Yields and molecular composition of gas phase and secondary organic aerosol from the photooxidation of the volatile consumer product benzyl alcohol: formation of highly oxygenated and hydroxy nitroaromatic compounds

- 5
- 6 Mohammed Jaoui¹, Kenneth S. Docherty², Michael Lewandowski¹, Tadeusz E. Kleindienst¹
- ⁷ ¹Center for Environmental Measurement & Modeling, U.S. Environmental Protection Agency, Research Triangle Park,
- 8 NC, 27711, USA
- ⁹ ²Jacobs Technology, Inc., Research Triangle Park, NC, 27709, USA
- 10 Correspondence: Mohammed Jaoui (Jaoui.mohammed@epa.gov)
- 11



21	Abstract. Recently, volatile chemical products (VCPs) have been increasingly recognized as important precursors for
22	secondary organic aerosol (SOA) and ozone in urban areas. However, their atmospheric chemistry, physical
23	transformation, and their impact on climate, environment, and human health remain poorly understood. Here, the yields
24	and chemical composition at the molecular level of gas and particle phase products originating from the photooxidation
25	of one of these VCPs, benzyl alcohol (BnOH), are reported. The SOA was generated in the presence of seed aerosol from
26	nebulized ammonium sulfate solution in a 14.5 m ³ smog chamber operated in flow mode. More than 50 organic
27	compounds containing nitrogen and/or up to seven oxygen atoms were identified by mass spectrometry. While a detailed
28	non-targeted analysis has been made, our primary focus has been to examine highly oxygenated and nitro-aromatic
29	compounds. The major components include ring-opening products with high oxygen to carbon ratio (e.g., malic acid,
30	tartaric acids, arabic acid, trihydroxy-oxo-pentanoic acids, and pentaric acid), and ring-retaining products (e.g.,
31	benzaldehyde, benzoic acid, catechol, 3-nitrobenzyl alcohol, 4-nitrocatechol, 2-hydroxy-5-nitrobenzyl alcohol, 2-
32	nitrophloroglucinol, 3,4-dihydroxy-5-nitrobenzyl alcohol). The presence of some of these products in the gas and particle
33	phases simultaneously provides evidence of their gas/particle partitioning. These oxygenated oxidation products made
34	dominant contributions to the SOA particle composition in both low and high NOx systems. Yields, organic mass to
35	organic carbon ratio, and proposed reaction schemes for selected compounds are provided. The aerosol yield was 5.2%
36	for BnOH/H ₂ O ₂ at SOA concentration of 52.9 μ g m ⁻³ and ranged between 1.7-8.1 % for BnOH/NOx at SOA concentration
37	of 40.0-119.5 μg m ⁻³ .
38	
39	
40	
41	
42	
43	
44	
45	
46	
47	Key words: Benzyl alcohol, highly oxygenated compounds, Consumer products, VCPs, Silylation, Yield, Nitroaromatic
48	compounds, SOA

49 1 Introduction

50 Modeling atmospheric organic aerosol (OA) using chemical transport models (CTMs) is complex, challenging, and 51 often can lead to model-measurement discrepancies (Zhao et al., 2016). Applying CTMs to urban areas reveals that traditional VOCs including combustion-related processes cannot account for the observed OA mass, leaving a substantial 52 fraction unresolved (Hayes et al., 2015). Recent studies suggest that this discrepancy is due in part to unaccounted, rapidly 53 54 reacting SOA and ozone precursors from unknown sources (Hodzic et al, 2009; Hayes et al., 2015; McDonald et al., 2018; 55 Akherati et al. 2019; Lu et al., 2020). Volatile chemical products (VCPs), such as personal care products, cleaning agents, 56 coatings, adhesives, and pesticides have emerged as possible sources in urban areas (McDonald et al., 2018). Their 57 emissions can be larger than those from usual sources, such as motor vehicles (Coggon et al., 2021). Laboratory, modeling, and field studies for VCPs have been conducted to assess their potential to affect ambient OA and ozone formation in 58 59 urban and suburban locations (McDonald et al., 2018; Khare et al., 2018; Stockwell et al., 2021; Seltzer et al., 2021; 60 Gkatzelis et al., 2021; Milani et al., 2021; Pennington et al., 2021; Coggon et al., 2021). The contribution of VCPs to 61 ambient OA is not fully understood and only limited modeling studies have been reported (Mohr et al., 2015; Vlachou et 62 al., 2018; Pennington et al., 2021; Oin et al., 2021; Seltzer et al., 2021). Additionally, few experimental and chamber 63 studies of VCPs have been conducted with limited characterization of aerosol products (Wu and Johnston, 2016, 2017; 64 Harrison and Well, 2012; Charan et al., 2020, 2021; Humes et al., 2022). For example, the analysis of SOA from the 65 oxidation of cyclic methyl siloxanes (Wu and Johnston, 2016, 2017; Fu et al. 2020; Alton and Browne, 2020; Charan et 66 al., 2021) and cyclic siloxanes (Janechek et al., 2019) has been conducted. Kinetic studies with limited products characterization have been reported for the oxidation of benzyl alcohol (BnOH) by hydroxyl radicals (Bernard et al., 67 68 2013; Wang, 2015; Harrison and Well, 2009, 2012). Recently, Humes et al. (2022) highlight the importance of oxygenated 69 aromatic VCPs emission to generate urban SOA and oxygenated products in both gas and aerosol phases. Therefore, 70 understanding the atmospheric chemistry of VCPs is important to assess their role in air quality and climate and to improve 71 SOA chemistry in CTMs thereby allowing for better estimates in health studies and source apportionment.

The challenges associated with evaluating VCP impacts on urban OA can be addressed by identifying atmospheric VCP concentrations and SOA markers linking those VCP to ambient particulate matter (PM). Benzyl alcohol (C_7H_8O) is an important ring containing VCP used as an organic intermediate and a solvent in a wide range of applications (Antonelli et al. 2002). BnOH is emitted also from flowers and flowering trees (Do et al., 1969; Horvat et al., 1990; Larsen and Poll, 1990; Humpf and Scheier, 1991; Boatright et al., 2004; Vallat and Dorn, 2005; Orlova et al.; 2006) and found in indoor 77 air (Weschler, 2011). Gas kinetic studies of loss rates and product distributions have been conducted using flow tubes and 78 environmental chambers. Bernard et al. (2013) examined the rate and mechanisms of the OH + BnOH reaction. Similarly, 79 Harrison and Wells (2009, 2012) investigated the rate constants for the BnOH reaction with ozone, OH and NO₃ radicals. 80 Carter et al. (2005) conducted chamber experiments to assess ozone and PM formation from BnOH and related compounds. Product studies from BnOH oxidation have focused mainly on gas phase (GP) products. Several carbonyl 81 82 products (benzaldehyde (BnAld), formaldehyde, glyoxal, butenedial, 4-oxopentanal, 3-hydroxy-2-propanaldehyde), and 83 benzyl nitrate, o-hydroxybenzyl alcohol, o-dihydroxy benzene were reported from the above studies. With respect to the 84 particle phase (PP), Charan et al. (2020) reported aerosol yields from BnOH oxidation together with a limited number of 85 SOA products. Finally, Wang (2015) conducted a theoretical study to elucidate the reaction mechanism of the oxidation 86 of BnOH with OH radicals.

87 In this study, we report a detailed non-targeted chemical analysis of GP and SOA products originated from the photooxidation of BnOH in the presence and absence of oxides of nitrogen (NOx), with the aim to better understand the 88 89 chemical composition at the molecular level. Gas chromatography-mass spectrometry (GC-MS) and high-performance 90 liquid chromatography were used for the identification of a range of organic compounds including oxygenated 91 nitroaromatics and related compounds bearing up to seven oxygen atoms. Nitroaromatics are pollutants of concern due to 92 their toxicity, light-absorption properties, and relatively long residence times in the environment. Highly oxygenated 93 compounds can partition into pre-existing particles or be involved in new particle formation. Also, in the present study, 94 SOA and secondary organic carbon (SOC) yields were measured with the results compared to published data. A chemical 95 mechanism is then proposed to represent and account for selected gas- and aerosol-phase products observed in this study. 96

97 **2** Experimental methods

All chemicals including N, O-*bis*(trimethylsilyl) trifluoroacetamide (BSTFA) derivatization reagent with 1% trimethylchlorosilane (TMCS) as catalyst and benzyl alcohol (99%), 2-methyl-4-nitrophenol, L-(+)-tartaric acid, D-(-)tartaric acid, and meso-tartaric acid were purchased from Aldrich Chemical Co. (Milwaukee, WI) at the highest purity (99.8%) available and were used without further purification. In addition to standards reported in our previous studies (Jaoui et al., 2004; 2018), 3-nitrobenzyl alcohol, benzoic acid, and 4-nitrocatechol were purchased from Tokyo Chemical Industry (OR, USA); while pentaric acid, 2,3-dihydroxy-4-methoxy-4-oxobutanoic acid, and arabic acid were obtained from Aurum Pharmatech, LLC (NJ, USA).

105 **2.1 Chamber description and operation**

106 All experiments were conducted in a 14.5 m³ fixed-volume chamber having TFE Teflon coated walls and maintained 107 at a positive pressure of 0.1 Torr. The chamber operation, procedures, and instrumentation have been described previously 108 (Kleindienst et al. 2006; 2009), and here just experiment-specific details are primarily included. A combination of 109 fluorescent bulbs having radiation from 300-400 nm was used to photolyze NO₂. In the absence of NO_x, the radiation 110 system was altered to include UV-313 sunlamps to adequately photolyze H₂O₂. The chamber was operated in steady-state 111 (SS or flow) mode to provide continuously stable effluent concentrations. Under these conditions, reactants and products 112 equilibrate with the chamber surfaces to minimize irreversible losses of gases and particles. The SS operation allows for 113 extended sampling periods to improve the accuracy and precision of the measurements (Shilling et al., 2008). Temperature, relative humidity, and UV light intensity were measured continuously with an uncertainty of 5%. Pre-114 115 experiment and post-experiment procedures (see section 2.5 below) were routinely carried out before and after each 116 experiment to minimize contamination in the chamber. The reactant generation system provided constant sources of zero 117 air, reactants, water vapor, and ammonium sulfate (AS) seed aerosol. The reactant flow of gases (e.g., NO_x) into the 118 chamber was regulated using mass flow controllers. BnOH was injected using a syringe pump, vaporized in a heated glass 119 bulb, and injected with zero air. For experiments in the absence of NOx, a 50% aqueous solution of H_2O_2 was vaporized 120 and injected using a second syringe pump, and photolyzed to produce OH radicals. Typical chamber AS concentrations 121 were approximately 1 μ g m⁻³. Each SS experiment went through an initial transient period of 18-24 h until the reactant 122 and product concentrations reached steady state.

123

124 **2.2 Gas-phase measurements**

125 A wide variety of instruments were used to measure the reactants and products. Nitric oxide (NO) and NOx were 126 measured with a TECO (Franklin, MA) oxides of nitrogen analyzer with an upstream nylon filter to remove nitric acid 127 produced from $OH + NO_2$. The NOx analyzer was calibrated with a NIST-traceable NO standard. Initial H_2O_2 128 concentrations were estimated by UV absorption using the ratio of the H₂O₂ to O₃ absorbances at 254 nm, as described 129 by Kleindienst et al. (2009). Experiments in the absence of NOx were conducted dry to avoid aqueous loss of H_2O_2 . BnOH 130 concentrations in the inlet and within the chamber were measured semi-continuously using an SRI Model 8610C compact 131 gas chromatograph with flame ionization detector (GC-FID; SRI Instruments, Torrance CA). The purity of the BnOH 132 was verified with GC-MS analysis.

133 Low molecular weight carbonyls and dicarbonyls were quantified by derivatization using 2,4-dinitrophenylhydrazine 134 (Smith et al., 1989). Samples were collected at 0.5 L min⁻¹ for 25 min and derivatized in a 4 mL solution of acidified 135 DNPH and then heated for 40 min at 70°C. Air samples were drawn for 20 min at a rate of 0.50 L min-1 through an impinger containing 5 mL of a DNPH solution in acetonitrile. The resulting solutions were analyzed by high-performance 136 137 liquid chromatography with a ultraviolet detector (HPLC/UV) (Smith et al., 1989). A 15-component hydrazone standard (comprising formaldehyde-, acctlehyde-, accolein, acetone-, propionaldehyde-, crotonaldehyde-, methacrolein-, 138 139 butyraldehyde-, 2-butanone-, BnAld-, glyoxal-, valeraldehyde-, m-tolualdehyde-, methylglyoxal-, and hexaldehyde; AccuStandard, Inc.) at a free carbonyl concentration of 30 µg mL⁻¹ for each component was used for calibration. Separate 140 141 dihydrazone standards of glyoxal-DNPH and methylglyoxal-DNPH were also formulated. Carbonyls were separated using a Hewlett-Packard (HP) 1100 HPLC system having an Agilent Zorbax ODS 4.6 x 250 mm, 5-µm column maintained 142 at 30°C eluted with binary acetonitrile-water gradient. A 10 µL injection volume was used for all standards and samples. 143 144 Carbonyls were quantified by UV absorption with a diode array detector set to 360 nm. Control and sample processing 145 were managed with HP ChemStation software. More highly oxidized gas-phase organic species were also collected with 146 a 60-cm, 4-channel XAD4-coated annular denuder for off-line analysis (Jaoui and Kamens, 2001). Once collected, the 147 denuders were extracted and analyzed according to the methodology described in section 2.4 below.

148

149 **2.3** Aerosol-phase: bulk parameter measurements

150 Organic carbon (OC) was measured using a semi-continuous elemental carbon-organic carbon (EC-OC) instrument 151 (Sunset Laboratories, Tigard, OR) (Offenberg et al., 2007). The pumping system draws chamber effluent through a quartz filter at a rate of 8 L min⁻¹ with carbon-strip denuder to remove gas-phase organics that might interfere with the 152 153 measurement. With a sample collection time of 0.5 h and an analysis time of 0.25 h, the duty cycle for the measurement 154 of OC was 0.75 h (Lewandowski et al., 2015). The aerosol volume, size distribution, and total number density were 155 measured using a scanning mobility particle sizer (SMPS), (Model 3071A, TSI, Inc., Shoreview, MN) and a condensation 156 particle counter (CPC) (Model 3010, TSI, Inc., Shoreview, MN). The SMPS operating conditions were as follows: sample flow 0.2 L min⁻¹; sheath flow 2 L min⁻¹; size scan from 19 to 982 nm. 157

158

159 2.4 Molecular characterization of GP and PP oxygenated organic products

160 A non-targeted chemical analysis was conducted focusing mainly on species bearing hydroxy and carboxylic groups 161 (Jaoui et al., 2004, 2013, 2018). For each experiment, six 47-mm glass fiber (GF) filters were taken for 24 h at a flow rate 162 of 16.7 L min⁻¹. A second set of samples used an in-line 60-cm XAD-4 coated annular denuder (followed by a GF filter) and analyzed for gas-phase organic products (Jaoui and Kamens, 2001). After collection, GF filters were extracted by 163 sonication with 5 mL methanol for 1 h, and denuders were extracted with 30 mL 1:1 dichloromethane/methanol mixture 164 165 (Jaoui and Kamens, 2001). Prior to extraction, denuders and GF filters were spiked with *cis*-ketopinic acid (KPA), trans-166 p-menth-6-ene-2,8-diol (PMD), and d₅₀-tetracosane (TCS) as internal/recovery standards (IS/RS). Denuder extraction solvents were rotary evaporated to ~1 mL and filtered using 0.45-µm PTFE syringe filters. A 2 µL portion of this extract 167 168 was analyzed by GC-MS (Jaoui and Kamens, 2001). The remaining denuder and filter extracts were evaporated to dryness 169 under a gentle stream of N₂ at room temperature using an N-Evap evaporation bath (Organomation Associates, Inc., 170 Berlin, MA), then derivatized with BSTFA (Jaoui et al., 2004). This technique provides a sensitive method for measuring 171 low levels of highly oxidized organic compounds, including semi- and intermediate-volatile compounds in the GP and 172 PP.

The GC-MS analysis was conducted on an Agilent GC (7890B) coupled with a quadrupole mass spectrometer (5977B). The injector, heated to 270 °C, was operated in splitless mode. Compounds were separated on a 60-m-long, 0.25mm-i.d. RTx-5MS column (Restek, Inc., Bellefonte, PA) with a 0.25-µm film thickness. The GC oven temperature was initiated at 84 °C, held for 1 min, then increased at 8 °C min⁻¹ to 200 °C, followed by a 2-min hold, then an increase at 10 °C min⁻¹ to 300 °C and a 15-min hold. The ion source, ion trap, and interface temperatures were 200, 200, and 300 °C, respectively. Mass spectra were collected in both the methane-chemical (CI) and electron ionization (EI) modes.

179

180 **2.5 Experimental and quality control procedures**

Before each experiment started, the chamber was flushed with zero air (hydrocarbon-free) for 24 h from an Aadco clean generator (Cleves, OH, USA). Experiments were conducted in either the absence or presence of NOx. For experiments with NOx, BnOH and NO were added to the chamber through flow controllers to the target concentration. For experiments in the absence of NOx, the photolysis of H_2O_2 was the source of OH. H_2O_2 as a 50 % aqueous solution was injected through a syringe pump into a heated glass bulb where it vaporized and then was mixed rapidly by the main dilution air flow. For these experiments, BnOH was added as described above. Ammonium sulfate seed aerosol was also introduced into the chamber for all experiments to serve as a condensing medium for semivolatile organic products that

188 might form. After the reactants reached steady state concentrations (24 hours), the background was characterized using all instruments to check for artifacts including background GP and PP species. Background chamber air was also 189 190 characterized using off-line analysis of denuder and/or filters as described above. Previous studies show that BnAld and 191 to a lesser extent benzoic acid, benzyl benzoate and dibenzyl ether present either as impurity, or as decomposition products 192 upon BnOH exposure to air at room temperature or sonication (Urakami et al., 2000; Ferri et al., 2006; Abend et al., 2004). 193 Here we investigated the effect of chamber air, sonication, and BSTFA derivatization on BnOH artifacts as described in 194 the supplementary information (SI) in section 1. A small amount of BnAld impurity was detected using the direct injection 195 (DI) method and estimated to be <0.1% in the purchased solution. When BnOH was exposed to clean air in the chamber 196 in the absence of light, sonication, and/or BSTFA derivatization, our results show additional low level of BnOH 197 conversion to BnAld and benzoic acid using DI and BSTFA methods, which is similar to the findings of Abend et al. 198 (2004), Urakami et al. (2000), and Ferri et al. (2006). Additional results and descriptions are provided in section S1 (SI). 199 Experiments were initiated by turning on the lights and allowing the irradiated chamber effluent to reach SS 200 conditions over a 24-h period which permits active sampling by the on-line instruments and the collection of denuder and 201 filter samples for subsequent off-line analysis. For organic intermediates wall losses are typically not an issue due to 202 reactions being conducted within a Teflon chamber. This potential issue is mitigated further from operating the chamber 203 in a SS mode where compound loss and re-evaporation quickly comes to steady state. Short lifetimes of radical 204 intermediates with other gas-phase constituents also render a negligible wall-loss. The stability of BnOH in the chamber 205 was investigated and BnOH was found to be highly stable with results given in the SI (section S1). Denuders and GF filter 206 samples were also analyzed to probe reproducibility of the analytical technique. The analysis showed consistent results. 207 Gas and particle samples from BnOH photooxidation are dominated by oxygenated species, several not having 208 authentic standards, and thus a portion of each sample was derivatized. Initially, we eliminated peaks detected in blank 209 and background samples. For compounds having standards, comparisons were made between the retention times and mass 210 spectra (CI and/or EI mode) of the chamber-derived peaks and those of the standards. For compounds not having 211 standards, individual peak identifications were associated with a product peak only if its retention time and mass spectrum 212 was consistent with the fragmentation pattern of the BSTFA-derivatized compound. All recorded spectra in this study were compared with those derived from reference standards, the literature, the NIST library, and an archive of mass 213 214 spectra from product compounds determined in our laboratory over the past twenty years.

216 3 Results and discussion

The initial conditions of the experiments conducted in this study are summarized in Table 1. Three NOx experiments were carried-out with initial BnOH ranging from 0.36 - 0.72 ppm and NO from 0.096 - 0.19 ppm. One experiment without NO_x was conducted with initial H₂O₂ and BnOH levels of 3.0 and 0.32 ppm, respectively. NOx experiments were conducted at ~30% RH, and the H₂O₂ experiment at < 4% RH to minimize H₂O₂ uptake onto chamber surfaces. Chamber temperatures were set to 25 °C. Each experiment was conducted for up to five days for samples requiring substantial masses or extended collection times and frequencies.

223 Steady state concentrations of NO, BnOH, O₃, and NOy for the four experiments are given in Table 2. The reacted 224 BnOH and NO were calculated from the difference between the initial and steady-state concentrations. For NOx experiments, the range of reacted BnOH concentrations was 0.22 - 0.34 ppm having a reproducibility of 20-30%. Under 225 these conditions, steady state concentrations of NOy, and O_3 were in the range of 0.08 - 0.16 and 0.011 - 0.15 ppm, 226 respectively. With NO present at steady-state, peroxy-peroxy (RO2-RO2) reactions were minimized. A constant aerosol 227 228 source was maintained for initial conditions given in Table 1. The major aerosol parameters measured (SOA, SOC, and 229 OM/OC) are given in Table 3. SOC uncertainties were taken from the reproducibility of the semi-continuous measurement 230 and typically better than 10% for a single run. For the organic mass (OM), the uncertainties are determined from the 231 reproducibility of side-by-side filter measurements which are typically better than 5%. An estimate of the systematic 232 errors due to minor changes in reactant concentrations, minor variations in chamber temperature, and similar factors bring 233 the total uncertainty to between 15-25% for these parameters (Kleindienst et al., 2009). SOA/SOC values were then 234 determined from the corrected data and given in Table 3. For experiments in the presence of NOx, SOA/SOC values 235 ranged from 1.7-2.0. Similarly, in the absence of NOx, the measured SOA/SOC value was 2.1.

236

237 **3.1 Secondary organic aerosol and secondary organic carbon yields**

Secondary organic aerosol yield (Y_{SOA}) and secondary organic carbon yield (Y_{SOC}) were calculated from the following respective relationships $Y_{SOA} = SOA/\Delta HC$ (1); $Y_{SOC} = SOC/\Delta HC_C$ (2) where SOA is the corrected organic aerosol mass concentration originated from filter measurements (6 filters) and ΔHC is the reacted BnOH concentration. SOC is the organic carbon concentration found in Table 3 and ΔHC_C is the reacted BnOH carbon concentration. SOA and SOC were corrected for wall loss to the chamber which had previously been determined for organic aerosol to be 0.067 h^{-1} (Kleindienst et al. 2012). Uncertainties in the yield come from the experimental uncertainties in SOA and SOC

244 production and the reacted BnOH concentrations. The uncertainty in the reacted BnOH results from the reproducibility 245 of the initial and steady-state values and is estimated to range from 20 - 30% given the low volatility of BnOH and challenges for introducing oxygenated species into the chamber in a consistent manner. Such challenges are also present 246 247 in a batch mode experiment given the difficulty to determine BnOH time profiles given its volatility and high reactivity toward oxidants (Shilling et al., 2008; Kroll et al., 2007). Similar findings have been reported for sesquiterpenes oxidation 248 249 (Jaoui et al., 2013). As a result, aerosol yields of higher accuracy are often reported to be associated with steady state as 250 opposed to batch mode experiments (Shilling et al., 2008). Moreover, in this work we explored the possibility of BnOH 251 being taken up by ammonium sulfate (AS) seed aerosol prior to start of the irradiation or by SOA after it is initiated. This 252 test was conducted using GC-MS analysis of derivatized (BSTFA) and underivatized denuder and GF filter extracts collected before and after the reaction starts (SI; Section S1). Under the experimental conditions used in this study, BnOH 253 254 was undetected in AS and SOA, thus limiting any participation in particle chemistry that may occur.

255 The production of aerosol, and thus the yield, were found to be highly sensitive to the precise initial conditions 256 (Tables 1, 3). Yields for the four experiments are shown in Table 3. Y_{SOA} values were determined for SOA concentrations from 39.6 - 119.5 μ g m⁻³ and ranged between 3.6 and 8.1%. Similarly, Y_{SOC} was measured for SOC concentrations from 257 $23.2 - 58.9 \ \mu gC \ m^{-3}$ and found to range between 2.7 and 5.1%. In the absence of NO_X, SOA and SOC yields were 5.2 258 and 3.1 % measured for SOA and SOC concentrations of 52.9 µg m⁻³ and 24.8 µgC m⁻³, respectively. For the two systems 259 at similar SOA concentrations ER890 and ER892, the SOA yield was higher for the experiment with NO_x. This may 260 261 result from the reaction of BnOH with NOx which tends to produce high levels of BnAld (Table 4), which may undergo 262 secondary reactions leading to additional SOA formation (see section 3.3). As expected, the data in Table 3 indicate that 263 Y_{SOA} and Y_{SOC} are lower at the lower SOA and SOC concentrations, respectively.

264 These SOA and SOC yields can be compared with other studies. Recently, Charan et al. (2020) reported SOA yields for the photooxidation of BnOH in the presence of NO_x with the initial OH coming from the photolysis of H_2O_2 . Their 265 chamber was operated in a batch mode and SOA yields approaching unity were reported. By contrast, three additional 266 267 studies reported much lower SOA yields of 9%, 30%, and 41% from McDonald et al. (2018), Carter et al. (2005), and Li et al. (2018), respectively. The yield reported by McDonald et al. (2018) was based on a multi-generation oxidation model; 268 that of Carter et al. (2005) was estimated as described in the original report, and that of Li et al. (2018) was based on 269 measurements in the presence of NOx and a surrogate urban hydrocarbon. The results of our study are much closer in 270 271 value to McDonald et al. (2018). The study by Charan et al. (2020) suggests that conditions can be found where BnOH

272 SOA yields are substantially greater than that found in this study and those previously reported. The major differences 273 between the Charan et al. study and the present work were the chamber-mode operation, the seed aerosol type and levels, 274 and the mix of oxidants used. While it can be difficult to compare the yields from the two studies some comments can be 275 made. (1) As noted, the Charan et al. yields result from conventional batch mode irradiations of BnOH, H_2O_2 and NO_X . (2) SOA levels were measured using an SMPS which measures aerosol volume which is then converted to aerosol mass 276 277 using a density of $1.4 \,\mu g \,nL^{-1}$. (3) Perhaps the biggest difference between the two studies is the use of an extremely high 278 initial seed aerosol mass, approximately two orders of magnitude higher than in this study. Thus, it is possible that an 279 adsorption mechanism played a part in contributing to the measured yields. (4) Finally, the use of high initial H_2O_2 280 concentrations relative to BnOH make it possible that H₂O₂ effectively competed with BnOH for OH via the reaction of $OH + H_2O_2 - -> HO_2 + H_2O$ thus generating a system rich in HO₂. Thus, aging process may be more prominent than in our 281 282 study. The uncertainties associated with SOA and reacted BnOH measurements and wall loss correction are unlikely to 283 account for the differences in the two studies. However, SOA yields have been reported to increase considerably as a 284 function of initial seed aerosol (Zhang et al. 2014), as well as to increase with the OH radical exposure (Wang et al., 285 2018). Qualitatively these two factors might bring the present SOA yields into reasonable agreement with Charan et al. 286 (2020), when data were extrapolated to similar seed aerosol and OH exposures. And as previously noted, in the low initial 287 seed aerosol used in our study no benzyl alcohol was detected in the seed aerosol or deposited on SOA. As further 288 plausibility for the results from the present study, Humes et al. (2022) recently reported yields from 12% - 18% for two 289 oxygenated aromatic species (1-phenoxy-2-propanol, and phenoxy-ethanol), compounds having similar structures to 290 BnOH.

291

292 **3.2 Reaction products identification**

Three methods were used in this study to identify oxygenated reaction products at the molecular level: (1) DNPH as derivatizing agent for small carbonyls (Smith et al. 1989); (2) BSTFA as derivatizing agent for hydroxyl and carboxylic compounds for GP and PP (Jaoui et al., 2004); and (3) direct injection (DI) method providing the capability for analysing slightly polar to non-polar compounds without the use of derivatization (Jaoui and Kamens, 2001). For the BSTFA and DI methods, the analysis of laboratory generated GP and PP products from BnOH oxidation shows a series of organic compounds containing nitro, ketone, carboxylic acid, and/or alcoholic functional groups. Many of these compounds do not have authentic standards and their identifications were based on the interpretation of the mass spectra of the derivatized

300 and/or underivatized compound (Jaoui and Kamens 2001; Jaoui et al., 2004, 2005). The identification should be regarded 301 as tentative except for compounds that have authentic standards. For the BSTFA method, the recognition of characteristic 302 ions was used to guide the analysis of mass spectra of the derivatives obtained in both electron ionization (EI) and chemical 303 ionization (CI) using methane as reagent gas. BSTFA reacts with -COOH and -OH groups to form BSTFA derivatives. 304 Characteristic ions are m/z 73, 75, 147, and 149. Adduct ions in CI from the derivatives include m/z M⁺ + 73, M⁺ + 41, M^{+} + 29, and M^{+} + 1; fragment ions include m/z M^{+} - 15, M^{+} - 73, M^{+} - 89, M^{+} - 117, M^{+} - 105, M^{+} - 133, and/or 305 306 M^+ – 207. The approach used for the identification is as follows: peaks detected in blank and background chamber samples 307 were eliminated first. A peak was associated with a reaction product only if its corresponding mass spectrum was 308 consistent with the fragmentation pattern of the BSTFA derivatization reagent. All recorded spectra were compared with 309 spectra derived from various reference compounds, authentic standard, NIST library, the PubChem website 310 (pubchem.ncbi.nlm.nih.gov), and/or by MS assignment. While the off-line technique is an integrated technique that 311 requires long sampling times, it does provide a sensitive method for products identification at the molecular level as well 312 as measuring low concentrations of highly oxidized organic compounds, and semivolatile compounds in the GP. Thus, 313 products found by this collection technique could be informative for possible precursors for the types of compounds that 314 may form in the PP. In the following discussion, data are first presented to support tentative identifications of oxidation 315 products in the GP and PP.

316 Gas phase products. GP measurements were made of major carbonyl products formed during the photooxidation of 317 BnOH including formaldehyde, acetaldehyde, acetone, methacrolein, 2-butanone, BnAld, glyoxal, and methylglyoxal. 318 Steady-state concentrations are given in Table 4. Under the conditions shown in Tables 1 and 2, high concentrations were observed for BnAld and glyoxal, and to a lesser extent formaldehyde in experiments with NOx, and high concentrations 319 320 of formaldehyde, acetaldehyde, and to a lesser extent BnAld and glyoxal in the experiment without NOx. BnAld level 321 was a factor of \sim 5 higher in NOx experiments compared to H₂O₂ experiments, and formaldehyde a factor of \sim 36 lower. 322 Glyoxal and methylglyoxal concentrations largely were similar in both NOx and H₂O₂ experiments. The formation of 323 BnAld, glyoxal, and formaldehyde as major products (Table 4) have already been reported from the oxidation of BnOH with yields of 25 (\pm 5), 20 (\pm 2), and 3.0% (\pm 0.2), respectively (Bernard et al., 2013; Harrison and Wells, 2012). 324

325 GP samples were also collected on five-channel annular denuders. Each denuder sample was extracted and analyzed 326 directly with GC-MS without derivatization. The remaining extract was silylated, and GC-MS analyzed qualitatively.

327 Typical total ion chromatograms (TIC) of GP products detected and identified in this study are shown in Figure 1. Figure 328 1 shows portions of three TIC in +EI of GP samples taken from experiments ER889 at steady state (underivatized: Figure 329 1a), ER892 (underivatized: Figure 1b), and ER889 (silvlated derivatives: Figure 1c). Peaks assigned in Figure 1 were 330 identified either by comparison with an authentic standard or by MS assignment. For clarity, only the main products are 331 shown, although several peaks could not be structurally identified. SOA generated from BnOH photooxidation is 332 dominated by oxygenated ring-opening products (see below). However, ring-retaining products were among the main 333 products observed in the GP including semivolatile organic compounds (SVOCs) (Figure 1c). Chromatograms associated 334 with the underivatized samples (Figure 1a, b) were used mainly to identify BnOH and BnAld in the system, although 335 several additional peaks absent in the background chromatogram were observed. At steady state, BnOH was not reacted completely as it was detected in both systems only in the GP using both DI and BSTFA methods (Figure 1). BnAld was 336 detected in both systems in the gas and particle phases, although it was not present with BSTFA method because of the 337 338 absence of OH or COOH groups. Figure S3 shows EI mass spectra of BnAld identified using authentic standard, and 339 those associated with three peaks eluting at 11.3, 12.0, and 12.8 min. Although no structural information could be 340 associated with these three peaks, molecular weights of 152, 152, 138 Da (all derivatized and underivatized masses are 341 Dalton (Da) but are not designated as such hereafter), were tentatively obtained.

342 Select GP products containing OH groups identified in the present study are summarized in Table 5. Table 5 contains 343 proposed structures, molecular weights of the silvlated derivatives (MW_{BSTFA}) and underivatized compounds (MW), 344 formula, and the 5 most intense ions associated with BSTFA derivatives in EI mode. Table 5 shows if GP products are detected also in the PP. Figure S4 shows EI mass spectra associated with selected peaks observed in Figure 1c, including 345 BnOH-1TMS, benzoic acid-1TMS, catechol-2TMS, and 2-hydroxybenzyl alcohol-2TMS. 2-Hydroxybenzyl alcohol-346 347 2TMS (2OHBnOH) peak eluted at 21.4 min was one of the largest peaks detected in the chromatogram in Figure 3c. The 348 20HBnOH-2TMS EI mass spectrum (Figure S4, bottom) shows strong characteristic fragments ions at m/z 73, 179 (M⁺⁻ 349 - 89), 253 (M⁺ - 15), 268 (M⁺), and its corresponding CI mass spectrum shows ions at m/z 253 (M⁺ - 15), 179 (M⁺ - 89) and weak adducts at M^{+} + 1, M^{+} + 29, and M^{+} + 41 that are consistent with the presence of two (-OH) groups, indicating 350 351 a BSTFA derivatized molecular weight of 268 Da. Bernard et al (2013) have identified 2OHBnOH and catechol in the GP of the reaction of BnOH and OH radicals. In our study, catechol was observed only in the H_2O_2 system in the PP. 352 353 Additional peaks were observed, which their mass spectra are consistent with products bearing OH and/or COOH groups, 354 however, their structural identification could not be obtained due to lack of authentic standards and the complexity of the

- interpretation of their mass spectra. BnAld was reported to undergo secondary reactions (Sankar et al., 2014) and may
- 356 play an important role as precursor for some oxygenated species observed in this study.

357 Particle phase products. One of the advantages of conducting experiments in SS mode is collecting sufficient gas and 358 aerosol masses on denuders and filters for qualitative and quantitative offline analysis. In this study, aerosol collected on 359 GF filters were solvent extracted, with the resulting extracts subjected to BSTFA derivatization followed by GC-MS analysis. SOA generated from both NOx and H₂O₂ systems was dominated by oxygenated organic compounds, for which 360 361 mass spectra for more than 50 species have been recorded. These species may have undergone several generations of 362 atmospheric oxidation. Several individual large peaks have been detected in addition to a significant number of small 363 peaks as shown in Figure 2. Figure 2 shows portions between 9 and 28 min of the TIC chromatograms of the silvlated 364 derivatives of the aerosol extracts associated with BnOH/NOx (top) and BnOH/H₂O₂ (bottom). The portion after 28 min 365 is discussed in the next section. The chromatograms in Figure 2 can be directly compared because the chamber air sampled 366 and the amount of extract analyzed for each system were the same. This evaluation revealed that more than 70% of peaks eluted from each system are identical, suggesting similar chemistry is involved in BnOH reaction products formed in the 367 368 presence and absence of NOx. In addition, a series of peaks dominated by fragments with odd m/z were observed only in 369 BnOH/NOx and their mass spectra were associated with nitrogen containing compounds as discussed in the NACs section 370 below. This suggests that the composition of a portion of SOA produced in the presence of NOx is different than that 371 formed in the absence of NOx, which can be clearly illustrated by the filters and extracts color shown in Figure 2 (bottom). 372 Consistent with the presence of nitroaromatics, filter F2 and methanol extract (E2) has lost most of the color seen in F1 373 and E1. The presence of NOx in the system produced material (filter F1) of a deep brown color. Most species structurally identified in this study have not been reported in the literature, and mass spectra associated with several peaks are provided 374 375 either in the main manuscript or in the SI. Additional reaction products (e.g., oligomers, organonitrates) might have been 376 present in the SOA but could not be detected based on the analytical techniques used in this study. Note that formulae, in 377 particular chemical structure, could not be obtained for several peaks recorded in this study due to challenges interpreting 378 their mass spectra. A set of compounds identified and detected before 28 min in the present study are summarized in 379 Table 5.

Ring retaining products (e.g., 2-hydroxy benzyl alcohol, benzoic acid, 4-hydroxy benzoic acid, and catechol) were detected in the PP in both systems. As noted above, some ring-retaining products were detected also in the GP as shown 382 in Table 5. Salicylaldehyde and 3-hydroxybenzaldehyde were present only in the GP. These two hydroxy-aldehydes may 383 undergo additional secondary reactions leading to some ring-opening products observed in this study. Representative EI 384 mass spectra of the TMS-derivatives associated with four compounds are shown in figure 3 including benzoic acid, 385 benzene-1,2-diol (catechol), 4-hydroxybenzoic acid, and 2-hydroxybenzyl alcohol. Additional EI and CI mass spectra are shown in figures S4 and S5 in the SI. The EI mass spectrum of the BSTFA derivative of 2-hydroxybenzyl alcohol 386 displayed in figure 3 shows abundant fragment ions at m/z 73, 147, 267 (M⁺), 253 (M⁺ - 15), and 179 (M⁺ - 89), and 387 388 weak ions at m/z 91, 223 and 163. The corresponding CI mass spectrum displayed in figure S4d shows abundant fragment 389 ions at m/2 268 (M⁺), 253 (M⁺ - 15), and 179 (M⁺ - 89) and adduct ions at m/2 293 (M⁺ + 29) and 309 (M⁺ + 41). This 390 fragmentation pattern is consistent with the presence of a compound with two hydroxyl groups and a benzene ring (m/z391 91) having molecular weight 268 for the BSTFA derivative, and MW 124 for its underivatized form. Similarly, the BSTFA 392 EI mass spectrum of 4-hydroxybenzoic acid (Figure 3c) shows characteristic fragment ions at m/z 73, 193 (M^{+,} - 89), 223 393 (M^{+,} - 60), 267 (M^{+,} - 15) and 282 (M^{+,}), and its CI mass spectrum fragment ions at m/z 73, 193, 67 and adducts at 283, 394 and 311. Again, these fragments and adducts are consistent with the presence of two (-OH) groups and a molecular weight of the derivatized compound of 282 and 138 for the underivatized compound. The presence of a peak at m/z 153 (M⁺ -395 396 117) is consistent with a compound bearing an organic acid group. The EI mass spectra recorded in this study for 2-397 hydroxybenzyl alcohol and 4-hydroxybenzoic acid are identical to the reference NIST spectrum (webook.nist.gov). Figure 398 S5 shows EI mass spectra associated with four peaks eluted at 12.86, 15.58, 16.24, and 19.78 min consistent with the 399 fragmentation pattern of BSTFA derivatives, although their structures could not be obtained.

400 Highly oxygenated compounds (HOCs). Recent studies show that highly oxygenated compounds (e.g., HOMs) play an important role in understanding SOA formation (Berndt et al. 2016, Jaoui et al., 2019; 2021 and references therein, Piletic 401 402 and Kleindienst, 2022). These compounds may result from several generations of atmospheric oxidation. In this study, 403 several ring-opening products eluted late in the chromatograms (RT > 25 min), with a relatively high O:C ratio of > 1.3404 likely contributes to their condensation in the PP, were detected. Three groups of these oxidation products were detected 405 in the PP in both systems. Figure 4 shows the portion between 25 and 34 min of selected GC-MS extracted-ion chromatograms where these groups (color coded) elute and uses the selected ions m/z 423, 437, and 525 (merged in one 406 407 chromatogram) to best illustrate them: (a) BnOH/NOx; (b) BnOH/H₂O₂; (c) chamber background. Groups 1, 2, and 3 consist of three (green), eight (blue) and four (red) peaks, respectively, and are completely absent from the background 408 409 chromatogram (figure 4c). Results from a comprehensive interpretation of EI and CI mass spectra associated with peaks

shown in figure 4 enabled the identification of several isomers associated with each group. Figure 5 displays three EI mass spectra associated with each group main peak, along with proposed structure and chemical formulae. Table 6 gives the major highly oxygenated compounds identified in this research, including the main peaks from each of these groups, in the order of their underivatized molecular weight. Table 6 gives the chemical formulas, O:C mass ratio, the five most abundant ions associated with each TMS derivative in methane-CI and EI modes, the molecular weights of the underivatized (MW) and TMS-derivatized compounds (MW_{BSTFA}), and the proposed chemical structures of the compounds.

417 Group 1 consists of meso-tartaric acid (mTA) (Rt 26.04 min), and L-/D-tartaric acids (ITA/dTA) (Rt 27.66 min) 418 identified based on authentic standards. The mass spectra of BSTFA derivatives of ITA and dTA standards (Figure S6, 419 SI) are very similar (eluting at the same time) and are only slightly different from the mTA (Figure S6: SI); however, 420 TA/dTA and mTA elute at two different retention times (Figure 4, S6). The peak associated with mTA, and TA/dTA are 421 among the largest peak observed in this portion of the chromatograms. Note, ITA isomer is the most abundant tartaric 422 acid present in nature (DeBolt et al., 2006). The fragments and adducts observed for the peak eluting at 25.19 min are 423 similar to those of mTA and d-/ITA and are consistent with the presence of four OH groups, a MW of 452 for the 424 derivatized compound and 164 for the underivatized compound, and a $C_4H_6O_6$ chemical formula. Tartaric acid has been 425 reported in ambient aerosol (Rohrl and Lammel, 2002; Gowda et al., 2016) and in chamber 1,3-butadiene SOA (Jaoui et 426 al., 2014). Recent studies suggest that tartaric acid and other hydroxy carboxylic acids undergo heterogeneous OH reaction 427 in aqueous solution, with the presence and position of OH group(s) playing an important role in fragmentation and 428 functionalization of organic aerosol (Cheng et al., 2016).

Group 2 consists of eight peaks (figure 4: blue) eluting between 28.5 and 31.5 min. The EI and CI mass spectra 429 430 associated with each peak display similar fragment and adduct ions across the range of 50 to 600 Da. The interpretation 431 of these mass spectra allows us to infer the molecular weight (MW) of the underivatized compounds as 164 and MW_{BSTFA} 432 of 452 for the TMS derivatives. The BSTFA CI mass spectrum of the peak eluted at 29.48 (largest peak) shows characteristic fragment ions at m/z 73, 437 [M^{+, -15]}, 363 [M^{+, -89]}, and 305 [M^{+, -105]}, and an adduct at 453 [M^{+, +1]}, 433 481 $[M^+ + 29]$, and 493 $[M^+ + 41]$. These fragments and adducts are consistent with the presence of four OH groups and 434 an MW of 452 for the derivatized compound and 164 for the underivatized compound. The presence of peaks at m/z 347 435 [M^{+.} - 105], and 335 (M^{+.} - 117) are consistent with a compound bearing alcoholic and carboxylic OH groups 436 437 simultaneously. This mass spectrum is similar to the one from methyltartaric acid reported previously from isoprene 438 oxidation by our group (Jaoui et al., 2019). The silylated methyltartaric acid mass spectrum (Jaoui et al., 2019) and mass
439 spectra associated with group 2 are only slightly different, however, they elute at different retention times. The peaks have
440 been tentatively identified as isomers of trihydroxy-oxo-pentanoic acid, with the structure of 4-oxo-D-arabonic acid
441 isomer shown in Table 6.

Group 3 consists of four peaks eluting between 32.5 and 34 min (figure 4: red). The EI and CI mass spectra associated 442 with each peak display similar fragment and adduct ions across the range of 50 to 600 Da. As a descriptive example, an 443 444 EI mass spectrum is shown in Figure 5 for peak eluted at 33.1 min. A comprehensive interpretation of EI and CI mass 445 spectra associated with group 3 peaks (Figures 4, 5), allows us to infer the molecular weight (MW) of the underivatized 446 compounds as 180, and MW_{BSTFA} of 540 for the TMS derivatives, with a chemical formulae $C_5H_8O_7$. The compounds corresponding to these four peaks were identified as isomers of C₅-trihydroxydicarboxylic acids. This identification is 447 448 tentative due to the absence of authentic standards, except for peak eluting at 33.47, which was identified as pentaric acid 449 (Table 6) based on authentic standard. The spectra of BSTFA derivatives of the remaining three red peaks are only slightly 450 different from the pentaric acid spectrum (Figure 6); however, they elute at different retention times. The EI mass spectra 451 are also similar to those reported in the literature for a set of C5-aldaric acids-TMS derivatives including xylaric, arabinaric 452 and ribaric acids (Hinton et al., 2008; https://pubchem.ncbi.nlm.nih.gov). Figure 6 shows the structure of pentaric acid 453 and its four isomers (a), the spectra associated with BSTFA derivative of pentaric acid observed in BnOH SOA (b: EI 454 mode), (c: CI-CH₄ mode), and standard (d: EI mode). Figure 6 also shows the structure of the main fragments observed 455 in BSTFA derivative of pentaric acid in EI mode including m/z at 540, 525, 407, 292, 147, and 73 Da. Pentaric acid and 456 its isomers (aldaric acids) are reported to be formed from the oxidation of aldopentose (Hinton, 2008; Derrien et al., 2018), 457 but no evidence has been provided for its presence in SOA samples. In the present study, we successfully identified aldaric acids from the oxidation of BnOH in SOA samples. 458

459

Nitroaromatic compounds (NACs). NACs of secondary origin are a possible contributor to urban OA and not only adversely affect human health and the environment but impact the aerosol optical properties and the atmospheric radiation balance. By understanding the sources of NACs in ambient particles and their chemical identities, we can evaluate their impact on the climate, environment, and human health. Recently, the analytical capabilities associated with BSTFA derivatization have been extended to NACs bearing hydroxyl and carboxylic acid groups (Jaoui et al., 2018). Mass spectra of most silylated NACs, especially methane-CI, are highly specific, reproducible, and produce characteristic fragments 466 useful in determining structural information and molecular weight, when authentic standards are not available (Jaoui et 467 al., 2018). In this study, a detailed analysis of mass spectra associated with peaks in chromatograms Figure 1c (GP) and Figure 2 top (PP) reveals the presence of several peaks presenting similar fragmentation patterns as those reported by 468 469 Jaoui et al. (2018) for species bearing hydroxyl, carboxylic, nitro groups, and benzene ring. Figure 7 shows the portion 470 between 23 and 42 min of two +EI extracted ion chromatograms for the BSTFA derivatives at m/z 210, 165 (IS), 299 (IS), 471 300, 298, 372, 388 (merged in one chromatogram) associated with BnOH/NOx (top) and BnOH/H₂O₂ (bottom). The EI 472 and/or CI mass spectra of selected nitroaromatic standards can be found in Jaoui et al. (2018), and additional representative 473 subset of the derivatives are displayed in Figures S7 (SI). For clarity, figure 7 inset shows an expanded portion of the top 474 chromatogram between 26.7 – 28 min. Table 7 contains proposed identification of NACs detected in this study, along with molecular weights, formulae, main 5 intense ions associated with CI and EI mass spectra of the derivatives, proposed 475 476 structure, and the GP to PP peak area ratio.

477 NACs with the highest confidence assignment are those identified by comparing their retention times, EI, and CI 478 mass spectra with those of reference standards, and NACs with low levels of confidence are those (1) that have been 479 identified previously in ambient PM or in smog chamber studies, (2) their EI mass spectra exist in the literature, or (3) 480 their molecular weights and numbers of OH, COOH, and NO₂ groups are simply consistent with the CI and EI mass 481 spectrum (Jaoui et al., 2018). A total of fourteen peaks associated with NACs were detected in this study. 3-Nitrobenzyl 482 alcohol, 4-nitrocatechol, 2-hydroxy-5-nitro benzyl alcohol, and 2-nitrophloroglucinol were identified based on authentic 483 standards. Three peaks eluted at 33.76, 34.70, and 34.76 having similar mass spectra as 2-nitrophloroglucinol (main peak) 484 eluted at 35.62 min were detected. They were tentatively associated with homologous series of 2-nitrophloroglucinol 485 including 3-nitrobenzene-1,2,4-triol, 5-nitropyrogallol, and 4-nitro-1,2,3-benzenetriol (not shown in Table 7). Similarly, three additional peaks having similar mass spectra as 2-hydroxy-5-nitrobenzyl alcohol were observed and were tentatively 486 487 associated with homologous series of 2-hydroxy-5-nitrobenzyl alcohol including 4-hydroxy-2-nitrobenzyl alcohol. The 488 EIC in figure 7 (top) includes a series of four peaks observed only in the PP eluting at 35.94, 36.60, 38.18 min, whose 489 mass spectra were consistent with the presence of molecular weight 185 and 401 for the underivatized and derivatized 490 compounds, respectively. Based on similarity of their mass spectra, they were tentatively identified as structural 491 homologue of 3,4-dihydroxy-5-nitrobenzyl alcohol (Table 7) with $C_7H_7NO_5$ formulae. As can be seen in figure 7 (bottom), 492 NACs peaks were not detected in BnOH/H₂O₂ SOA extract, consistent with the formation of NACs in the presence of 493 NOx. All NACs were detected in both GPs and PP (Table 7), except 2-nitrophloroglucinol and 3,4-dihydroxy-5494 nitrobenzyl alcohol and their isomers were observed only in the PP consistent with their low volatility. This result suggests 495 that NACs may be formed in the GP, and partition to the PP for those with low volatility, although PP reactions may 496 occur as suggested by Charan et al. (2020) who analyzed only PP. 4-Nitrocatechol and 2-nitrophloroglucinol were among the largest NAC peaks observed in our study (Figure 7). All three experiments conducted in this study were analyzed for 497 498 NACs to probe reproducibility of the BSTFA method and showed consistent results. 2-Nitrophloroglycinol, 4nitrocatechol and other NACs has been reported in PM collected in Pico Mountain Observatory, Pico Island in the Azores 499 500 archipelago by Ikemori et al., (2019). A series of NACs have been reported recently by Charan et al. (2020) in BnOH 501 SOA using off-line UPLC/ESIQ-ToFMS (ultra-high-performance liquid chromatography electrospray ionization 502 quadruple time of flight mass spectrometry), and the structure assigned to formulas obtained from MassLynk software 503 was based on expected oxidation products and MS/MS analysis. These observations support the identification of NACs 504 reported in this study. 4-Nitrophenol was reported in the GP by Bernard et al., (2013) at low yield and by Charan et al. 505 2020 in SOA from the OH radical oxidation of BnOH but was not detected either in the GP or the PP in this study.

506

507 **3.3. Mechanism of product formation**

508 Based on known GP reactions for aromatic compounds, a schematic representation for the reaction of BnOH with 509 OH is presented in schemes 1-3. It is developed to understand the chemistry leading to the main GP and PP products 510 identified experimentally in this study including HOCs and NACs. These schemes incorporate the latest experimental, 511 quantum and kinetic developments of the fate of peroxy/alkoxy benzoyl radicals including autooxidation (Wang, 2015; 512 Sankar et al. 2014, and Namysl et al. 2020). The lines shown in these schemes are either one step or multistep pathways. Rate constants at room temperature of BnOH with OH radical, O₃, and NO₃ radical of 2.8 10⁻¹¹, 6 x 10⁻¹⁹ (upper limit), 513 and 4.0 x 10⁻¹⁵ cm³ molecule⁻¹ s⁻¹, respectively, have been reported in the literature (Harrison and Wells, 2009, 2012; 514 515 Bernard et al., 2013). This suggests that the day-time oxidation of BnOH will be mainly initiated by OH radicals. The reaction for O₃ and NO₃ radical are not included in schemes 1-3, although they are expected to be formed as minor 516 517 products in our systems.

518 The reaction of BnOH with OH radicals is initiated primarily by H atom abstraction from the external CH₂ group

519 leading to BnAld, and OH addition to the aromatic *ipso* (C1) and *ortho* (C2 or C6) positions to form two alkyldihydroxy

- 520 adducts R1 and R2 (scheme 1). The OH addition to the *para* (C3, C5) and *meta* (C4) position was reported to be not
- 521 favourable based on theoretical study of Wang, (2015). Scheme 1 shows mechanistic pathways leading to the formation



Scheme 1. Initial reaction pathways proposed to produce selected products detected in this study (blue color) in the gas or PP (Table 5). R1-26OO and R2-13OO intermediates undergo further reactions leading to ring-opening products as shown in scheme 3.

527 of stable products (blue) including BnAld, 4-hydroxybenzyl alcohol, 2-hydroxybenzyl alcohol, phenol, formaldehyde, 528 and catechol. The initial branching ratios shown in scheme 1 are those reported by Wang (2015), obtained by combining 529 quantum chemistry calculations and experimental work from the literature. BnAld was observed in this study in the 530 presence and absence of NOx, and its secondary chemistry may lead in part to oxygenated compounds observed in this 531 study (Bernard et al., 2013). Due to large number of possible intermediates formed (Wang, 2015), only selected pathways 532 energetically favourable leading to some products observed in this study are considered. We refer the readers to Wang 533 (2015) paper for an in-depth theoretical analysis of mechanistic pathways leading to the formation of selected reaction 534 products. The adduct R1 reacts rapidly through addition of O_2 to the ortho (C2) to produce peroxy radicals R1-2OO, The 535 O₂ addition to para position (C4) leading to R1-4OO peroxy radicals (not shown in Scheme 1) was found to be endothermic, therefore negligible (Wang, 2015). Radicals R1-2OO undergo intramolecular H-shifts or ring closures to 536 537 form a stable bicyclic intermediate R1-2600 (red). Similarly, R2 reacts rapidly with O₂ to form peroxy radical R2-100 538 intermediate, which itself undergoes intramolecular H-shifts or ring closures to form a stable bicyclic intermediate R2-539 1300 (red). R1-2600 and R2-1300 intermediates undergo further reactions leading to ring-opening products as shown 540 in scheme 3 below. 2-Hydroxybenzyl alcohol was proposed by Wang (2015) to form through the reaction of R2 with O_2 541 involving rapid direct H-abstraction. A possible formation pathway of phenol is decomposition of the peroxy radicals R1-542 200 and R2-100 through CH₂OH radical elimination (Bernard et al., 2013). CH₂OH radical reacts rapidly with O₂ to 543 produce formaldehyde. Catechol was proposed to originate from the reaction of OH radicals with phenol (Atkinson et al., 544 1992) and with 2-hydroxybenzyl alcohol (Bernard et al., 20013).

545 NACs observed in this study (Table 7) are expected to be formed through reaction of OH radicals with BnOH in the 546 presence of NO₂. Scheme 2 briefly summarizes the main mechanistic pathways leading to BnOH NACs, which follow 547 similar chemistry as those reported for toluene, benzene, and xylenes (Jenkin et al., 2003; Vidovic et al., 2018) and 548 summarized by Wang et al., (2019). The steps shown in scheme 2 are multi-steps and the reader should consult the 549 reference papers above for more in depth information. NACs are proposed to originate from secondary reactions of 550 catechol, 1,3,5-trihydroxy benzene, and 4-hydroxybenzyl alcohol with OH radicals in the presence of NO2 (scheme 2). 551 These intermediates are proposed to be originated from R1, R2, and R3 adducts. Additional pathways could be initiated via less well understood aqueous-phase nitration (Kroflic et al., 2018). 4-Nitrocatechol is proposed to be initiated through 552 the reaction of catechol with OH radicals in the presence of NOx (Finewax et al., 2018). 4-Hydroxy-2-nitrobenzyl alcohol 553 554 is proposed to be likely originated from the alkyldihydroxy-para-adduct formed from the OH addition to para position

- 555 (scheme 2). 2-Nitrophloroglucinol and 3,4-dihydroxy-5-nitrobenzyl alcohol follow similar reactions involving R3 adduct,
- 556 OH radicals, and NO₂. According to Wang (2015) calculation, at high NO₂ (100 ppbv) the reaction of R1 with NO₂ can
- 557 compete to a minor degree with the reaction with O₂, therefore R1 possibly forms minor amounts of nitrobenzyl alcohol.





560 Scheme 2. Proposed mechanism for selected NAC species observed in this study.

561 HOCs were detected in the PP from the oxidation of BnOH in both low and high NOx systems (Table 6). Mechanistic 562 pathways based on theoretical studies leading to several HOCs (e. g. HOMs) from the atmospheric oxidation of biogenic 563 and aromatic hydrocarbons have been reported recently in the literature involving unimolecular reaction through autoxidation, and peroxy and/or alkoxy radical isomerization (Wang, 2015; Jaoui et al., 2021; Piletic and Kleindienst, 564 2022). The formation of selected HOCs observed in this study is consistent with the following pathways proposed in 565 scheme 3 involving R2-1300 radical as the starting material. R1-2600 adduct undergoes similar reactions leading to 566 567 butenedial and 2,3-epoxy-butanedial as shown in scheme S1 (SI). Tartaric acid, 2,3,5-trihydroxy-4-oxo-pentanoic acid, and pentaric acid, observed in this study for the first time, are proposed to be initiated by the oxidation of butenedial/2,3-568 569 epoxy-butanedial, 5-hydroxy-4-oxo-2-penatenal, and 4-hydroxy-2,3-epoxypentandial (scheme 3). The mechanism



571 **Scheme 3.** Proposed mechanism for selected highly oxygenated compounds observed in this study.

573 leading to butenedial/2,3-epoxybutanal, 5-hydroxy-4-oxo-2-penatanal (scheme S1) was reported by Wang (2015) from 574 the OH oxidation of BnOH, therefore is not shown in scheme 3. Tartaric acid present in the PP at high level (Figure 4), is proposed to be formed through the oxidation of butenedial and/or 2,3-epoxybutanal through classical oxidation of 575 576 aldehydes and alkenes to carboxylic acid (not shown in scheme 3). Similarly, 2,3,5-trihydroxy-4-oxo-pentanoic acid and 577 pentaric acid are proposed to rise from the oxidation of 5-hydroxy-4-oxo-2-pentenal, and 4-hydroxy-2,3-578 epoxypentanedial, respectively following similar mechanistic pathways reported by Jaoui et al. (2021) for the formation 579 of methyltartaric acid from 4-hydroxy-2-methyl-but-2-enal involving peroxy and alkoxy radical isomerization (not 580 reported here). In this study, a new mechanism is proposed in scheme 3 leading to the formation of 4-hydroxy-2,3-581 epoxypentanedial, which is the starting material for pentaric acid formation. It involves several intermediate steps 582 including unimolecular H migration (e.g., 1,5-H shift), ring opening and decomposition. Formaldehyde and glyoxal 583 observed in this study are also shown in scheme 3.

584

585 **4. Summary**

586 In the present manuscript, laboratory experiments were conducted to investigate SOA formation from the oxidation 587 of benzyl alcohol in the presence and absence of NOx. Chamber aerosol collected under these conditions has been 588 analyzed for organic mass to organic carbon ratio, and aerosol yield. In addition, the chemical composition of the gas 589 phase and SOA was analyzed using derivative-based methods followed by gas chromatography-mass spectrometry and 590 high-performance liquid chromatography analysis of the derivative compounds. More than 50 oxygenated organic 591 compounds in the gas and particle phases were identified. While a detailed non-targeted analysis has been made, our 592 primary focus has been to examine highly oxygenated and nitroaromatic compounds. The major components include ring-593 opening products with high oxygen to carbon ratio (e. g. malic acid, tartaric acid, arabic acid, 2,3,5-trihydroxy-4-oxo-594 pentanoic acid, and pentaric acid) and ring-retaining products (e. g. benzaldehyde, benzoic acid, catechol, 3-nitrobenzyl 595 alcohol, 4-nitrocatechol, 2-hydroxy-5-nitrobenzyl alcohol, 2-nitrophloroglucidol, 5-(hydroxymethyl)- 3-nitro-1,2-benzyl 596 diol). The presence of some of these products in the gas and particle phases simultaneously provides evidence of their 597 gas/particle partitioning. These oxygenated oxidation products made dominant contributions to the SOA particle 598 composition in both low and high NOx systems. Yields, organic mass to organic carbon ratio, and proposed reaction 599 schemes for selected compounds are provided.

600	Finally, a set of reaction pathways are proposed that accounts for selected reaction products observed in this study
601	from BnOH photooxidation in the presence of OH radicals, including NACs and HOCs. The proposed mechanism is
602	based on (1) theoretical studies reported previously in the literature and (2) mechanisms associated with aromatics
603	oxidation (e.g., benzene, toluene, xylenes). New pathways were proposed for the formation of newly observed highly
604	oxygenated compounds tartaric acid, 2,3,5-trihydroxy-4-oxopentanoic acid, and pentaric acid. Butenedial/2,3epoxy-
605	butandial, 5-hydroxy-4-oxo-2-pentenal, and 4-hydroxy-2,3-expoxypentanedial were proposed as the starting intermediate
606	species leading to these highly oxygenated compounds. While theoretical studies involving unimolecular reactions were
607	developed focusing mainly on ring-containing products (Wang, 2015, Piletic and Kleindienst, 2022), similar theoretical
608	investigations focusing on linear species (Jaoui et al., 2021) as HOCs reported in this study will help strengthen the
609	pathways proposed here.
610	The results of this study potentially have atmospheric implications for areas impacted by benzyl alcohol including
611	urban and indoor areas and contribute to understanding the formation of ambient SOA from oxygenated anthropogenic
612	precursors. Nitroaromatics are pollutants of concern due to their toxicity, light-absorption properties, and relatively long
613	residence times in the environment. HOCs may partition into pre-existing particles or be involved in new particle
614	formation.
615 616 617	
618	Data Availability. The data used in this study can be found at: https://catalog.data. gov/dataset/epa-sciencehub. DOI:
619	10.23719/1527893.
620	Commeting interests. The outhors declars no commeting financial interest
621	Competing interests. The authors declare no competing mancial interest.
622	Disalainer This work has been subjected to the U.S. Environmental Protection Agency's administrative review and
624	<i>Discutter</i> . This work has been subjected to the U.S. Environmental Protection Agency's administrative review and
024	approved for publication. The views expressed in this article are those of the authors and do not necessarily represent the
625	views or policies of the U.S. Environmental Protection Agency. Mention of trade names does not constitute endorsement
626	or recommendation of a commercial product by U.S. EPA.
627	
628	

630 References

- Abend, A. M., Chung, L., Bibart, R. T., Brooks, M., and McCollum, D. G.: Concerning the stability of benzyl alcohol:
 formation of benzaldehyde dibenzyl acetal under aerobic conditions, J. Pharm. Biomed. Anal., 34, 5, 957-962,
 doi:10.1016/j.jpba.2003.11.007, 2004.
- 634
- Akherati, A., Cappa, C. D., Kleeman, M. J., Docherty, K. S., Jimenez, J. L., Griffith, S. M., Dusanter, S., Stevens, P. S.,
- and Jathar, S. H.: Simulating secondary organic aerosol in a regional air quality model using the statistical oxidation
- 637 model Part 3: Assessing the influence of semi-volatile and intermediate-volatility organic compounds and NOx, Atmos.
- 638 Chem. Phys., 19, 4561–4594, 2019.
- 639
- 640 Alton, M. W., and Browne, L. C.: Atmospheric chemistry of volatile methyl siloxanes: kinetics and products of oxidation
- by OH radicals and Cl atoms, Environ. Sci. Technol., 54, 5992–5999, 2020.
- 642
- Antonelli, L., Mapelli, E., Strini, A., Cerulli, T., Leoni, R., and Stella S.: Laboratory and real scale comparative study of
 benzyl alcohol emission from a two-component epoxy paint, Proceedings: Indoor Air, 584-589, 2002.
- 645
- Atkinson, R., Aschmann, S. M., and Arey, J.: Reactions of OH and N03 Radicals with Phenol, Cresols, and 2-Nitrophenol
 at 296 ± 2 K, Environ. Sci. Technoi. 1992, 26, 1397-1403, 1992.
- 648
- Bernard, B., Magneron, I., Eyglunent, G., Daële, V., Wallington, T. J., Hurley, M. D., and Mellouki, A.: Atmospheric
 chemistry of benzyl alcohol: kinetics and mechanism of reaction with OH radicals, Environ. Sci. Technol., 47,
 3182–3189, 2013.
- 652
- Berndt, T., Herrmann, H., Sipila, M., and Kulmala, M.: Highly oxidized second-generation products from the gas-phase
 reaction of OH radicals with isoprene. J. Phys. Chem. A, 120 (51), 10150–10159, 2016.
- 655
- 656 Boatright, J., Negre, F., Chen, X., Kish, C. M., Wood, B., Peel, G., Orlova, I., Gang, D., Rhodes, D., and Dudareva, N.:
- Understanding in vivo benzenoid metabolism in petunia petal tissue, Plant Physiol., 135, 1993–2011, 2004.

659	Carter, W. P. L., Malkina, I. L., Cocker III, D. R., and Song, C.: Environmental chamber studies of VOC species in
660	architectural coating and mobile source emissions, South Coast Air Quality Management District Contract No. 03468,
661	2005.
662	
663	Charan, S. M., Buenconsejo, R. S., and Seinfeld, J. H.: Secondary organic aerosol yields from the oxidation of benzyl
664	alcohol, Atmos. Chem. Phys., 20, 13167–13190, doi:10.5194/acp-2020-49, 2020
665	
666	Charan, S. M., Huang, Y., Buenconsejo, R. S., Li, Q., Cocker III, D. R., and Seinfeld, J. H.: Secondary organic aerosol
667	formation from the oxidation of decamethylcyclopentasiloxane at atmospherically relevant OH concentrations, Atmos.
668	Chem. Phys., 22, 917-928, doi:10.5194/acp-22-917-2022, 2021.
669	
670	Cheng, C. T., Chan, M. N., and Wilson, K. R.: Importance of unimolecular HO2 elimination in the heterogeneous OH
671	reaction of highly oxygenated tartaric acid aerosol, J. Phys. Chem. A, 120, 5887-5896, doi: 10.1021/acs.jpca.6b05289,
672	2016.
673	
674	Coggon, M. M., Gkatzelis, G. I., McDonald, B. C., Gilman, J. B., Schwantes, R. H., Abuhassan, N., Aikin, K. C., Arendd,
675	M. F., Berkoff, T. A., Brown, S. S., Campos, T. L., Dickerson, R. R., Gronoff, G., Hurley, J. F., Isaacman-VanWertz, G.,
676	Koss, A. R., Lia, M., McKeen, S. A., Mosharyd, F., Peischl, J., Pospisilova, V., Renh, X., Wilson, A., Wu, Y., Trainer,
677	M., and Warneke, C.: Volatile chemical product emissions enhance ozone and modulate urban chemistry,
678	doi:10.1073/pnas.2026653118, PNAS, 118, 32, e2026653118, 2021.
679	
680	DeBolt, S., Cook, D. R., and Ford, C. M.: L-Tartaric acid synthesis from vitamin C in higher plants, PNAS, 103 (14) 5608-
681	5613, doi: 10.1073/pnas.0510864103, 2006.

- 682
- Derrien, E., Ahmar, M., Martin-Sisteron, E., Raffin, G., Queneau, Y., Marion, P., Beyerle, M., Pinel, C., and Besson, M.:
- 684 Oxidation of aldoses contained in softwood hemicellulose acid hydrolysates into aldaric acids under alkaline or

- noncontrolled pH conditions, Industrial & Engineering Chemistry Research, 57 (13), 4543-4552, doi.:
 10.1021/acs.iecr.8b00239, 2018.
- 687
- Do, J. Y., Salunkhe, D. K., and Olson, L. E.: Isolation, identification and comparison of the volatiles of peach fruit as
 related to harvest maturity and artificial ripening, J. Food Sci., 34, 618–621, 1969.
- 690
- Ferri, D., Mondelli, C., Krumeich, F., and Baiker, A.: Discrimination of active palladium sites in catalytic liquid-phase
 oxidation of benzyl alcohol, J. Phys. Chem. B., 110, 46, 22982-22986, doi:10.1021/jp065779z, 2006.
- 693
- Finewax, Z., de Gow, J. A., and Ziemann, P. J.: Identification and Quantification of 4-Nitrocatechol Formed from OH
 and NO₃ Radical-Initiated Reactions of Catechol in Air in the Presence of NO_x: Implications for Secondary Organic
 Aerosol Formation from Biomass Burning, Environmental Science & Technology, 52 (4), 1981-1989, doi.:
 10.1021/acs.est.7b05864, 2018.
- 698
- Fu, Zi., Xie, H., Elm, J., Guo, X., Fu, Zh., and Chen, J.: Formation of low-volatile products and unexpected high
 formaldehyde yield from the atmospheric oxidation of methylsiloxanes, Environ. Sci. Technol., 54, 12, 7136-7145,
 doi:10.1021/acs.est.0c01090, 2020.
- 702
- Gkatzelis, G. I., Coggon, M. M., McDonald, B. C., Peischl, J., Aikin, K. C., Gilman, J. B., Trainer, M., and Warneke, C.:
 Identifying volatile chemical product tracer compounds in U.S. Cities, Environ. Sci. Technol.,
 doi:10.1021/acs.est.0c05467, 55, 188–199, 2021.
- 706
- Gowda, D., Kawamura, K., and Tachibana, E.: Identification of hydroxy- and keto-dicarboxylic acids in remote marine
 aerosols using gas chromatography/quadruple and time-of-flight mass spectrometry, Rapid Communications in Mass
 Spectrometry, 30(7), 992–1000, doi: 10.1002/rcm.7527, 2016.
- 710
- 711 Harrison, J. C., and Wells, J. R.: Gas-phase chemistry of benzyl alcohol: reaction rate constants and products with OH
- 712 radical and ozone, Atmos. Environ., 43, 798–804, 2009.

- Harrison, J. C., and Wells, J. R.: 2-Butoxyethanol and benzyl alcohol reactions with the nitrate radical: rate coefficients
 and gas-phase products, Int. J. Chem. Kinet., 44, 778–788, 2012.
- 716
- 717 Hayes, P. L., Carlton, A. G., Baker, K. R., Ahmadov, R., Washenfelder, R. A., Alvarez, S., Rappenglück, B., Gilman, J.
- B., Kuster, W. C., de Gouw, J. A., Zotter, P., Prévôt, A. S. H., Szidat, S., Kleindienst, T. E., Offenberg, J. H., Ma, P. K.,
- and Jimenez, J. L.: Modeling the formation and aging of secondary organic aerosols in Los Angeles during CalNex 2010,
- 720 Atmos. Chem. Phys., 15, 5773–5801, 2015.
- 721
- 722 Hinton, M. R.: Xylaric acid, D-arabinaric acid (D-lyxaric acid), L-arabinaric acid (L-lyxaric acid), and Ribaric acid-1,4-
- 723 lactone; Synthesis and isolation-synthesis of polyhydroxypolyamides therefrom, Theses, Dissertations, & Professional
- Papers. 1202, https://scholarworks.umt.edu/etd/1202, 2008.
- 725
- Hodzic, A., Jimenez, J. L., Madronich, S., Aiken, A. C., Bessagnet, B., Curci, G., Fast, J., Lamarque, J.-F., Onasch, T.
 B., Roux, G., Schauer, J. J., Stone, E. A., and Ulbrich, I. M.: Modeling organic aerosols during MILAGRO: importance
 of biogenic secondary organic aerosols, Atmos. Chem. Phys., 9, 6949–6981, 2009.
- 729
- Horvat, R. J., Chapman, G. W., Jr., Robertson, J. A., Meredith, F. I., Scorza, R., Callahan, A. M., and Morgens, P.:
 Comparison of the volatile compounds from several commercial peach cultivars, J. Agric. Food Chem., 38, 234–237,
 1990.
- 733
- Humes, M. B., Wang, M., Kim, S., Machesky, J. E., Gentner, D. R., Robinson, A. L., Donahue, N. M., and Presto, A. A.:
 Limited secondary organic aerosol production from acyclic oxygenated volatile chemical products, Environ. Sci. Technol.
 56, 4806–4815, 2022.

- Humpf, H. U., and Schreier, P.: Bound aroma compounds from the fruit and the leaves of blackberry (Rubus laciniata L.),
 J. Agric. Food Chem., 39, 1830–1832, 1991.
- 740

741 Ikemori, E., Nakayama, T., and Hasegawa, H.: Characterization and possible sources of nitrated mono- and di-aromatic
742 hydrocarbons containing hydroxyl and/or carboxyl functional groups in ambient particles in Nagoya, Japan, Atmos.

743 Environ., 211, 91-102, 2019.

744

Janechek, N. J., Marek, R. F., Bryngelson, N., Singh, A., Bullard, R. L., Brune, W. H., and Stanier, C. O.: Physical
properties of secondary photochemical aerosol from OH oxidation of a cyclic siloxane, Atmos. Chem. Phys., 19, 1649–
1664, 2019.

748

Jaoui, M., and Kamens, R. M.: Mass balance of gaseous and particulate products analysis from α -pinene/NO_x/air in the presence of natural sunlight, J. Geophys. Res., 106, D12, 12,541-12,558, doi:10.1029/2001JD900005, 2001.

751

Jaoui, M., Kleindienst, T. E., Lewandowski, M., and Edney, E. O.: Identification and quantification of aerosol polar
oxygenated compounds bearing carboxylic and/or hydroxyl groups, 1. Method development, Anal. Chem., 76, 4765–
4778, 2004.

755

Jaoui, M., Kleindienst, T. E., Lewandowski, M., Offenberg, J. H., and Edney, E. O.: Identification and quantification of aerosol polar oxygenated compounds bearing carboxylic or hydroxyl groups. 2. Organic tracer compounds from monoterpenes, Environ. Sci. Technol., 39, 5661–5673, 2005.

759

- Jaoui, M., Kleindienst, T. E., Docherty, K. S., Lewandowski, M., and Offenberg, J. H.: Secondary organic aerosol
 formation from the oxidation of a series of sesquiterpenes: a-cedrene, b-caryophyllene, a-humulene and a-farnesene with
 O₃, OH and NO₃ radicals, Environ. Chem. 10, 178–193, doi:10.1071/EN13025, 2013.
- 763
- Jaoui, M., Lewandowski, M., Docherty, K., Offenberg, J. H., and Kleindienst, T. E.: Atmospheric oxidation of 1,3-
- butadiene: characterization of gas and aerosol reaction products and implications for PM_{2.5}, Atmos. Chem. Phys., 13681–
- 766 13704, doi: 10.5194/acp-14-1368114, 2014.

768 Jaoui, M., Lewandowski, M., Offenberg, H. J., Colon, M., Docherty, K. S., and Kleindienst, T. E: Characterization of

Jaoui, M., Szmigielski, R., Nestorowicz, K., Kolodziejczyk, A., Sarang, K., Rudzinski, K. J., Konopka, A., Bulska, E.,

Lewandowski, M., And Kleindienst, T. E.: Organic hydroxy acids as highly oxygenated molecular (HOM) tracers for

- aerosol nitroaromatic compounds: Validation of an experimental method, Mass Spectrom., 53, 680–692, 2018.
- 770

771

⁷⁷³ aged isoprene aerosol, Environmental Science & Technology, 53 (24), 14516-14527, doi: 10.1021/acs.est.9b05075, 2019. 774 775 Jaoui, M., Piletic, I., Szmigielski, R., Rudzinski, J. K., E, Lewandowski, M., Riedel, T. P., and Kleindienst, T. E.: Rapid 776 production of highly oxidized molecules in isoprene aerosol via peroxy and alkoxy radical isomerization pathways in low 777 and high NO_x environments: Combined laboratory, computational and field studies, Science of The Total Environment, 778 775, 145592, doi: 10.1016/j.scitotenv.2021.145592, 2021 779 780 Jenkin, M. E., Saunders, S. M., Wagner, V., and Pilling, M. J.: Protocol for the development of the Master Chemical 781 Mechanism, MCM v3 (Part B): tropospheric degradation of aromatic volatile organic compounds, Atmos. Chem. Phys., 782 3, 181–193, https://doi.org/10.5194/acp-3-181-2003, 2003. 783 784 Khare, P., and Gentner, D. R.: Considering the future of anthropogenic gas-phase organic compound emissions and the 785 increasing influence of non-combustion sources on urban air quality, Atmos. Chem. Phys., 18, 5391–5413, 2018. 786 787 Kleindienst, T. E., Edney, E. O., Lewandowski, M., Offenberg, J. H., and Jaoui M.: Secondary organic carbon and aerosol 788 yields from the irradiations of isoprene and \Box -pinene in the presence of NO_x and SO₂, Environ. Sci. Technol., 40, 3807– 789 3812, 2006. 790 791 Kleindienst, T. E., Lewandowski, M., Offenberg, J. H., Jaoui, M., and Edney, E. O.: The formation of secondary organic 792 aerosol from the isoprene + OH reaction in the absence of NO_x, Atmos. Chem. Phys., 9, 6541–6558, 2009. 793

794	Kleindienst, T. E., Jaoui, M., Lewandowski, M., Offenberg, J. H., and Docherty, K. S.: The formation of SOA and
795	chemical tracer compounds from the photooxidation of naphthalene and its methyl analogs in the presence and absence
796	of nitrogen oxides, Atmos. Chem. Phys., doi:10.5194/acp-12-8711-2012, 12, 8711-8726, 2012.

Kroflic, A., Hus, M., Grilc, M., and Grgic, I.: Underappreciated and complex role of nitrous acid in aromatic nitration
under mild environmental conditions: the case of activated methoxyphenols, Environ. Sci. Technol., 52, 13756–13765,
https://doi.org/10.1021/acs.est.8b01903, 2018.

- 801
- Kroll, J. H., Chan, A. W. H., Ng, N. L., Flagan, R. C., and Seinfeld, J. H.: Reactions of semivolatile organics and their
 effects on secondary organic aerosol formation, Environ. Sci. Technol., 41, 3545–3550, 2007.
- 804
- Larsen, M., and Poll, L.: Odor thresholds of some important aroma compounds in raspberries, Z. Lebensm. Unters. Forsch,
 191, 129–131, 1990.
- 807
- Lewandowski, M., Jaoui, M., Offenberg, J. H., Krug, J. D., and Kleindienst, T. E.: Atmospheric oxidation of isoprene and
 1,3-butadiene: influence of aerosol acidity and relative humidity on secondary organic aerosol, Atmos. Chem. Phys., 15,
 3773–3783, doi:10.5194/acp-15-3773-2015, 2015.
- 811
- Li, W., Li, L., Chen, C-L, Kacarab, M., Peng, W., Price, D., Xu, J., and Cocker III, D. R.: Potential of select intermediate-

volatility organic compounds and consumer products for secondary organic aerosol and ozone formation under relevant
urban conditions, Atmos. Environ., 118, 109-117, 2018.

- 815
- Lu, Q., Murphy, B. N., Momei Q., Adams, P. J., Zhao, Y., Pye, H. O. T., Efstathiou, C., Allen, C., and Robinson, A. L.:

817 Simulation of organic aerosol formation during the CalNex study: updated mobile emissions and secondary organic

818 aerosol parameterization for intermediate-volatility organic compounds, Atmos. Chem. Phys., 20, 4313–4332, 2020.

- 820 McDonald, B. C., De Gouw, J. A., Gilman, J. B., Jathar, S. H., Akherati, A., Cappa, C. D., Jimenez, J. L., Lee-Taylor, J.,
- 821 Hayes, P. L., McKeen, S. A., Cui, Y. Y., Kim, S. W., Gentner, D. R., Isaacman-VanWertz, G., Goldstein, A. H., Harley,

- R. A., Frost, G. J., Roberts, J. M., Ryerson, T. B., and Trainer, M.: Volatile chemical products emerging as largest
 petrochemical source of urban organic emissions, Science, 359, 760–764, 2018.
- 824
- Milani, A., Al-Naiema, I. M., and Stone, E. A: Detection of a secondary organic aerosol tracer derived from personal care
 products, Atmos. Environ., 246, 118078, 2021.
- 827
- 828 Mohr, C., DeCarlo, P. F., Heringa, M. F., Chirico, R., Richter, R., Crippa, M., Querol, X., Baltensperger, U., and Prévôt,
- A. S. H.: Spatial variation of aerosol chemical composition and organic components identified by positive matrix
- factorization in the Barcelona region, Environ. Sci. Technol., 49, 10421–10430, 2015.
- 831
- Namysl, S., Pelucchi, M., Maffei, L. P., Herbinet, O., Stagni, A., Faravelli, T., and Battin-Leclerc, F.: Experimental and
 modeling study of benzaldehyde oxidation, Combustion and Flame, 211, 124–132, 2020.
- 834
- Offenberg, J. H., Lewandowski, M., Edney, E. O., Kleindienst, T. E., Jaoui, M.: Investigation of a systematic offset in the
 measurement of organic carbon with a semicontinuous analyzer, J. A&WMA, 57:5, 596-599, doi:10.3155/10473289.57.5.596, 2007
- 838
- 839 Orlova, I., Marshall-Colón, A., Schnepp, J., Wood, B., Varbanova, M., Fridman, E., Blakeslee, J. J., Peer, W. A., Murphy,
- A. S., Rhodes, D., Pichersky, E., and Dudareva, N.: Reduction of Benzenoid synthesis in petunia flowers reveals multiple
 pathways to benzoic acid and enhancement in auxin transport, Plant Cell, 18, 3458–3475, 2006.
- 842
- Pennington, E. A., Seltzer, K. M., Murphy, B. N., Qin, M., Seinfeld, J. H., Pye, H. O. T.: Modeling secondary organic
 aerosol formation from volatile chemical products, Atmos. Chem. Phys., doi:10.5194/acp-21-18247-18261-2021, 1824718261, 2021.
- 846
- 847 Piletic, I. R., and Kleindienst, T. E.: Rates and Yields of Unimolecular Reactions Producing Highly Oxidized Peroxy
- 848 Radicals in the OH-Induced Autoxidation of α -Pinene, β -Pinene, and Limonene, *The Journal of Physical Chemistry A*.
- 849 126 (1), 88-100, doi: 10.1021/acs.jpca.1c07961, 2022.

851	Qin, M. M., Murphy, B. N., Isaacs, K. K., McDonald, B. C., Lu, Q. Y., McKeen, S. A., Koval, L., Robinson, A. L.,
852	Efstathiou, C., Allen, C., and Pye, H. O. T.: Criteria pollutant impacts of volatile chemical products informed by near-
853	field modelling, Nature Sustainability, 4, 129–137, https://doi.org/10.1038/s41893-020-00614-1, 2021.
854	
855	Rohl, A., and Lammedl. G.: Determination of malic acid and other C4 dicarboxylic acids in atmospheric aerosol samples,
856	Chemosphere, 46(8), 1195-1199, doi: 10.1016/s0045-6535(01)00243-0, 2002.
857	
858	Sankar, S., Nowicka, E., Carter, E., Murphy, D. M., Knight, D. W., Bethell, D., and Hutchings, G. J.: The benzaldehyde
859	oxidation paradox explained by the interception of peroxy radical by benzyl alcohol, Nature Communic,
860	doi:10.1038/ncomms4332, 5, 3332, 2014.
861	
862	Seltzer, K. M., Murphy, B. N., Pennington, E. A., Allen, C., Talgo, K., and Pye, H. O. T.: Volatile chemical product
863	enhancements to criteria pollutants in the United States, Environ. Sci. technol., doi:10.1021/acs.est.1c04298, 2021.
864	
865	Shilling, J. E., Chen, Q., King, S. M., Rosenoern, T., Kroll, J. H., Worsnop, D. R., McKinney, K. A., and Martin, S. T.:
866	Particle mass yield in secondary organic aerosol formed by the dark ozonolysis of α -pinene, Atmos. Chem. Phys., 8,
867	2073–2088, doi:10.5194/acp-8-2073-2008, 2008.
868	
869	Smith, D. F., Kleindienst, T. E., and Hudgens, E. E.: Improved high-performance liquid chromatographic method for
870	artifact free measurements of aldehydes in the presence of ozone using 2,4-dinitrophenylhydrazine, J. Chromatogr. A,
871	483, 431–436, 1989.
872	
873	Stockwell, C. E., Coggon, M. M., Gkatzelis, G. A., Ortega, J., McDonald, B. C., Peischl, J., Aikin, K., Gilman, J. B.,
874	Trainer, M., and Warneke, C.: Volatile organic compound emissions from solvent- and water borne coatings:
875	compositional differences and tracer compound identifications, Atmos. Chem. Phys., 21, 6005-6022, doi:10.5194/acp-
876	21-6005-2021, 2021.
877	

878	Urakami, K., Kobayashi, C., Miyazaki, Y., Nishijima, K., and Yoshimura, Y.: Degradation products generated by
879	sonication of benzyl alcohol, a sample preparation solvent for the determination of residual solvents in pharmaceutical
880	bulks, on capillary gas chromatography, Chem. Pharm. Bull., 48, 1299–1303, 2000.
881	
882	Vallat, A., and Dorn, S.: Changes in volatile emissions from apple trees and associated response of adult female codling
883	moths over the fruit-growing season, J. Agric. Food Chem., 53, 4083-4090, 2005.
884	
885	Vidovic, K., Lasic Jurkovic, D., Sala, M., Kroflic, A., and Grgic, I.: Nighttime aqueous-phase formation of nitrocatechols
886	in the atmospheric condensed phase, Environ. Sci. Technol., 52, 9722-9730, https://doi.org/10.1021/acs.est.8b01161,
887	2018.
888	
889	Vlachou, A., Daellenbach, K. R., Bozzetti, C., Chazeau, B., Salazar, G. A., Szidat, S., Jaffrezo, J. L., Hueglin, C.,
890	Baltensperger, U., El Haddad, I., and Prévôt, A. S.: Advanced source apportionment of carbonaceous aerosols by coupling
891	offline AMS and radiocarbon size-segregated measurements over a nearly 2-year period. Atmos. Chem. Phys., 18, 6187-
892	6206, 2018.
893	
894	Wang, L.: The atmospheric oxidation mechanism of benzyl alcohol initiated by OH radicals: the addition channels, Chem.
895	Phys. Chem., 16 (7), 1542-1550, doi:10.1002/cphc.201500012, 2015.
896	
897	Wang, N., Jorga, S. D., Pierce, J. R., Donahue, N. M., and Pandis, S. N.: Particle wall-loss correction methods in smog
898	chamber experiments, Atmos. Meas. Tech., 11, 6577-6588, doi:10.5194/amt-11-6577-2018, 2018.
899	
900	Wang, Y., Hu, M., Wang, Y., Zheng, J., Shang, D., Yang, Y., Liu, Y., Li, X., Tang, R., Zhu, W., Du, Z., Wu, Y., Guo, S.,
901	Wu, Z., Lou, S., Hallquist, M., and Yu, J. Z.: The formation of nitro-aromatic compounds under high NOx and
902	anthropogenic VOC conditions in urban Beijing, China, Atmos. Chem. Phys., 19, 7649–7665, 2019.
903	
904	Weschler, C. J.: Chemistry in indoor environments: 20 years of research, Indoor Air, 21 (3), 205-218, 2011.
905	
	35

906 Wu, Y., and Johnston, M. V.: Molecular characterization of secondary aerosol from oxidation of cyclic methylsiloxanes,

```
907 J. Am. Soc. Mass. Spectr., 27, 402–409, doi:10.1007/s13361-015-1300-1, 2016.
```

- 909 Wu, Y., and Johnston, M. V.: Aerosol formation from OH oxidation of the volatile cyclic methyl siloxane (cVMS)
- 910 Decamethylcyclopentasiloxane, Environ. Sci. Technol., 51, 4445–4451, doi:10.1021/acs.est.7b00655, 2017.
- 212 Zhao, B., Wang, S., Donahue, N. M., Jathar, S. H., Huang, X., Wu, W., Hao, J., and Robinson, A. L.: Quantifying the
- 913 effect of organic aerosol aging and intermediate volatility emissions on regional-scale aerosol pollution in China, Sci.
- 914 Rep., 6, 28815, doi:10.1038/srep28815, 2016.

- 916 Zhang, X., Cappa, C. D., Jathar, S. H., McVay, R. C., Ensberg, J. J., Kleeman, M. J., and Seinfeld, J. H.: Influence of
- 917 vapor wall loss in laboratory chambers on yields of secondary organic aerosol, PNAS, doi:10.1073/pnas.1404727111,
- 918 111 (16), 5802-5807, 2014.

Exp. IDs	BnOH	H ₂ O ₂	NO ^a	Seed surface	BnOH/NO	Т	RH
	(ppb)	(ppm)	(ppb)	area (nm² cm⁻³)	(ppb/ppb)	(°C)	(%)
ER-889	385	-	178	4.67 x 10 ⁷	2.2	24.5	31.0
ER-890	355	-	96	4.94 x 10 ⁷	3.7	24.5	31.1
ER-891	723	-	188	9.88 x 10 ⁷	3.8	24.6	31.3
ER-892	319	3.04	-	1.36 x 10 ⁶	-	25.7	< 4.0

Table 1. Initial conditions for BnOH experiments in the presence and absence of NO.

935 T: temperature; RH: relative humidity. Seed aerosol at $1 \ \mu g \ m^{-3}$. ^a: The initial NOx during the irradiations was greater than 98% NO.

Table 2. Steady-state GP and reacted BnOH and NO concentration during the irradiations.

Exp. IDs	NO Reacted NO		BnOH	Reacted BnOH	BnOH/NO	O ₃	NOy
	(ppb)	(ppb)	(ppb)	(ppb)	ratio	(ppb)	(ppb)
					(ppb/ppb)		
ER889	78	100	132	253	1.7	30	163
ER890	9	87	132	223	14.7	147	80
ER891	29	159	387	336	13.4	11	146
ER892	-	-	85	234	-	28	-

939 Table 3. Formation and yields of SOA (Y_{SOA}) and SOC (Y_{SOC}). All organic and carbon aerosol masses are corrected for

940 a wall loss of 0.067 h^{-1}	(Kleindienst et al., 2012).
-----------------------------------	-----------------------------

Exp. IDs SOA		SOC	SOC SOA/SOC		Y _{SOC}	
	$(\mu g/m^3)$	$(\mu gC/m^3)$		(%)	(%)	
ER889	39.6	23.2	1.7	3.6	2.7	
ER890	56.1	30.3	1.9	5.7	4.0	
ER891	119.5	58.9	2.0	8.1	5.1	
ER892	52.9	24.8	2.1	5.2	3.1	

Table 4. Steady state carbonyl concentrations (ppmV) during BnOH oxidation (FH: formaldehyde; AH: acetaldehyde;

Ac: acetone; MA: methacrolein; BN: 2-butanone; BnAld: benzaldehyde; G: glyoxal; MG:	methylglyoxal).
---	-----------------

Exp. ID	FH	AH	Ac	MA	BN	BnAld	G	MG
ER889	2.4	1.2	1.0	1.0	0.6	23.09	5.0	0.6
ER890	1.5	2.8	-	-	2.9	18.2	3.8	0.4
ER891	5.1	2.5	1.3	2.0	1.4	30.8	8.6	0.6
ER892	181.7	23.5	-	0.8	0.8	5.2	7.8	1.6

- 976 **Table 5.** Summary of selected reaction products detected and identified either in gas-phase (GP), particle phase (PP) or
- both from BnOH/NOx, and BnOH/H₂O₂ experiments. Tables 6 and 7 shows additional aerosol species with high oxygen
- 978 to carbon ratio and/or nitro group. NA: not applicable. ^a: underivatized m/z are given. *: identified with authentic standard.

IUPAC/common nomenclature Benzyl alcohol (BnOH)	Formula C7H8O	m/z BSTFA (EI) 165, 91, 135, 180, 73	MW [MW _{BSTFA}] (g mol ⁻¹) 108 [180]	Proposed Structure	Detected GP
Phenol	C ₆ H ₆ O	73, 151, 166, 94, 65	94 (166)	HO	GP, PP
Benzaldehyde (BnAld)	C ₇ H ₆ O	106, 105, 77, 77, 51	106 (NA)	T T	GP, PP
Benzene-1,2-diol (catechol)	C ₆ H ₆ O ₂	239, 255, 80, 283, 73	110 (254)	н. о	РР
Benzoic acid	C ₇ H ₆ O ₂	179, 105, 135, 77, 194	122 (194)		GP, PP
Salicylaldehdye	C7H6O2	179, 105, 135, 77, 194	122 (194)	H.O.H.	GP
3-Hydroxy benzaldehyde	C ₇ H ₆ O ₂	179, 105, 135, 77, 194	122 (194)	T-O T-O	GP
2-Hydroxybenzyl alcohol (salicyl alcohol)	C ₇ H ₈ O ₂	73, 253, 179, 268, 147	124 (268)	т. О О-т	GP, PP
4-Hydroxybenzyl alcohol	C ₇ H ₈ O ₂	73, 179, 253, 268, 147	124 (268)	E B	GP, PP
4-Hydroxybenzoic acid (<i>p</i> -salicylic acid)	C ₇ H ₆ O ₃	267, 223, 193, 282, 73	138 (282)		PP [H ₂ O ₂]

- **Table 6.** Highly oxygenated products (O:C > 1.3) identified in benzyl alcohol photooxidation in the presence of NOx, or
- 981 H₂O₂. *: identified with authentic standard. *L*-Tartaric acid and *D*-tartaric acid co-elute. The structure of 4-oxo-D-arabonic
- 982 acid isomer and 2,3,5-Trihydroxy-4-oxopentanal isomer are shown for trihydroxy-oxo-pentanoic acid, and trihydroxy-
- 983 oxo-pentanal, respectively. Four peaks with similar fragments/adducts as pentaric acid were observed.

Nomenclature	Chemical	O/C Ratio	m/z BSTFA Derivative	MW	Proposed Structure
	Formulae	(by wt)	(CI-CH4); (EI)	(MW _{BSTFA})	
Epoxysuccinic acid (2	$C_4H_4O_5$	1.7	187, 261, 73, 277, 173	132	o o
peaks)			73, 173, 261, 129, 143	(276)	H.
2-Hydroxybutanedioic acid*	$C_4H_6O_5$	1.7	233, 335, 73, 307, 351	134	0
(malic acid)			73, 147, 233, 245, 335	(350)	H.O.
					о _н .о
Trihydroxy-oxo-pentanal (5	$C_5H_8O_5$	1.3	73, 275, 203, 349, 393	148	о о ^{,н}
peaks)			147, 73, 349, 233, 259	(364)	H.O
					H, O H
meso-Tartaric acid*	$C_4H_6O_6$	2.0	423, 321, 277, 439, 73	150	0 0 ^H
			73, 147, 292, 219, 423	(438)	H.O.H
					н
L-Tartaric acid*	$C_4H_6O_6$	2.0	423, 321, 277, 439, 73	150	0 0 ^H
			73, 147, 292, 219, 423	(438)	Н. О. Н.
					н
Trihydroxy-oxo-pentanoic	$C_5H_8O_6$	1.6	73, 437, 363, 481, 493	164	он о
acid (8 peaks)			217, 73, 147, 437, 292	(452)	о стран
					он он
D-Arabinonic acid* (Arabic	C5H10O6	1.6	361, 217, 73, 435, 525	166	H. Q.H
acid)			204, 437, 73, 147, 319	(526)	H-O H
					H [,] O O
Pentaric acid* (4 peaks)	$C_5H_8O_7$	1.9	525, 333, 407, 435, 73	180	H _. oo,H
			73, 292, 189, 407, 525	(540)	н.0.0.н
					о _н , оо
				1	

988 Table 7. NACs identified in benzyl alcohol photooxidation in the preser	nce of NOx.
---	-------------

Nomenclature	Chemical Formula Rt (min)	<i>m/z</i> BSTFA Derivative (CH ₄ -CI) (EI)	MW (MW _{bstfa})	Observed in GP; PP [GP/PP ratio]	Proposed Structure
3-nitrobenzyl	C ₇ H ₇ NO ₃	226, 210, 180, 136, 73	153	PP, GP	_OH
alcohol ^a	(25.93)	210, 180, 165, 194, 73	(225)	[1.71]	NO ₂
4-nitrocatechol ^a	C ₆ H ₅ NO ₄	300, 284, 328, 254, 73	155	PP, GP	ОН
	(30.86)	73, 284, 299, 269, 223	(299)	[0.08]	ОН
					NO ₂
2-hydroxy-5-nitro	C ₇ H ₇ NO ₄	314, 298, 268, 342, 73	169	PP, GP	$O_2N_{\rm N}$
benzyl alcohol ^a (4 isomers)	(34.26)	298, 283, 191, 314, 73	(313)	[0.08]	ОН
2-nitro	C ₆ H ₅ NO ₅	388, 372, 416, 428, 73	171	PP	
phloroglucinol ^a (4 isomers) ^b	(35.62)	73, 372, 387, 284, 306	(387)		NO2 OH
3,4-dihydroxy-5-	C ₇ H ₇ NO ₅	388, 372, 416, 428, 73	185	PP	0
nitrobenzyl alcohol (4 isomers) ^c	(38.18)	73, 224, 3876, 401, 356	(401)		HO OH

^a: identified using authentic standards. ^b: Three additional peaks eluted at 33.76, 34.70, 34.76 min with similar mass spectra as those recorded for 2-nitrophloroglucinol standard were detected, and the structure given here is for 2-nitrophloroglucinol. ^c: Three additional peaks eluted at 35.94, 36.60, 38.18 min with similar mass spectra were detected.



Figure 1. Portion of GC-MS total ion chromatogram in EI mode of GP underivatized denuder extract (a) ER-889 (presence of NOx),
(b) ER-892 (absence of NOx), and (c) ER889- (presence of Nox) as BSTFA derivatives.



1013 methanol extract appearance: BnOH/NOx filter (F1); BnOH/H₂O₂ (F2). The same volume of air was sampled on each filter.



Figure 3. Positive EI mass spectra of BSTFA derivatives of selected ring-containing products: benzoic acid, benzene-1,2-diol, 4-1015 hydroxybenzoic acid; and 2-hydroxybenzyl alcohol.



1024Figure 4. Portion (25-34 min) of GC-MS extracted ion chromatograms (CI-CH4) at m/z 423 (green); m/z 437 (blue); and m/z 525 (red)1025merged in one chromatogram (a) BnOH in the presence of NOx; (b) BnOH in the presence of H2O2 and absence of NOx; (c) Chamber1026background. Red and top blue: right axis.



Figure 5. Mass spectra (methane-CI) of ester TMS derivatives of meso-tartaric acid (top right), trihydroxy-oxo-pentanoic acid (middle right), (c) pentaric acid (bottom right), along with the portion of GC-MS extracted ion chromatograms shown in figure 6. Chemical formulae and chemical structure associated with each group is given in the middle column.



Figure 6. Molecular structures of pentaric acid and its isomers (a); mass spectra of TMS derivatives of pentaric acid acquired for smog
 chamber SOA (EI: b, CI: c) and authentic standard (d: EI); Major pentaric acid fragments observed in EI mode (e).



Figure 7. Portion of GC-MS extracted ion chromatograms (EI mode) at *m/z* 210, 165 (IS), 299 (IS), 300, 298, 372, 388 associated with
 nitroaromatic compounds merged in one chromatogram (red) BnOH in the presence of NOx (ER889); (blue) BnOH in the presence of
 H₂O₂ and absence of NOx (ER892).