of 10²–10³ s⁻¹ (Matti Maricq, 2007). Even if these exhausts are directly injected into the reactor without any pre-treatment, uptake onto the particles still cannot play a major role in the fate of gas-phase radical and NO_x species, since VOCs and NO_x in raw exhausts, which are

proportionally orders-of-magnitude higher, still dominate the fate of oxidants. Dilution of combustion emissions simultaneously lowers condensational sinks and the sinks of oxidants due to chemical reactions, with their relative importance remaining the same as in undiluted emissions."

We acknowledge that a strong dilution leads to a lower PM loading than in raw exhausts, which slows down the uptake. But this "lower" PM loading may still be much higher than typical ambient values after dilution. So it may not necessarily be "low PM loading" compared to ambient conditions, as suggested by the Referee below (R2.8).

We have added some text to L566 to acknowledge the PM loading change by dilution:

"Note that a strong dilution lowers aerosol mass loading in vehicle emissions. As a result, condensation of gases onto particles is slower than in raw exhausts. However, condensational sinks after dilution may still be significantly higher than typical ambient values (Matti Maricq, 2007; Donahue et al., 2016)."

R2.6) To address author response 2.2 ii), clearly gas-phase species have a strong impact on OA, being 100% responsible for all SOA. Mass transfer of semi-volatile and low volatility gas-phase species (in the case of exhaust experiments, almost completely due to condensation on existing primary PM) has a determinant effect on PM mass. Gas phase oxidants may have a limited impact on OA chemistry, but gas phase species have a profound effect on OA. Given the paper title relates to modelling the chemistry in OFRs (not modelling the oxidants), it is not solely transfer of radical species between phases that is of concern.

As discussed in response to R2.5, we agree with this and did not intend to say otherwise. However this mostly concerns physics ("mass transfer" in the Referee's words) and not chemistry. We have modified that paragraph to clarify the fact that physical uptake of semivolatile and low-volatility gas-phase species have a strong impact on PM mass:

"Gas-phase radical and NO_x/NO_y species only has limited impacts on OA *chemistry* in this study. Heterogeneous oxidation of OA by OH is generally slow. Significant OA loss due to heterogeneous oxidation can only be seen at equivalent photochemical ages as high as weeks (Hu et al., 2016). The enhancement of heterogeneous oxidation due to NO is remarkable only at OH concentration close to the ambient values but not at typical values in OFR (Richards-Henderson et al., 2015).

It is well known that the aerosol concentration can have a major impact on the physical uptake of semivolatile and low-volatility gas-phase species. However this process is not explicitly modeled in this study."

Also, we have modified text to L148 for more clarity:

"Also, particle-phase chemistry and physical and chemical interactions of gas-phase species with particles are not considered in this study."

R2.7) In the context of the above discussion, I do not understand the final paragraph of the authors suggested added text in point 2.2. I think this requires further explanation before inclusion in the paper.

The core idea of that paragraph is that the details of VOC product evolution have a strong influence on the rate of OH loss, but this is highly complex and not well captured even by models as explicit as MCM. We surrogated VOC evolution by SO_2 for simplicity and efficiency and this introduces much more uncertainty on (the temporal variation of) OHR_{ext} due to VOC (and hence the radical chemistry) than mass transfer to the particle phase does. Thus, there is no strong need for explicitly including gas-particle mass transfer in the present modeling work.

We have modified that paragraph (L166) below for more clarity:

"As OHRext plays a major and even dominant role in OH loss, it is an important approximation that the real OHRext decay (due to not only primary VOC oxidation and subsequent oxidation of higher generation products, but also wall loss, partitioning to the particle phase, reactive uptake etc.) is surrogated by that of SO₂ (see Fig. S2 of Peng et al. 2015). Gas-phase measurements in literature laboratory studies revealed that there is a large variability of the evolution of total OHR_{ext} during oxidation of primary VOCs and subsequent oxidation of their intermediate products, depending on the type of precursors (Nehr et al., 2014; Schwantes et al., 2017). This variability is obviously mainly due to the formation of different types and amounts of oxidation intermediates/products contributing to OHR_{ext}. This variation is highly complex due to the large number of possible oxidation intermediates and the limited knowledge of detailed higher-generation mechanisms, and thus is difficult to accurately capture even if modeling with a mechanism as explicit as Master Chemical Mechanism is performed (Schwantes et al., 2017). Therefore, it is justified to use a lumped surrogate to model the OHR_{ext} decay for simplicity and efficiency. This approximation is a major contributor to uncertainty of our model. The uncertainties due to both the types of oxidation intermediates/products."

R2.8) Whilst I do not expect the paper to explicitly address coupling of the gaseous and particulate processes, I would expect the current manuscript to at least acknowledge the interaction between the gas phase chemistry and gaseous losses associated with condensation and the resultant increase in PM mass. The authors should state that their study is completely relevant for low PM loadings in OFRs, but care should be taken when applying it to high ambient PM concentration or direct emission studies (both raw and diluted). Clearly the authors are aware of the necessity to include coupled multiphase processes and should be commended in their work with Jeff Pierce's group on this.

We have acknowledged (see the response to R2.5) that dilution changes PM loadings and hence mass transfer rates, and, when diluted, sources have lower PM loadings in OFRs than in raw emissions. However, we believe that the statement that our study is only valid to low PM loadings is incorrect, as discussed in detail in our response to R2.5. Thus no changes have been made in response to this point.

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1 Modeling of the chemistry in oxidation flow reactors with high initial NO

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7 Abstract. Oxidation flow reactors (OFRs) are increasingly employed in atmospheric chemistry research 8 because of their high efficiency of OH radical production from low-pressure Hg lamp emissions at both 9 185 and 254 nm (OFR185) or 254 nm only (OFR254). OFRs have been thought to be limited to studying 10 low-NO chemistry (where peroxy radicals (RO₂) react preferentially with HO₂) because NO is very rapidly 11 oxidized by the high concentrations of O₃, HO₂, and OH in OFRs. However, many groups are performing 12 experiments aging combustion exhaust with high NO levels, or adding NO in the hopes of simulating high-NO chemistry (where RO_2 + NO dominates). This work systematically explores the chemistry in 13 14 OFRs with high initial NO. Using box modeling, we investigate the interconversion of N-containing 15 species and the uncertainties due to kinetic parameters. Simple initial injection of NO in OFR185 can 16 result in more RO2 reacted with NO than with HO2 and minor non-tropospheric photolysis, but only 17 under a very narrow set of conditions (high water mixing ratio, low UV intensity, low external OH 18 reactivity (OHR_{ext}), and initial NO concentration (NOⁱⁿ) of tens to hundreds of ppb) that account for a 19 very small fraction of the input parameter space. These conditions are generally far away from 20 experimental conditions of published OFR studies with high initial NO. In particular, studies of aerosol 21 formation from vehicle emissions in OFR often used OHR_{ext} and NOⁱⁿ several orders of magnitude higher. 22 Due to extremely high OHRext and NOⁱⁿ, some studies may have resulted in substantial non-tropospheric 23 photolysis, strong delay to RO_2 chemistry due to peroxynitrate formation, VOC reactions with NO_3 24 dominating over those with OH, and faster reactions of OH-aromatic adducts with NO₂ than those with 25 O2, all of which are irrelevant to ambient VOC photooxidation chemistry. Some of the negative effects 26 are worst for alkene and aromatic precursors. To avoid undesired chemistry, vehicle emissions generally 27 need to be diluted by a factor of >100 before being injected into OFR. However, sufficiently diluted 28 vehicle emissions generally do not lead to high-NO chemistry in OFR, but are rather dominated by the 29 low-NO RO2+HO2 pathway. To ensure high-NO conditions without substantial atmospherically irrelevant 30 chemistry in a more controlled fashion, new techniques are needed.

31 1 Introduction

The oxidation of gases that are emitted into the atmosphere, in particular volatile organic compounds (VOCs), is one of the most important atmospheric chemistry processes (Haagen-Smit, 1952; Chameides et al., 1988). VOC oxidation is closely related to radical production and consumption (Levy II, 1971), O₃ production, and formation of secondary aerosols (Odum et al., 1996; Hoffmann et al., 1997; Volkamer et al., 2006; Hallquist et al., 2009), which have impacts on air quality and climate (Lippmann, 1991; Nel, 2005; Stocker et al., 2014).

Chemical reactors are critical tools for research of VOC oxidation. Oxidation reactions of interest often have typical timescales of hours to weeks. Studying these processes in ambient air can be confounded by dispersion and changes in ambient conditions, which often occur on similar timescales. Chemical reactors allow for decoupling of these two types of processes. Also, they should be able to simulate the different regimes of reactions occurring in the atmosphere, e.g., VOC oxidation under low and high-NO conditions (peroxy radical fate dominated by reaction with HO₂ or with NO) representing remote and urban areas, respectively (Orlando and Tyndall, 2012).

45 Large environmental chambers are a commonly used reactor type (Carter et al., 2005; Wang et al., 2011). They typically employ actinic wavelength (>300 nm) light sources (e.g., outdoor solar radiation 46 47 and UV blacklights) to produce oxidants and radicals and have large volumes (on the order of several 48 cubic meters or larger). However, the capability of generating sustained elevated levels of OH, the most 49 important tropospheric oxidant, is usually limited in chambers, resulting in OH concentrations similar to those in the atmosphere (10⁶-10⁷ molecules cm⁻³; Mao et al., 2009; Ng et al., 2010), and consequently, 50 51 long simulation times (typically hours) to reach OH equivalent ages of atmospheric relevance (George 52 et al., 2007; Kang et al., 2007; Carlton et al., 2009; Seakins, 2010; Wang et al., 2011). The partitioning of 53 gases and aerosols to chamber walls (usually made of Teflon) in timescales of tens of minutes to hours 54 makes it difficult to conduct very long experiments that simulate high atmospherically-relevant 55 photochemical ages (Cocker et al., 2001; Matsunaga and Ziemann, 2010; Zhang et al., 2014; Krechmer 56 et al., 2016). In addition, the long simulation times and large size of chambers and auxiliary equipment 57 are logistically difficult for field deployment, and their cost limits the number of laboratories equipped 58 with them.

59 Given the limitations of environmental chambers, a growing number of experimenters have 60 instead employed oxidation flow reactors (OFRs). OFRs have a much smaller size (of the order of 10 L), 61 efficiently generate OH via photolysis of H₂O and/or O₃ by more energetic 185 and 254 nm photons 62 from low-pressure Hg lamps, and overcome the abovementioned shortcomings of chambers due to a 63 much shorter residence time (George et al., 2007; Kang et al., 2007, 2011; Lambe et al., 2011). Moreover, 64 OFRs are able to rapidly explore a wide range of OH equivalent ages within a short period (2 hr), during 65 which significant changes of ambient conditions can usually be avoided in the case of field deployment 66 (Ortega et al., 2016; Palm et al., 2016, 2017). Because of these advantages, OFRs have recently been 67 widely used to study atmospheric chemistry, in particular secondary organic aerosol (SOA) formation 68 and aging, in both the laboratory and the field (Kang et al., 2011; Li et al., 2013; Ortega et al., 2013,

69 2016; Tkacik et al., 2014; Palm et al., 2016).

70 In addition to experimental studies using OFRs, there has also been some progress in the 71 characterization of OFR chemistry by modeling. Li et al. (2015) and Peng et al. (2015) developed a box 72 model for OFR HO_x chemistry that predicts measurable quantities [e.g., OH exposure (OH_{exp}, in 73 molecules cm 3 s] and O₃ concentration (abbr. O₃ hereinafter, in ppm)] in good agreement with 74 experiments. This model has been used to characterize HO_x chemistry as a function of H₂O mixing ratio 75 (abbr. H₂O hereinafter, unitless), UV light intensity (abbr. UV hereinafter, in photons cm⁻² s⁻¹), and 76 external OH reactivity [in s⁻¹, OHR_{ext}= $\sum k_i c_i$, i.e., the sum of the products of concentrations of externally 77 introduced OH-consuming species (c_i) and rate constants of their reactions with OH (k_i)]. Based on this 78 characterization, Peng et al. (2015) found that OH suppression, i.e., reduction of OH concentration 79 caused by OHRext, is a common feature under many typical OFR operation conditions. Peng et al. (2016) 80 systematically examined the relative importance of non-OH/non-tropospheric reactants on the fate of VOCs over a wide range of conditions, and provided guidelines for OFR operation to avoid non-81 82 tropospheric VOC photolysis, i.e., VOC photolysis at 185 and 254 nm.

83 In previous OFR modeling studies, NO_x chemistry was not investigated in detail, since in such in 84 typical OFR experiments with large amounts of oxidants (e.g., OH, HO_2 , and O_3), NO would be very rapidly oxidized and thus unable to compete with HO₂ for reaction with peroxy radicals (RO₂). Li et al. 85 (2015) estimated an NO (NO₂) lifetime of ~0.5 (~1.5) s under a typical OFR condition. From these 86 87 estimates, OFRs processing ambient air or laboratory air without large addition of NOx were assumed to be not suitable for studying oxidation mechanisms relevant to polluted conditions under higher NO 88 concentrations. OFRs have recently been used to conduct laboratory experiments with very high initial 89 NO_x levels (Liu et al., 2015) and deployed to an urban tunnel, where NO_x was high enough to be a major 90 91 OH reactant (Tkacik et al., 2014). The former study reported evidence for the incorporation of nitrogen 92 into SOA. Besides, OFRs have been increasingly employed to process emissions of vehicles, biomass 93 burning, and other combustion sources (Table 1), where NO can often be hundreds of ppm (Ortega et 94 al., 2013; Martinsson et al., 2015; Karjalainen et al., 2016; Link et al., 2016; Schill et al., 2016; Alanen et 95 al., 2017; Simonen et al., 2017). It can be expected that such a high NO input together with very high 96 VOC concentrations would cause a substantial deviation from good OFR operation conditions identified 97 in Peng et al. (2016). Very recently, N₂O injection has been proposed by Lambe et al. (2017) as a way to 98 study oxidation of VOCs under high NO conditions in OFR. As more OFR studies at high NO_x level are 99 conducted, there is growing need to understand the chemistry of N-containing species in OFRs and 100 whether it proceeds along atmospherically-relevant channels.

101 In this study, we present the first comprehensive model of OFR NO_V chemistry. We extend the 102 model of Li et al. (2015) and Peng et al. (2015) by including a scheme for NO_V species. Then this model 103 is used to investigate i) if an OFR with initial NO injection results in NO significantly reacting with RO_2 104 under any conditions, ii) if previously published OFR experiments with high initial NO concentrations 105 led to RO_2 +NO being dominant in VOC oxidation without negative side effects (e.g., non-tropospheric 106 reactions), iii) how to avoid undesired chemistry in future studies. The results can provide insights into the design and interpretation of future OH-oxidation OFR experiments with large amounts of NO_xinjection.

109 2 Methods

110 The physical design of the OFR modeled in the present work, the chemical kinetics box model, and 111 the method of propagating and analyzing the parametric uncertainties on the model have already been 112 introduced previously (Kang et al., 2007; Li et al., 2015; Peng et al., 2015). We only provide brief 113 descriptions for them below.

114 2.1 Potential Aerosol Mass flow reactor

115 The OFR modeled in this study is the "Potential Aerosol Mass" (PAM) flow reactor, firstly introduced by Kang et al. (2007). The PAM OFR is a cylindrical vessel with a volume of ~13 L, equipped 116 117 with low-pressure Hg lamps (model no. 82-9304-03, BHK Inc.) to generate 185 and 254 nm UV light. 118 This popular design is being used by many atmospheric chemistry research groups, particularly those 119 studying SOA (Lambe and Jimenez, 2017 and references therein). When the lamps are mounted inside 120 Teflon sleeves, photons at both wavelengths are transmitted and contribute to OH production ("OFR185 mode"). In OFR185, H₂O photolyzed at 185 nm produces OH and HO₂, while O₂ photolyzed at the same 121 122 wavelengths results in O_3 formation. $O(^{1}D)$ is produced via O_3 photolysis at 254 nm and generates additional OH through its reaction with H₂O. 185 nm lamp emissions can be filtered by mounting the 123 124 lamps inside quartz sleeves, leaving only 254 nm photons to produce OH ("OFR254 mode"). In this mode, 125 injection of externally formed O_3 is necessary to ensure OH production. As the amount of O_3 injected is 126 a key parameter under some conditions (Peng et al., 2015), we adopt the notation OFR254-X to denote OFR254 experiments with X ppm initial O_3 ($O_{3,in}$). In this study, we investigate OFR experiments with NO 127 128 injected and thus utilize "OFR185-iNO" to describe the OFR185 mode of operation with initially (at the 129 reactor entrance) injected NO. The same terminology is used for the OFR254 mode. For instance, the 130 initial NO injection into OFR254-7 is denoted as OFR254-7-iNO.

131 2.2 Model description

132 The basic framework of the box model used in this study, a standard chemical kinetics model, is 133 the same as in Peng et al. (2015). Plug flow is assumed in the model, since approximately taking 134 residence time distribution into account leads to similar results under most conditions but at much 135 higher computational expense (Peng et al., 2015). In addition to the reactions in the model of Peng et 136 al. (2015), including all HOx reactions available in the JPL Chemical Kinetic Data Evaluation (Sander et al., 137 2011), all gas-phase NO_y reactions available in the JPL database except those of organic nitrates and 138 peroxynitrates are also considered in the current reaction scheme. An updated JPL evaluation was published recently (Burkholder et al., 2015), with slightly different (~20%) rate constants for 139 140 $NO_2+HO_2+M \rightarrow HO_2NO_2+M$ and $NO_2+NO_3 \rightarrow N_2O_5$. The updated rate constants only result in changes of 141 ~10-20% of the concentrations of the species directly consumed/produced by these reactions. These 142 changes are smaller than the parametric uncertainties of the model (see Section 3.1.3). For other species, concentration changes are negligible. $HO_2NO_2+M\rightarrow HO_2+NO_2+M$ and $N_2O_5+M\rightarrow NO_2+NO_3+M$, 143 144 are also included in the scheme, with kinetic parameters from the IUPAC Task Group on Atmospheric

145 Chemical Kinetic Data Evaluation (Ammann et al., 2016). As in Peng et al. (2015, 2016), SO₂ is used as a 146 surrogate of external OH reactants (e.g., VOCs). NOy species, although also external OH reactants, are 147 explicitly treated in the model and not counted in OHRext in this work. Therefore, OHRext stands for non-148 NOy OHRext only hereinafter, unless otherwise stated. 149 Also, particle-phase chemistry and physical and chemical interactions of gas-phase species with 150 particles are not considered in this study. We have made this assumption because: 151 The presence of aerosols has typically negligible impacts on the gas-phase chemistry of radicals, 152 NO_x/NO_y, and OH reactants studied here. Condensational sink (CS) of ambient aerosols can 153 rarely exceed 1 s⁻¹ even in polluted areas and is usually 1-3 orders of magnitude lower 154 (Donahue et al., 2016; Palm et al., 2016). Thus, even under the assumption of unity uptake 155 coefficient, CS cannot compete with OHR_{ext} (usually on the order of 10 s⁻¹ or higher) in OH loss. 156 Uptake of NO onto aerosols only occurs through the reaction with RO2 on particle surface 157 (Richards-Henderson et al., 2015), which is formed very slowly (see below) compared to gas-158 phase HOx and NOx chemistry. Uptake of HO2, O3, NO3 etc. is even more unlikely to be of 159 importance due to lower uptake coefficients (Moise and Rudich, 2002; Moise et al., 2002; 160 Hearn and Smith, 2004; Lakey et al., 2015). Combustion exhausts can have high aerosol 161 loadings with condensational sinks on the order of 10²-10³ s⁻¹ (Matti Maricq, 2007). Even if 162 these exhausts are directly injected into the reactor without any pre-treatment, uptake onto 163 the particles still cannot play a major role in the fate of gas-phase radical and NO_x species, since 164 VOCs and NOx in raw exhausts, which are proportionally orders-of-magnitude higher, still 165 dominate the fate of oxidants. Dilution of combustion emissions simultaneously lowers 166 condensational sinks and the sinks of oxidants due to chemical reactions, with their relative 167 importance remaining the same as in undiluted emissions. 168 Gas-phase radical and NO_x/NO_y species only has limited impacts on OA chemistry in this study. 169 Heterogeneous oxidation of OA by OH is generally slow. Significant OA loss due to 170 heterogeneous oxidation can only be seen at photochemical ages as high as weeks (Hu et al., 171 2016). The enhancement of heterogeneous oxidation due to NO is remarkable only at OH 172 concentration close to the ambient values but not at typical values in OFR (Richards-Henderson 173 et al., 2015). 174 It is well known that the aerosol concentration can have a major impact on the physical uptake of Deleted: It 175 semivolatile and low-volatility gas-phase species. However this process is not explicitly modeled in this 176 study. 177 As OHRext plays a major and even dominant role in OH loss, it is an important approximation that 178 the real OHRext decay (due to not only primary VOC oxidation and subsequent oxidation of higher 179 generation products, but also wall loss, partitioning to the particle phase, reactive uptake etc.) is 180 surrogated by that of SO2_ (see Fig. S2 of Peng et al. 2015). Gas-phase measurements in literature Deleted: 181 laboratory studies revealed that there is a large variability of the evolution of total OHRext during 182 pxidation of primary VOCs and subsequent oxidation of their intermediate products, depending on the

Deleted: Also, particle-phase processes and interactions of gasphase species with particles are not considered in this study. We have made this assumption because:

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192 type of precursors (Nehr et al., 2014; Schwantes et al., 2017). This variability is obviously mainly due to 193 the <u>formation</u> of different types and amounts of oxidation intermediates/products contributing to 194 OHRext, This variation is highly complex due to the large number of possible oxidation intermediates and

the limited knowledge of detailed higher-generation mechanisms, and thus is difficult to accurately 196 capture even if modeling with a mechanism as explicit as Master Chemical Mechanism is performed

197 (Schwantes et al., 2017). Therefore, it is justified to use a lumped surrogate to model the OHRext decay

198 for simplicity and efficiency. This approximation is a major contributor to uncertainty of our model. The

199 uncertainties due to both the types of oxidation intermediates/products,

195

200 A residence time of 180 s and typical temperature (295 K) and atmospheric pressure (835 mbar) 201 in Boulder, CO, USA are assumed for all model cases. The lower-than-sea level pressure only leads to 202 minor differences in the outputs (Li et al., 2015). We explore physical input cases evenly spaced in a 203 logarithmic scale over very wide ranges: H₂O of 0.07%-2.3%, i.e., relative humidity (RH) of 2-71% at 204 295 K; 185 nm UV of 1.0x10¹¹-1.0x10¹⁴ and 254 nm UV of 4.2x10¹³-8.5x10¹⁵ photons cm⁻² s⁻¹; OHRext of 205 1-16000 s⁻¹; O_{3,in} of 2.2-70 ppm for OFR254; initial NO mixing ratio (NOⁱⁿ) from 10 ppt to 40 ppm. 206 Besides, conditions with OHR_{ext}=0 are also explored. UV at 254 nm is estimated from that at 185 nm 207 according to the relationship determined by Li et al. (2015). Several typical cases within this range as well as their corresponding 4 or 2-character labels (e.g., MMOV and HL) are defined in Table 2. Literature 208 209 studies are modeled by adopting all reported parameters (e.g., residence time, H_2O , and $O_{3,in}$) and 210 estimating any others that may be needed (e.g., UV) from the information provided in the papers.

211 In this study, OH equivalent ages are calculated under the assumption of an ambient OH concentration of 1.5x10⁶ molecules cm⁻³ (Mao et al., 2009). Conditions leading to a ratio of RO₂ reacted 212 213 with NO over the entire residence time [r(RO₂+NO)] to that with HO₂ [r(RO₂+HO₂)] larger than 1 are 214 regarded as "high NO" (under the assumption of constant OHR_{ext} from VOCs, see Section S1 for more 215 details), where [r(X)] is the total reactive flux for reaction X over the entire residence time. F185_{exp}/OH_{exp} 216 and $F254_{exp}/OH_{exp}$ are used as measures of the relative importance of VOC photolysis at 185 and 254 217 nm to their reactions with OH, respectively [F185_{exp} (F254_{exp}) are 185 (254) nm photon flux exposure, 218 i.e., product of 185 (254) nm photon flux and time]. Readers may refer to Figs. 1 and 2 of Peng et al. 219 (2016) for the determination of the relative importance of non-tropospheric (185 and 254 nm) 220 photolysis of individual VOCs. Although the relative importance of non-tropospheric photolysis depends 221 on individual VOCs, in the present work, we set criteria on $F185_{exp}/OH_{exp}<3x10^3$ cm/s and 222 $F254_{exp}/OH_{exp}<4x10^5$ cm/s to define "good" conditions and $F185_{exp}/OH_{exp}<1x10^5$ cm/s and 223 $F254_{exp}/OH_{exp}<1x10^7$ cm/s (excluding good conditions) to define "risky" conditions. Conditions with higher F185_{exp}/OH_{exp} or F254_{exp}/OH_{exp} are defined as "bad". Under good conditions, photolysis of most 224 225 VOCs has a relative contribution <20% to their fate; under bad conditions, non-tropospheric photolysis is likely to be significant in all OFR experiments, since it can hardly be avoided for oxidation 226 227 intermediates, even if the precursor(s) does not photolyze at all. Under risky conditions, some species 228 photolyzing slowly and/or reacting with OH rapidly (e.g., alkanes, aldehydes, and most biogenics) still 229 have a relative contribution of photolysis <20% to their fates, while species photolyzing more rapidly

Deleted: evolution Deleted: . but not Deleted: changes in CS, wall conditions etc. Also this variability Deleted: MCM Deleted: It Deleted: thus Deleted: introduced by this approximation include those

Deleted: and all interactions of VOCs with aerosols, walls etc. And the uncertainties due to the former dominate over those due to the latter

240 and/or reacting with OH more slowly (e.g., aromatics and other highly conjugated species and some

saturated carbonyls) will undergo substantial non-tropospheric photolysis. Note that these definitions
are slightly different than in Peng et al. (2016). All definitions of the types of conditions are summarized

in Table 3.

244 2.3 Uncertainty analysis

245 We apply the same method as in Peng et al. (2014, 2015) to calculate and analyze the output 246 uncertainties due to uncertain kinetic parameters in the model. Random samples following log-normal 247 distributions are generated for all rate constants and photoabsorption cross sections in the model using 248 uncertainty data available in the JPL database (Sander et al., 2011) or estimated based on IUPAC data 249 (Ammann et al., 2016). Then, Monte Carlo Uncertainty Propagation (BIPM et al., 2008) is performed for 250 these samples through the model to obtain the distributions of outputs. Finally, we compute squared 251 correlation coefficients between corresponding input and output samples and apportion the relative 252 contributions of individual kinetic parameters to the output uncertainties based on these coefficients 253 (Saltelli et al., 2005).

254 3 Results and discussion

In this section, we study the NO_y chemistry in OFR while considering relevant experimental issues.
 Based on these results, we propose some guidelines for OFR operation for high-NO OH oxidation of
 VOCs.

258 3.1 NO_y chemistry in typical OFR cases with initial NO injection

NO was thought to be unimportant (i.e., unable to significantly react with RO₂) in OFRs with initial NO injection (OFR-iNO) based on the argument that its lifetime is too short due to large amounts of O₃ OH, and HO₂ to compete with RO₂+HO₂ (Li et al., 2015). We evaluate this issue below by calculating NO effective lifetime (τ_{NO} , in s), defined as NO exposure (NO_{exp}, in molecules cm⁻³ s) divided by initial NO concentration, under various conditions. This definition cannot effectively capture the true NO average lifetime if it is close to or longer than the residence time. In this case, τ_{NO} close to the residence time will be obtained, which is still long enough for our characterization purposes.

266 3.1.1 OFR185-iNO

267 In OFR185-iNO, NO is not oxidized extremely quickly under all conditions. For instance, under a 268 typical condition in the midrange of the phase space shown in Fig. 1a, τ_{NO} ~13 s. This lifetime is much shorter than the residence time, but long enough for OH_{exp} to reach ~3x10¹⁰ molecules cm⁻³ s, which is 269 270 equivalent to an OH equivalent age of ~6 hrs. Such an OH equivalent age is already sufficient to allow 271 some VOC processing and even SOA formation to occur (Lambe et al., 2011; Ortega et al., 2016). Within 272 τ_{NO} , NO suppresses HO₂ through the reaction NO+HO₂ \rightarrow NO₂+OH, leading to NO_{exp}/HO_{2exp} of ~700 during 273 this period, high enough for RO₂ to dominantly react with NO. Meanwhile, NO+HO₂→NO₂+OH enhances 274 OH production, which helps OH_{exp} build up in a relatively short period. In addition, non-tropospheric 275 photolysis of VOCs at 185 and 254 nm is minor (F185_{exp}/OH_{exp} ~ 600 cm/s, Fig. 1a), because of enhanced 276 OH production and moderate UV. Therefore, such an OFR condition may be of some interest for high-277 NO VOC oxidation. We thus analyze the NO_v chemistry in OFR185-iNO in more detail below, by taking

278 the case shown in Fig. 1a as a representative example.

279 In OFR185-iNO, HO_x concentrations are orders-of-magnitude higher than in the atmosphere 280 while the amount of O₃ produced is relatively small during the first several seconds after the flow enters 281 the reactor. As a result, NO is not oxidized almost exclusively by O3 as in the troposphere, but also by 282 OH and HO₂ to form HONO and NO₂, respectively (Fig. 1a). The large concentration of OH present then oxidizes HONO to NO₂, and NO₂ to HNO₃. Photolysis only plays a negligible role in the fate of HONO and 283 284 NO₂ in OFRs, in contrast to the troposphere, where it is the main fate of these species. This is because 285 the reactions of HONO and NO2 with OH are greatly accelerated in OFR compared to those in the 286 troposphere, while photolysis not (Peng et al., 2016). The interconversion between NO₂ and HO₂NO₂ is 287 also greatly accelerated (Fig. 1a), since a large amount of HO₂ promotes the formation of HO₂NO₂, 288 whose thermal decomposition and reaction with OH in turn enhance the recycling of NO2. Though not 289 explicitly modeled in this study, RO2 are expected to undergo similar reactions with NO2 to form 290 reservoir species, i.e., peroxynitrates (Orlando and Tyndall, 2012). Peroxynitrates that decompose on 291 timescales considerably longer than OFR residence times may serve as effectively permanent NO_v sinks 292 in OFRs (see Section 3.4.1).

293 Interestingly but not surprisingly, the NO_v chemistry shown in Fig. 1a is far from temporally 294 uniform during the OFR residence time (Fig. S1a). Within τ_{NO} , NO undergoes an e-fold decay as it is 295 rapidly converted into NO2 and HONO, whose concentrations reach maxima around that time. After 296 most NO is consumed, HONO and NO₂ also start to decrease, but significantly more slowly than NO, 297 since they do not have as many and efficient loss pathways as NO. The reaction of OH with HONO, the dominant fate of HONO, is slower than that with NO (Fig. 1a). The net rate of the NO2-to-HO2NO2 298 conversion becomes low because of the relatively fast reverse reaction (Fig. 1a). Besides, the total loss 299 300 of NO₂ is partially offset by the production from HONO. The generally stable concentrations of HONO and NO₂ (Fig. S1a) result in their respective reaction rates with OH that are comparable during and after 301 302 τ_{NO} (Fig. 1a), as OH variation is also relatively small during the entire residence time (Fig. S1b). However, 303 the NO₂-to-HO₂NO₂ conversion after τ_{NO} is much faster than during it (Fig. 1a), resulting from 304 substantially decreased NO and HO₂ concomitantly increasing >1 order of magnitude after τ_{NO} (Fig. 305 S1a,b). HNO₃ and HO₂NO₂, which are substantially produced only after NO₂ is built up, have much higher 306 concentrations later than within τ_{NO} .

307 Under other OFR185-iNO conditions than in Fig. 1a, the major reactions interconverting NOv 308 species are generally the same, although their relative importance may vary. At lower NOⁱⁿ, the 309 perturbation of HO_x chemistry caused by NO_y species is smaller. Effects of NOⁱⁿ less than 1 ppb (e.g., typical non-urban ambient concentrations) are generally negligible regarding HO_x chemistry. Regarding 310 311 NO_y species, the pathways in Fig. 1a are still important under those conditions. At higher NOⁱⁿ (e.g., >1 312 ppm), one might expect NO₃ and N₂O₅ to play a role (as in OFR254-iNO; see Section 3.1.2 below), since 313 high NO_y concentrations might enhance self/cross reactions of NO_y. However, this would not occur 314 unless OH production is high, since relatively low O₃ concentrations in OFR185-iNO cannot oxidize NO₂ 315 to NO₃ rapidly. Also, a large amount of NO_v can lead to significant OH suppression. That would in turn

 $\label{eq:slow} 316 \qquad \text{slow down the NO}_3 \ \text{production from HNO}_3 \ \text{by OH}. \ \text{This is especially true when an OFR is used to oxidize}$

317 the output of highly concentrated sources (e.g., from vehicle exhausts). When sources corresponding

318 to OHR_{ext} of thousands of s⁻¹ and NO^{in} of tens of ppm are injected into OFR185 (Fig. 1b), they essentially

319 inhibit active chemistry except NO consumption, as all subsequent products are much less abundant

320 compared to remaining NO (Fig. S1c).

321 3.1.2 OFR254-iNO

322 The ppm-level O_{3,in} used in the OFR254-iNO mode of operation has a strong impact on its NO_y 323 chemistry. An O_{3,in} of 2.2 ppm (lowest in this study) is already enough to shorten τ_{NO} to ~1 s, preventing 324 NO from playing a role in the chemistry under most explored conditions. The reaction fluxes under a 325 typical $O_{3,in}$ of 7 ppm are shown in Fig. 1c. A reactive flux from NO+O₃ \rightarrow NO₂ makes the reaction of NO 326 with other oxidants (OH, HO₂ etc.) negligible. The HNO₃ production pathway from NO₂ is similar to that 327 in OFR185-iNO. The interconversion between NO2 and HO2NO2 is also fast over the residence time, and 328 even faster than in OFR185-iNO during τ_{NO} , since a high concentration of O₃ also controls the OH-HO₂ interconversion and makes HO2 more resilient against suppression due to high NO (Fig. S1f; Peng et al., 329 330 2015). A major difference in the NO_y chemistry in OFR254-iNO (Fig. 1c) compared to OFR185-iNO (Fig. 331 1a) is significant NO₃/N₂O₅ chemistry due to high O₃ in OFR254-iNO, which accelerates the oxidation of 332 NO_2 to NO_3 . Interconversion between NO_2+NO_3 and N_2O_5 also occurs to a significant extent because of 333 high NO_2 . Under the conditions of Fig. 1c, NO_3 can also be significantly consumed by HO_2 . Unlike 334 OFR185-iNO, OFR254-iNO can substantially form NO₃ from HNO₃ under conditions that are not on the 335 extremes of the explored physical condition space, e.g., at higher UV and lower NOⁱⁿ (e.g., Fig. S2). In 336 the case of very high NOⁱⁿ (equal to or higher than $O_{3,in}$), all O_3 can be rapidly destroyed by NO. As a 337 consequence, OH production is shut down and these cases are of little practical interest (Fig. S3h).

338 3.1.3 Uncertainty analysis

339 The results of uncertainty propagation confirm that the output uncertainties due to uncertain 340 kinetic parameters are relatively low compared to other factors (e.g., non-plug flow in OFR; Peng et al., 341 2015) and the overall model accuracy compared to experimental data (a factor of 2-3; Li et al., 2015). 342 For OFR185-iNO, NO, NO₃, and OH exposures have relative uncertainties of ~0–20%, ~40–70%, and ~15– 343 40%, respectively. The uncertainties in OH exposure are very similar to those in the cases without NO_x 344 (Peng et al., 2015). The contribution of NO_y reactions to OH_{exp} uncertainty is negligible, except for some 345 contribution of OH+NO \rightarrow HONO in a few cases with high NOⁱⁿ (Fig. 2). The uncertainties on NO_{exp} are 346 dominated by the reactions producing HO_x and O₃, i.e., the major consumers of NO. For NO₃ exposure, 347 a few major production and loss pathways (e.g., $NO_2+NO_3 \rightarrow N_2O_5$, $N_2O_5 \rightarrow NO_2+NO_3$, and $HO_2+NO_3 \rightarrow OH+NO_2+O_2)$ dominate its uncertainties. OFR254-iNO has a simpler picture of parametric 348 349 uncertainties in terms of composition. O₃ controls the NO oxidation under most conditions and this 350 reaction contributes most of output uncertainties for NO exposures. $HO_2+NO_3 \rightarrow OH+NO_2+O_2$ dominates 351 the uncertainty on NO₃ exposure. The levels of those uncertainties are lower than in OFR185-iNO (<2% 352 for NO exposure; <60% in all cases and <25% in most cases for NO₃ exposure). Thus, model uncertainties 353 in OFR254-iNO are not shown in detail.

354 3.2 Different conditions types

Having illustrated the main NO_v chemical pathways for typical cases, we present the results of the exploration of the entire physical parameter space (see Section 2.2). Note that the explored space is indeed very large and gridded logarithmically uniformly in every dimension. Therefore, the statistics of the exploration results can be useful to determine the relative importance of the conditions types defined in Section 2.2 and Table 3.

360 It has been shown that during τ_{NO} , RO₂ can react dominantly with NO (Section 3.1.1), while to 361 determine if a condition is high-NO (see Table 3), the entire residence time is considered. This is done 362 because for VOC oxidation systems of interest, there will be significant oxidation of the initial VOC and 363 its products under low-NO conditions, if τ_{NO} is shorter than the reactor residence time. After most NO 364 is consumed, the longer the remaining residence time, the more RO2 will react with HO2 and the more 365 likely that an input condition is classified as low-NO. For a condition to be high-NO, a significantly long τ_{NQ} is required. Figure 3 shows the fractional occurrence distribution of good/risky/bad conditions in 366 367 the entire explored condition space over logarithm of r(RO₂+NO)/r(RO₂+HO₂), which distinguishes high-368 and low-NO conditions. In OFR254-iNO, τ_{NO} is so short that no good high-NO condition is found in the 369 explored range in this study (Fig. 3a). A fraction of explored conditions are bad high-NO. These 370 conditions result from a full consumption of O₃ by NO. Then very little HO_x is produced (right panels in 371 Fig. S3h), but the fate of any RO₂ formed is dominated by RO₂+NO (right panels in Fig. S3i). However, 372 also due to negligibly low OH concentration, little RO2 is produced and non-tropospheric photolysis of 373 VOCs is also substantial compared to their reaction with OH under these conditions, classifying all of 374 them as "bad" (Fig. 3a).

375 In OFR185-iNO, in addition to the typical case shown in Fig. 1a, many other cases have a τ_{NO} of 376 ~10 s or longer (Figs. S3b and S4), which allow the possibility of high-NO conditions. Indeed, ~1/3 of 377 explored conditions in OFR185-iNO with a residence time of 3 min are high-NO (Fig. 3b). Most of these 378 high-NO conditions are also classified as bad, similar with those in OFR254-iNO. More importantly, in 379 contrast to OFR254-iNO, good and risky high-NO conditions also comprise an appreciable fraction of 380 the OFR185-iNO conditions. It is easily expected that very high OHR_{ext} and NOⁱⁿ lead to bad high-NO 381 conditions (all panels in Fig. 4), since they strongly suppress HO_x, which yields bad conditions and in 382 turn keep NO destruction relatively low. Besides, the occurrence of bad high-NO conditions is reduced 383 at high UV (bottom panels in Fig. 4), which can be explained by lowered NO due to high O₃ production 384 and fast OH reactant loss due to high OH production. Good high-NO conditions are rare in the explored 385 space. They are only 1.1% of total explored conditions (Fig. 3b) and present under very specific conditions, i.e., higher H₂O, lower UV, lower OHR_{ext}, and NOⁱⁿ of tens to hundreds of ppb (Figs. 4 and 386 387 S5). Since a very high NO can suppress OH, to obtain both a significant NO level and a good conditions, NOⁱⁿ can only be tens to hundreds of ppb. As NOⁱⁿ is lower and OH is higher than under bad high-NO 388 conditions, UV should be lower than bad high-NO conditions to keep a sufficiently long presence of NO. 389 Thus, UV at 185 nm for good high-NO conditions are generally lower than 10¹² photons cm⁻² s⁻¹ (Fig. S5). 390 391 In addition, a low OHR_{ext} (generally <50 s⁻¹) and a higher H_2O (the higher the better, although there is 10

no apparent threshold) are also required for good high-NO conditions (Fig. S5), as Peng et al. (2016)
pointed out. Risky high-NO conditions often occur between good and bad high-NO conditions, e.g., at
lower NOⁱⁿ than bad conditions (e.g., Cases ML, MM, HL, and HM in Fig. 4, see Table 2 for the typical
case label code), at higher OHR_{ext} and/or NO_{in} than good conditions (e.g., Cases ML and MM), and at
lower H₂O than good conditions (e.g., Case LL).

The trend of the distributions of good, risky, and bad low-NO conditions is generally in line with the analysis in Peng et al. (2016). For low-NO conditions, NO_y species can be simply regarded as external OH reactants, as in Peng et al. (2016). As H_2O decreases and/or OHR_{ext} or NOⁱⁿ increases, a low-NO condition becomes worse (good->risky->bad) (Figs. 4 and 5). In OFR185-iNO, increasing UV generally makes a low-NO condition better because of an OH production enhancement (Fig. 4); while in OFR254iNO, increasing UV generally makes a low-NO condition worse (Fig. 5), since at a higher UV, more O₃ is destroyed and the resilience of OH to suppression is reduced.

404 As discussed above, the fraction of high-NO conditions also depends on OFR residence time. A 405 shorter residence time is expected to generally lead to a larger fraction of high-NO conditions, since the 406 time spent in the reaction for $t > \tau_{NO}$ is significantly smaller. Thus, we also investigate an OFR185-iNO 407 case with a residence time of 30 s. In Fig. 3b, compared to the case with a residence time of 3 min, the 408 distributions of all condition types (good/risky/bad) of the 30 s residence time case shift toward higher 409 $r(RO_2+NO)/r(RO_2+HO_2)$. Nevertheless, shortening the residence time also removes the period when the 410 condition is better (i.e., less non-tropospheric photolysis), when external OH reactants have been partially consumed and OH suppression due to $\mathsf{OHR}_{\mathsf{ext}}$ has been reduced later in the residence time. As 411 412 a result, the fractions of good and risky conditions decrease (Fig. 3b). With the two effects (higher 413 r(RO2+NO)/r(RO2+HO2) and more significant non-tropospheric photolysis) combined, the fraction of 414 good high-NO conditions increases by a factor of ~3. An even shorter residence time does not result in 415 a larger good high-NO fraction, since the effect of enhancing non-tropospheric photolysis is even more 416 apparent.

417 3.3 Effect of non-plug flow

418 We performed model runs where the only change with respect to our box model introduced in 419 Section 2.2 is that the plug-flow assumption is replaced by the residence time distribution (RTD) 420 measured by Lambe et al. (2011) (also see Fig. S8 of Peng et al. (2015)). The chemistry of different air 421 parcels with different residence times is simulated by our box model and outputs are averaged over the 422 RTD. Lateral diffusion between different air parcels is neglected in these simulations.

423 OH_{exp} calculated from the mode with RTD (OH_{exp,RTD}) is higher than that calculated from the plug-424 flow model (OH_{exp,PF}) in both OFR185-iNO and OFR254-iNO (Table 4 and Fig. S6). Under most explored 425 conditions deviations are relatively small, which leads to an overall positive deviation of OH_{exp,RTD} from 426 OH_{exp,PF} by ~x2 (within the uncertainties of the model and its application to real experimental systems). 427 For OFR185-iNO, most conditions (~90%) in the explored space lead to <x3 differences between OH_{exp,PF} 428 and OH_{exp,RTD}, while for a small fraction of cases the differences can be larger (Fig. S6). The larger 429 deviations are mainly present at high UV, OHR_{ext}, and NOⁱⁿ, where conditions are generally "bad" and in 430 which experiments are of little atmospheric relevance. Under these specific conditions, external OH 431 reactants and NO_y can be substantially destroyed for the air parcels with residence times longer than 432 the average, while this is not the case for the average residence time. This feature was already described 433 by Peng et al. (2015) (see Fig. S10 of that study). Although only non-NO_y external OH reactants were 434 considered in that study, the results are the same. In the present study, a higher upper limit of the 435 explored OHRext range (compared to Peng et al., 2015, due to trying to simulate extremely high OHRext used in some recent literature studies) large amounts of NO_y and cause somewhat larger deviations. In 436 437 OFR254-iNO, OH is less suppressed at high OHRext and NOⁱⁿ than in OFR185-iNO because of high O₃ 438 (Peng et al., 2015), OH_{exp.RTD} deviations from OH_{exp.PF} are also smaller (Table 4).

439 Based on the outputs of the model with RTD, similar mapping of the physical input space as Figs. 440 4 and 5 can be done (Figs. S7 and S8). Overall, the mapping of the RTD model results is very similar with 441 that of the plug-flow model. The conditions appear to be only slightly better in a few places of the explored space than those from the plug-flow model, which can be easily explained by the discussions 442 443 above. Besides, the mapping in Figs. S7 and S8 also appear to be slightly more low-NO, for the same reasons discussed above. After NO is destroyed at long residence times, HO₂, suppressed by NO, also 444 445 recovers as OH. r(RO₂+NO)/r(RO₂+HO₂) is obviously expected to be smaller than in the plug-flow model 446 in general.

447 Note that most conditions that appear to be better in the RTD model results are already 448 identified as bad by the plug-flow model. Those conditions look slightly better only because of their 449 better *RTD-averaged* F185_{exp}/OH_{exp} and F254_{exp}/OH_{exp}. However, each of those cases is actually 450 composed of both a better part at longer residence times and also a worse part at shorter residence 451 times. Under those conditions, the reactor simultaneously works in two distinct regimes, one of which 452 is bad due to heavy OH suppression. Such conditions are obviously not desirable for OFR operation.

453 3.4 Possible issues related to high-NO_x levels

In the discussion above, we focused on obtaining high-NO conditions and considered only one experimental issue (non-tropospheric photolysis) that had been previously investigated in Peng et al. (2016) and is not specific for experiments with high NO injection. We discuss additional potential reasons why the OFR-iNO chemistry can deviate strongly from tropospheric conditions, as specifically related to high-NO_x level in this subsection.

459 3.4.1 NO2

460 NO2 reacts with RO2 to form peroxynitrates, generally regarded as reservoir species in the 461 atmosphere as most of them thermally decompose very quickly compared to atmospheric time scales. 462 However, in OFRs, with residence times on the order of minutes, some peroxynitrates may no longer be 463 considered as fast decomposing. This is especially true for acylperoxy nitrates, whose lifetimes can be 464 hours at room temperature (Orlando and Tyndall, 2012). Acylperoxy nitrates are essentially sinks instead 465 of reservoirs in OFRs for both NO2 and RO2. RO2 is estimated to be as high as several ppb in OFRs by our model (e.g., ~6 ppb RO₂ in OFR185 at H₂O=1%, UV at 185 nm=1x10¹³ photons cm⁻² s⁻¹, OHR_{ext}=1000 s⁻¹, 466 467 and NOⁱⁿ=0), while high-NO experiments can yield far higher NO₂. If all RO₂ were acylperoxy, the RO₂ 468 chemistry could be rapidly shut down by NO₂, as rate constants of these RO₂ + NO₂ reactions are around 469 10⁻¹¹ cm³ molecule⁻¹ s⁻¹ (Orlando and Tyndall, 2012). Nevertheless, acylperoxy nitrates are not expected 470 to typically be the dominant component of peroxynitrates, since acyl radicals are not a direct oxidation 471 product of most common VOCs and can only be formed after several steps of oxidation (Atkinson and 472 Arey, 2003; Ziemann and Atkinson, 2012). Most alkylperoxy nitrates retain their short-lived reservoir 473 characteristics in OFRs due to their relatively short thermal decomposition time scales (on the order of 474 0.1 s; Orlando and Tyndall, 2012). Even so, OFR experiments can be seriously hampered at extremely 475 high NO2. If NO2 reaches ppm levels, the equilibrium between RO2+NO2 and alkylperoxy nitrate 476 $(RO_2+NO_2\leftrightarrow RO_2NO_2)$ is greatly shifted toward the alkylperoxy nitrate side, as the forward and reverse rate constants are on the order of 10⁻¹² cm³ molecule⁻¹ s⁻¹ and 1 s⁻¹, respectively (Orlando and Tyndall, 477 478 2012). This results in a substantial decrease in effective RO₂ concentration, or in other words, a 479 substantial slow-down of RO₂ chemistry.

480 Parts per million levels of NO₂ may impose an additional experimental artifact in the oxidation 481 chemistry of aromatic precursors, OH-aromatic adducts, i.e., the immediate products of aromatic 482 oxidation by OH, undergo addition of O₂ and NO₂ at comparable rates under ppm levels of NO₂ (rate constants of the additions of O_2 and NO_2 are on the order of 10^{-16} and 10^{-11} molecules cm⁻³ s⁻¹, 483 respectively ;Atkinson and Arey, 2003). However, only the former addition is atmospherically relevant 484 485 (Calvert et al., 2002). Liu et al. (2015) performed OFR254-iNO experiments with toluene over a range of 486 NOⁱⁿ of 2.5–10 ppm, encompassing the NO concentration range at which the reactions of OH-toluene 487 adduct with O_2 and with NO_2 are of equal importance (~5 ppm; Atkinson and Arey, 2003). This suggests 488 that nitroaromatics, whose formation was reported in the study of Liu et al. (2015), might have been formed in substantial amounts in that study through the addition of NO₂ to the OH-toluene adduct. 489

490 **3.4.2** NO₃

As discussed in Section 3.1, NO₃ can be formed in significant amounts in OFRs with high NO injection. Although NO₃ is also present in the atmosphere, especially during nighttime, significant VOC oxidation by both OH and NO₃ results in more complex chemistry that may complicate the interpretation of experimental results. NO₃ oxidation-only OFR has been previously realized experimentally via thermal dissociation of injected N₂O₅ (Palm et al., 2017). We discuss below how to avoid significant VOC oxidation by NO₃ and achieve OH-dominated VOC oxidation in OFRs with high NO injection.

498 If NO_{3exp}/OH_{exp} > 0.1, NO₃ can be a competitive reactant for biogenic alkenes and dihydrofurans, 499 which have a C=C bond for NO₃ addition, and phenols, which have activated hydroxyl for fast hydrogen abstraction by NO₃ (Atkinson and Arey, 2003), while for lower NO_{3exp}/OH_{exp}, OH is expected to dominate 500 501 the oxidation of all VOCs, as shown in Fig. 6. Oxidation for VOCs without alkene C=C bonds and phenol 502 hydroxyl (such as alkanes and (alkyl)benzenes) is dominated by OH unless NO_{3exp}/OH_{exp} > 1000. Despite 503 its double bond, ethene reacts as slowly with NO3 as alkanes, likely due to lack of alkyl groups enriching electron density on the C=C bond, which slows NO3 addition. We calculate NO3exp/OHexp for OFR185-504 505 iNO and OFR254-iNO and plot histograms of this ratio in Fig. 6. Many experimental conditions lead to 506 high enough NO_{3exp}/OH_{exp} that NO₃ is a competitive sink for alkenes, while only under very extreme 507 conditions can NO₃ be a competitive sink for species without C=C bonds. High-NO conditions in OFR185-508 iNO have lower NO_{3exp}/OH_{exp} (~10⁻²-10²) than in OFR254-iNO (~10¹-10⁵) (Figs. 6 and S3d,g,j). This 509 difference in NO_{3exp}/OH_{exp} is due to the different levels of O_3 in the two modes, as high O_3 promotes 510 NO₂-to-NO₃ oxidation. Note that low-NO conditions in both OFR185-iNO and OFR254-iNO can also reach high NO_{3exp}/OH_{exp} as some high-NO conditions have. This is because in OFR185-iNO a large part of NO₃ 511 is formed by OH oxidation, resulting in NO_{3exp}/OH_{exp} being largely influenced by NOⁱⁿ but not by other 512 513 factors mainly governing OH (Fig. S3d); and under low-NO conditions in OFR254-iNO, NO3 can form 514 rapidly from NO₂+O₃, while OH can be heavily suppressed by high OHR_{ext} (Fig. S3g,j).

515 Most of the species shown in Fig. 6 are primary VOCs, except phenols and a dihydrofuran, which 516 can be intermediates of the atmospheric oxidation of (alkyl)benzenes (Atkinson and Arey, 2003) and 517 long-chain alkanes (Aimanant and Ziemann, 2013; Strollo and Ziemann, 2013; Ranney and Ziemann, 2016), respectively. Nevertheless, only the phenol production may occur in high-NO OFRs, as the 518 519 particle-phase reaction in the photochemical formation of dihydrofurans from alkanes is too slow 520 compared to typical OFR residence times (Ranney and Ziemann, 2016). Therefore, the impact of NO₃ 521 oxidation on VOC fate needs to be considered only if the OFR input flow contains high NO mixed with biogenics and/or aromatics [(alkyl)benzenes and/or phenols]. However, (alkyl)benzenes were likely to 522 523 be major SOA precursors in, to our knowledge, the only few literature OFR studies with high NO levels (Ortega et al., 2013; Tkacik et al., 2014; Liu et al., 2015). In the study of the air in a traffic tunnel (OFR185-524 525 iNO mode; Tkacik et al., 2014), where toluene is usually a major anthropogenic SOA precursor as in other urban environments (Dzepina et al., 2009; Borbon et al., 2013; Hayes et al., 2015; Jathar et al., 526 527 2015), NO_x was several hundreds of ppb. This resulted in an estimated NO_{3exp}/OH_{exp} range of ~0.1–1, where up to ~30% of cresols (intermediates of toluene oxidation) may have been consumed by NO₃. 528 Dihydrofurans may also have formed in the tunnel air (but outside the OFR) in the presence of $NO_{\rm x}$ 529 530 (Aimanant and Ziemann, 2013; Strollo and Ziemann, 2013) and, after entering the OFR, they would have 531 been substantially (up to ~50%) consumed by NO₃. In the laboratory experiment of Liu et al. (2015) with 532 toluene, the injection of as much as 10 ppm NO elevated NO_{3exp}/OH_{exp} to ~100, where cresols from 533 toluene oxidation reacted almost exclusively with NO₃ in addition to being photolyzed.

534 3.4.3 A case study

535 We use a case study of an OFR254-13-iNO laboratory experiment with a large amount of toluene 536 (5 ppm) and NOⁱⁿ (10 ppm) to illustrate how very high VOC and NO concentrations cause multiple types 537 of atmospherically irrelevant reactions in OFR. Due to very high OHR_{ext} and NOⁱⁿ, photolysis of toluene 538 at 254 nm may have been important (Peng et al., 2016). In case of a high (close to 1) quantum yield, up 539 to ~80% of the consumed toluene in their experiments could have been photolyzed (Scheme 1). Of the 540 rest of reacted toluene, ~10% undergoes H-abstraction by OH from the methyl group in the model, 541 leading to an RO2 similar to alkyl RO2 and likely proceeding with normal RO2 chemistry. ~90% of the toluene formed an OH-adduct (Calvert et al., 2002). As discussed above, 70% of this adduct (depending 542 543 on NOⁱⁿ) is predicted to recombine with NO₂ producing nitroaromatics because of the ppm-level NO_x.

544 The adduct could also react with O2 via two types of pathways, of which one was addition forming a 545 special category of RO2 (OH-toluene-O2 adducts) potentially undergoing ring-opening (Atkinson and 546 Arey, 2003; Orlando and Tyndall, 2012; Ziemann and Atkinson, 2012), the other H-elimination by O2 547 producing cresols. Again, like toluene, cresols may have been substantially photolyzed. As a result of 548 NO_{3exp}/OH_{exp} ~100, only a minor portion of cresol could have undergone OH addition and then H-549 elimination again. This pathway leads to the formation of methyldihydroxybenzenes and other OHoxidation products (Atkinson and Arey, 2003). The rest of cresols may have formed methylphenoxy 550 551 radicals, nevertheless, dominantly via H-abstraction by NO₃, since H-abstraction by OH was even a minor 552 pathway compared to the OH-addition one (Atkinson et al., 1992). In summary, the model results 553 suggest that there were two possible routes leading to nitroaromatic formation. However, one of them 554 (recombination of OH-aromatic adducts with NO₂) is likely of little atmospheric relevance due to very 555 high NO_x needed, and the other (H-abstraction from cresol) occurs in the atmosphere but is not a major 556 fate of aromatics (Calvert et al., 2002).

557 3.5 Implications for OFR experiments with combustion emissions as input

558 Emissions from combustion sources, e.g., vehicles and biomass burning, usually contain VOCs 559 and NO_x at very high concentrations (Table 1). An injection of this type of emissions (typically with OHR_{ext} of thousands of s⁻¹ or larger and NOⁱⁿ of tens of ppm or larger) in OFRs without any pretreatment is likely 560 to cause all experimental issues discussed in Peng et al. (2016) and this paper, i.e., strong OH 561 562 suppression, substantial non-tropospheric photolysis, strong RO₂ suppression by NO₂ whether RO₂ is 563 acyl RO2 or not, fast reactions of NO2 with OH-aromatic hydrocarbon adducts, substantial NO3 contribution to VOC fate, and even a near-total inhibition of OFR chemistry due to complete titration of 564 565 O3 by NO in the case of OFR254. We take the study of Karjalainen et al. (2016), who used an OFR to 566 oxidize diluted car exhaust in real-time, as an case study to investigate the extent to which these issues 567 may affect typical combustion source studies and to explore approaches to mitigate the problems.

568 During the first 200 s of their experiment (defined as the "cold start" period when the catalyst is 569 cold and emissions are high), NO and total hydrocarbon in the emissions of the test vehicle reached 570 ~400 and ~600 ppm, respectively. We first simulate the oxidation of those emissions without any 571 dilution (even though x12 dilution was used in their experiments) to explore the most extreme 572 conditions. Our model simulation indicates that such an extremely concentrated source would generally 573 lead to bad high- or low-NO conditions (depending on NO concentration) in their OFR (Fig. 7), even 574 though it was run at relatively high H₂O and UV. OH suppression can be as high as 3 orders of magnitude; 575 VOC fates by non-tropospheric photolysis and reactions of alkenes and phenols with NO3 can be nearly 100%; up to ~1/3 of OH-toluene adduct may be recombined with NO2 instead of forming an adduct with 576 577 O2. After the test vehicle entered the "hot stabilized" stage (200-1000 s), its VOC emissions (on the order of ppm) were still too high for an undiluted OFR to yield a good condition (Fig. S9). OH suppression 578 579 can still reach 2 orders of magnitude; non-tropospheric photolysis, and sometimes reactions with NO₃, 580 can still dominate over reactions with OH in VOC fates; reactions of OH-toluene adduct with NO2 can 581 still be substantial at some small NO emission spikes. Moreover, although NO emissions were roughly

at ppm level even during the hot stabilized period, NO effective lifetime may be very short during thatperiod, leading to low-NO conditions in their OFR.

584 As suggested in Peng et al. (2016) for low-NO OFR, dilution of sources can also mitigate strong 585 deviations on OFR-iNO chemistry vs. atmospherically-relevant conditions. A dilution by a factor of 12, 586 as actually used by Karjalainen et al. (2016), appears to be sufficient to bring most of the hot stabilized 587 period under good conditions (Fig. S9). However, most VOC, or in other words, most SOA formation potential, was emitted during the cold start period, when risky and bad conditions still prevailed (Figs. 588 589 7 and 8). Even if the emissions are diluted by x100, the cold-start emission peak (Fig. 7) is still under 590 risky conditions. Although bad conditions are eliminated and good condition is present during most of 591 time, this emission peak under risky condition may contribute >50% to total SOA formation potential 592 (Fig. 8). For SOA formed under good condition to be dominant, a dilution factor >400 would be needed. 593 Note that a strong dilution lowers aerosol mass loading in vehicle emissions. As a result, condensation 594 of gases onto particles is slower than in raw exhausts. However, condensational sinks after dilution may 595 still be significantly higher than typical ambient values (Matti Maricq, 2007; Donahue et al., 2016).

596Note that the emissions of the test vehicle of Karjalainen et al. (2016) are rather clean compared597to the typical 2013 US on-road fleet (i.e., all at the hot stabilized stage) measured by Bishop and598Stedman (2013) (Figs. 9 and S10). For emissions of an average on-road fleet, a dilution by a factor of599100 or larger would be necessary to ensure that most emissions would be processed in OFR185 under600good conditions at the highest H2O and UV in this study (Figs. 9b and S10b,e,h). In the case of lower H2O601and/or UV, an even larger dilution factor would be required.

Conducting OFR185-iNO experiments at high UV lowers the dilution factor needed for good 602 603 conditions. However, it also renders good high-NO condition impossible (see Section 3.2 and Fig. S4). If 604 one wants to oxidize vehicle exhausts in a high-NO environment in OFR, as in an urban atmosphere, OFR185 at low UV is necessary. Consequently, a much stronger dilution is in turn necessary to keep the 605 606 operation condition still good. Nevertheless, not all vehicle emissions can be moved into good high-NO 607 region through a simple dilution (Figs. 9c and S10c, f,i). Furthermore, a low UV would seriously limit the 608 highest OH_{exp} that OFR can achieve (~3x10¹¹ molecules cm⁻³ s for modeled good high-NO conditions in 609 this study), while a much higher OH_{exp} would be desirable to fully convert SOA formation potential into 610 measurable SOA mass. If both good high-NO condition and high OH_{exp} are required, new techniques 611 (e.g., injection of N₂O at percent level proposed by Lambe et al. (2017)) may be necessary.

612 4 Conclusions

In this study, OFR chemistry involving NO_y species was systematically investigated over a wide range of conditions. NO initially injected into the OFR was found to be rapidly oxidized under most conditions. In particular, due to high O₃ concentrations, NO lifetime in OFR254-iNO was too short to result in a significant RO₂ consumption by NO compared to that by HO₂ under all conditions with active chemistry. Nevertheless, it is not completely impossible for OFR185-iNO to have a significant RO₂ fate by NO and minor non-tropospheric photolysis at the same time ("good high-NO conditions"). According to our simulations, these conditions are most likely present at high H₂O, low UV, low OHR_{extr} and NOⁱⁿ

620 of tens to hundreds of ppb.

621 However, many past OFR studies with high NO injection were conducted under conditions 622 remarkably different from the abovementioned very narrow range. NOⁱⁿ and/or OHR_{ext} in those studies 623 were often much higher than good high-NO conditions require (particularly, >3 orders of magnitude in 624 some OFR studies using combustion emissions as input). In addition to non-tropospheric organic 625 photolysis, OFR oxidation of highly concentrated sources can cause multiple large deviations from 626 tropospheric OH oxidation, i.e., RO2 suppression by high NO2, substantial nitroaromatic formation from 627 the recombination of NO2 and OH-aromatic adducts, and fast reactions of VOCs with NO3 compared to 628 those with OH.

629 Working at lower NO_x (sub-ppm level) and VOC concentrations or dilution can mitigate these 630 experimental problems. In general, a strong dilution (by a factor of >100) is needed for OFR that process 631 typical on-road vehicle emissions. Humidification can also make good conditions more likely. By these 632 measures, good conditions can be guaranteed, as long as NO and/or precursor concentrations are 633 sufficiently low, while high-NO conditions cannot be ensured. To aid design and interpretation of OFR 634 experiments with high NO injection, we provide our detailed modeling results in a visualized form (Fig. S3). For OFR users in need for both high OH_{exp} and high NO, simple NO injection is not a good option. 635 636 New techniques (e.g., injection of N₂O proposed by Lambe et al. (2017) or other innovations) may be 637 necessary to meet this need. 638

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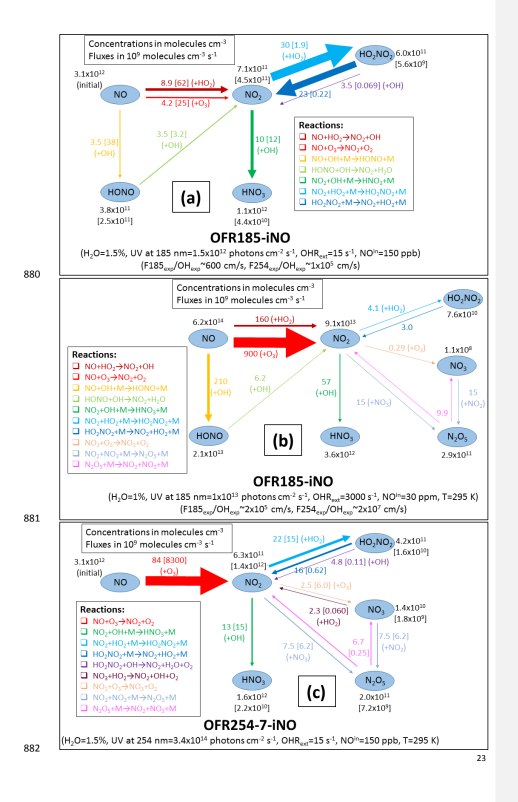
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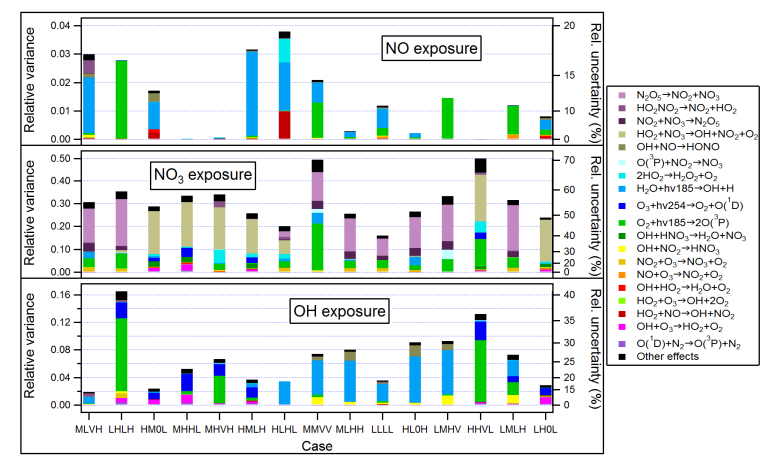
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883 Figure 1. Schematics of main N-containing species and their major interconversion pathways under 884 typical input conditions for (a) OFR185-iNO with NOⁱⁿ=150 ppb, (b) OFR254-7-iNO with NOⁱⁿ=150 ppb, 885 and (c) OFR185-iNO with NOⁱⁿ=30 ppm. Species average concentrations (in molecules cm⁻³) are shown 886 in black beside species names. Arrows denote directions of the conversions. Average reaction fluxes (in 887 units of 10^9 molecules cm⁻³ s⁻¹) are calculated according to the production rate, and shown on or beside 888 the corresponding arrows and in the same color. Within each schematic, the thickness of the arrows is 889 a measure of their corresponding species flux. Multiple arrows in the same color and pointing to the 890 same species should be counted only once for reaction flux on a species. Note that all values in these 891 schematics are average ones over the residence time, except for those in square brackets in panels a 892 and b, which are average values within approximate NO effective lifetime (τ_{NO} , or more accurately, an 893 integer multiple of the model's output time step closest to NO effective lifetime). All concentrations and 894 fluxes have two significant digits.



- 896 Figure 2. Relative variances (left axes)/uncertainties (right axes) of several outputs (i.e., NO, NO₃, and OH exposures) of Monte Carlo uncertainty propagation, and relative
- 897 contributions of key reactions to these relative variances in several typical cases (denoted in 4-character labels, see Table 2 for the typical case label code) in OFR185-iNO.
- 898 Relative variances are shown in linear scales (left axis), while corresponding relative uncertainties, equal to relative variances' square roots, are indicated by the non-linear
- right axis. Only the reactions with a contribution of no less than 0.04 to at least one relative variance are shown.

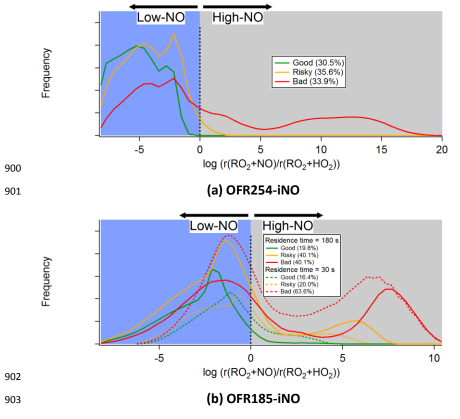


Figure 3. Frequency occurrence distributions of good, risky, and bad conditions (see Table 3) over
 logarithm of the ratio between RO₂ reacted with NO and with HO₂ (see Section S1 for more detail) for
 (a) OFR254-iNO (only the case with a residence time of 180 s) and (b) OFR185-iNO (including two cases
 with residence times of 180 and 30 s). Low and high-NO regions (see Table 3) are colored in light blue
 and grey, respectively.

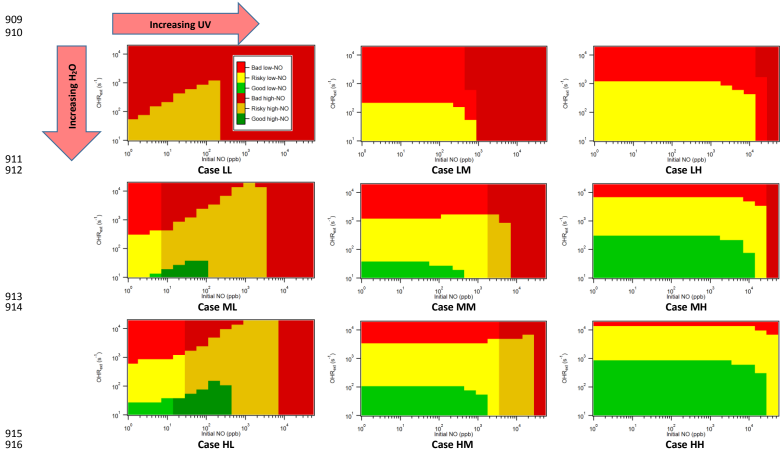




Figure 4. Image plots of the condition types defined in Table 3 vs. external OH reactivity (excluding N-containing species) and initial NO for several typical cases in OFR185-iNO (see Table 2 for the case label code).

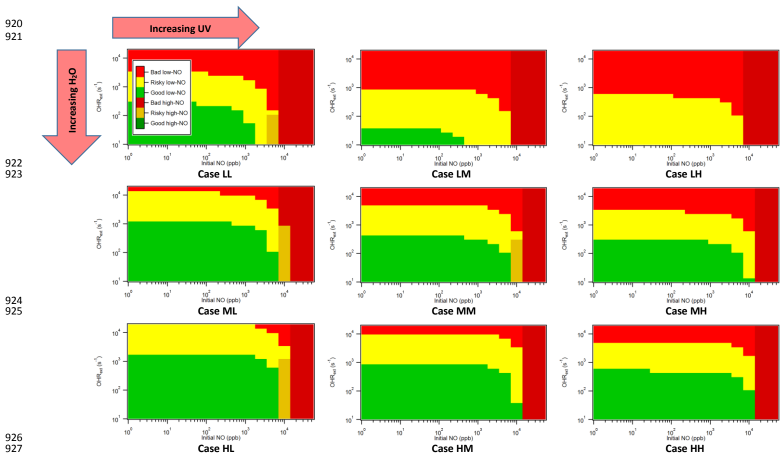


Figure 5. Same format as Fig. 4, but for OFR254-22-iNO.

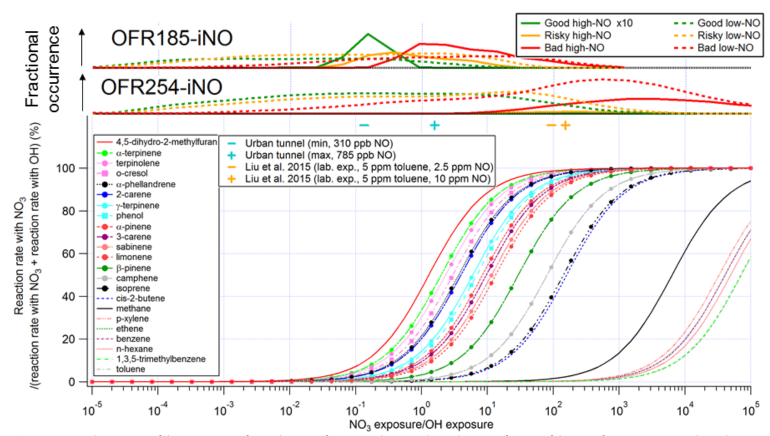
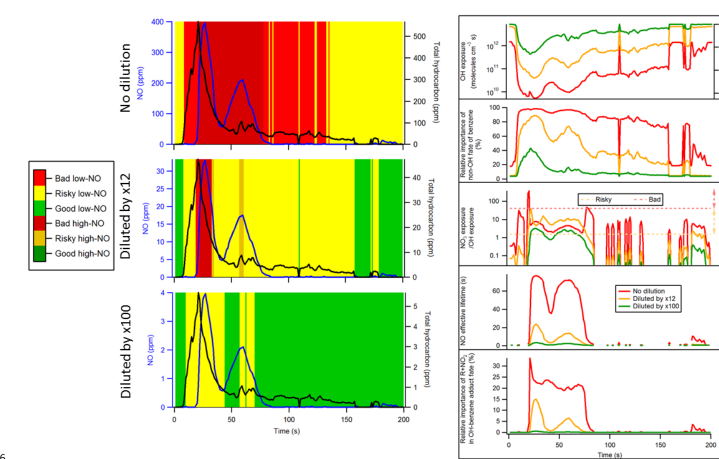


Figure 6. Fractional importance of the reaction rate of several species of interest with NO₃ vs. that with OH, as a function of the ratio of exposure to NO₃ and OH. The curves of biogenics and phenols are highlighted by solid dots and squares, respectively. The turquoise and orange markers show the ranges of modeled exposure ratios between NO₃

932 and OH of a source study in an urban tunnel (Tkacik et al., 2014) and a laboratory study (Liu et al., 2015) using OFR, respectively. In the upper part of the figure, the modeled

- 933 frequency distributions of ratios of NO₃ exposure to OH exposure under good/risky/bad high/low-NO conditions for OFR185-iNO and OFR254-iNO are also shown. See Table
- 934 3 for the definitions of the three types of conditions. All curves, markers, and histograms in this figure share the same abscissa.





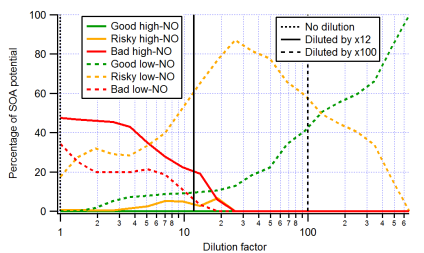
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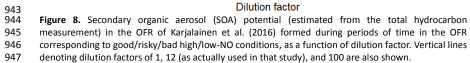
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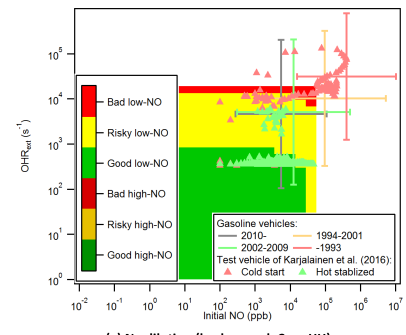
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e of remaining suppression

Figure 7. (left) NO and total hydrocarbon during the first 200 s of the test of Karjalainen et al. (2016) in the cases of no dilution, dilution by a factor of 12 (as actually done in that study), and dilution by a factor of 100. Different periods of time are colored according to corresponding emissions (i.e., input conditions for OFR), classified as good/risky/bad high/low-NO. (right) OH exposure/percentage of remaining OH after suppression, relative importance of non-OH fate of benzene, exposure ratio of NO₃ to OH, NO effective lifetime, and relative importance of reaction of OH-toluene adduct with NO₂ in the fate of this adduct in the OFR of Karjalainen et al. (2016) during the first 200 s of their test in the cases of no dilution, dilution by a factor of 12, and dilution by a factor of 100. Horizontal orange and red dashed lines in the middle right panel denote "risky" and "bad" regions for exposure ratio of NO₃ to OH, respectively. Above the orange (red) dashed line, reaction with NO₃ contributes >20% to the fate of phenol (isoprene).

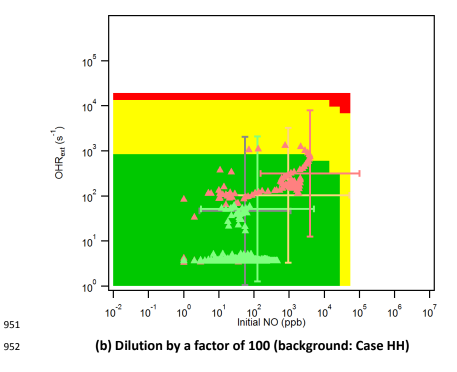




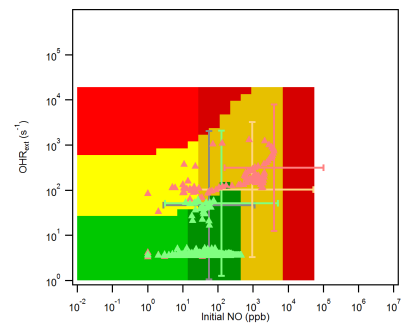




(a) No dilution (background: Case HH)







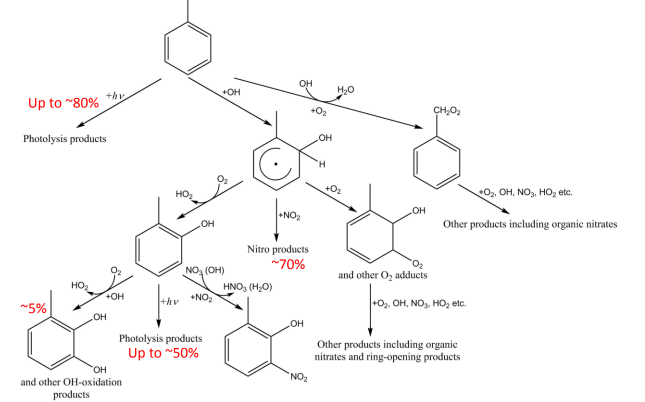


(c) Dilution by a factor of 100 (background: Case HL)

955 Figure 9. Location of individual 1 s datapoints vs. OFR185-iNO reaction conditions. Datapoints are shown 956 from the test vehicle of Karjalainen et al. (2016), as well as average exhaust from gasoline vehicle on-957 road emissions measured by Bishop and Stedman (2013). On-road emissions are classified by vehicle 958 year and the distribution of each category is shown as a cross representing 1 standard deviation (with 959 log-normal distribution assumed). The X and Y axes are NO and external OH reactivity (excluding N-960 containing species) due to vehicle emissions in OFR in the cases of (a) no dilution and (b,c) dilution by a 961 factor of 100. The Karjalainen et al. (2016) points are classified as cold start (during first 200 s) and hot 962 stabilized (during 200–1000 s). In addition, the same image plots as the panels of Cases HH (high H_2O 963 and high UV, see Table 2 for the case label code) and HL in Fig. 4 (OFR185-iNO) are shown as background 964 for comparison.

966 Scheme 1. Possible major reactions in an OFR254-13-iNO with 5 ppm toluene and 10 ppm initial NO. Branching ratios in red are estimated by the model and/or according to

967 Calvert et al. (2002), Atkinson and Arey (2003), Ziemann and Atkinson (2012), and Peng et al. (2016). Note that addition/substitution on the aromatic ring may occur at other
 968 positions. Intermediates/products shown here are the isomers that are most likely to form. Branching ratios shown in red are not overall but from immediate reactant.



Study	Source type	Temperature	Relative	Dilution factor	External OH reactivity of	Source NO _x
		(K)	humidity (%)		undiluted source (s ⁻¹)	concentration (ppm)
Link et al. (2016)	Diesel vehicle emission		50	45-110	~5000*1	436 ^{*1}
Martinsson et al. (2015)	Biomass burning emission			1700	156400 ^{*1}	154
Karjalainen et al. (2016)	Gasoline vehicle emission	295	60	12	~73000 ^{*2,a}	~400 ^{*1,b}
Liu et al. (2015)	Purified gas	293	13	1	~1400 ^{*1,a}	10 ^{*1,b}
Tkacik et al. (2014)	Tunnel air	293	42	1	~60 ^{*1,a}	~0.8*1
Ortega et al. (2013)	Biomass burning emission	290	30	~500	~15-500	~0.2

970
 Table 1. Experimental conditions of several OFR studies with high NO injection.

971 *1 maximum value in the study

 \ast_2 value at the moment of maximum NO emission

^{*a} NO_y species excluded ^{*b} NO only

Table 2. Code of the labels of typical cases. A case label can be composed of four characters denoting the water mixing ratio, the photon flux, the external OH reactivity 976

excluding N-containing species, and the initial NO mixing ratio, respectively. A case label can also be composed of two characters denoting the water mixing ratio and the 977

978 photon flux.

	Water mixing ratio	Photon flux	External OH reactivity (no ON)	Initial NO mixing ratio		
 Options	L=low (0.07%)	L=low (10 ¹¹ photons cm ⁻² s ⁻¹ at 185 nm; 4.2x10 ¹³ photons cm ⁻² s ⁻¹ at 254 nm)	0	0		
	M=medium	M=medium (10 ¹³ photons cm ⁻² s ⁻¹ at 185 nm;	L=low	L=low		
	(1%)	1.4x10 ¹⁵ photons cm ⁻² s ⁻¹ at 254 nm)	(10 s ⁻¹)	(10 ppb)		
	H=high	H=high (10 ¹⁴ photons cm ⁻² s ⁻¹ at 185 nm;	H=high	H=high		
	(2.3%)	8.5x10 ¹⁵ photons cm ⁻² s ⁻¹ at 254 nm)	(100 s ⁻¹)	(316 ppb)		
			V=very high (1000 s ⁻¹)	V=very high (10 ppm)		
Example —	LH0V:	low water mixing ratio, high photon flux, no external OH reactivity (excluding ON), very high initial NO mixing ratio				
	ML:	medium water mixing ratio, low photon flux				

979 980

Condition	Good	Risky	Bad	
Criterion	F185 _{exp} /OH _{exp} <3x10 ³ cm s ⁻¹ and F254 _{exp} /OH _{exp} <4x10 ⁵ cm s ⁻¹	$F185_{exp}/OH_{exp}<1x10^5$ cm s ⁻¹ and $F254_{exp}/OH_{exp}<1x10^7$ cm s ⁻¹ (excluding good conditions)	F185 _{exp} /OH _{exp} ≥1x10 ⁵ cm s ⁻¹ or F254 _{exp} /OH _{exp} ≥1x10 ⁷ cm s ⁻¹	
Condition	High-NO	High-NO		
Criterion* $\frac{r(RO_2+NO)}{r(RO_2+HO_2)} > 1$		$\frac{r(RO_2+NO)}{r(RO_2+HO_2)} \le 1$		

* See Section S1 for detail.

984 **Table 4.** Statistics of the ratio between OH exposures calculated in the model with the Lambe et al. (2011) residence time distribution (OH_{exp,RTD}) and in the plug-flow model

985 (OH_{exp,PF}). The geometric mean, uncertainty factor (geometric standard deviation), and percentage of outlier cases (>3 or <1/3) are shown for OFR185-iNO, OFR254-70-iNO,

986 and OFR254-7-iNO.

	Geometric mean	Uncertainty factor	Outlier cases (%)
OFR185-iNO	1.91	1.64	11
OFR254-7-iNO	1.59	1.51	7
OFR254-70-iNO	1.48	1.29	3