Long-term total OH reactivity measurements in a boreal forest

Arnaud P. Praplan¹, Toni Tykkä¹, Dean Chen², Michael Boy², Ditte Taipale², Ville Vakkari^{1,3}, Putian Zhou², Tuukka Petäjä², and Heidi Hellén¹

¹Finnish Meteorological Institute, P.O. Box 503, 00101 Helsinki, Finland

²Institute for Atmospheric and Earth System Research/Physics, Faculty of Science, P.O. Box 64, 00014 University of Helsinki, Finland

³Unit for Environmental Sciences and Management, North-West University, ZA-2520 Potchefstroom, South Africa

Correspondence: A. P. Praplan (arnaud.praplan@fmi.fi)

Abstract.

Total hydroxyl radical (OH) reactivity measurements were conducted at the second Station for Measuring Ecosystem-Atmosphere Relations (SMEAR II), a boreal forest site located in Hyytiälä, Finland, from April to July 2016. The measured values were compared with OH reactivity calculated from a combination of data from the routine trace gas measurements (sta-

- 5 tion mast) as well as online and offline analysis with gas chromatography coupled to mass spectrometry (GC-MS) and offline liquid chromatography. Up to 104 compounds, mostly Volatile Organic Compounds (VOCs) and oxidised VOCs, but also inorganic compounds, were included in the analysis, even though the data availability for each compound varied with time. The monthly averaged experimental total OH reactivity was found to be higher in April and May (ca. 17 s⁻¹) than in June and July (7.4 and 12.3 s⁻¹, respectively). The measured values varied much more in spring with high reactivity peaks in late afternoon,
- 10 with values higher than in the summer, in particular when the soil was thawing. Total OH reactivity values generally followed the pattern of mixing ratios due to change of the boundary layer height. The missing reactivity fraction (defined as the different between measured and calculated OH reactivity) was found to be high. Several reasons that can explain the missing reactivity are discussed in detail such as 1) missing measurements due to technical issues, 2) not measuring oxidation compounds of detected biogenic VOCs, 3) missing important reactive compounds or classes of compounds with the available measurements.
- 15 In order to test the second hypothesis, a one-dimensional chemical transport model (SOSAA) has been used to estimate the amount of unmeasured oxidation products and their expected contribution to the reactivity for three different short periods in April, May, and July. However, only a small fraction (3.1–7.3 %) of the missing reactivity can be explained by modelled secondary compounds (mostly oxidised VOCs). These findings indicate that compounds measured but not included in the model as well as unmeasured primary emissions contribute the missing reactivity. In the future, non-hydrocarbon compounds from
- 20 other sources than vegetation (e.g. soil) should be included in OH reactivity studies.

1 Introduction

Terrestrial vegetation is responsible for about 90% of the emissions of Biogenic Volatile Organic Compounds (BVOCs) into the atmosphere (Guenther et al., 1995). Isoprene and monoterpenes are the most abundant BVOCs globally (44 and 17%,

respectively; Guenther et al., 2012). These compounds are very reactive and their lifetimes range from minutes to hours, thus

25 influencing tropospheric chemistry.

Total hydroxyl radical (OH) reactivity measurements can be used as a method to assess our understanding of tropospheric chemistry (Kovacs and Brune, 2001; Williams and Brune, 2015). Many observations of total OH reactivity have been performed in the past few decades and compared to calculated OH reactivity derived from known chemical composition of the atmosphere. While for urban environment the unexplained (or *missing*) reactivity fraction remains low, it is often more than 50 % in forested

30 environments (see the review by Yang et al., 2016). Based on these observations, Ferracci et al. (2018) modelled the global OH reactivity, as well as hypothetical missing chemical sink, which was found to be mostly localized above forested areas and in a few areas with large anthropogenic emissions.

Large fractions of missing reactivity were first observed in a forest in northern Michigan (Di Carlo et al., 2004) and later observed as well in other forested environments (e.g. Hansen et al., 2014; Nakashima et al., 2014; Ramasamy et al., 2016;

- 35 Zannoni et al., 2016). Also in the tropical forest of Borneo up to 70 % of the measured total OH reactivity remained unexplained (Edwards et al., 2013). In addition, Nölscher et al. (2016) identified a large difference of missing OH reactivity between the dry and wet seasons in the Amazon rainforest, with 79 % on average and between 5 to 15 %, respectively. They identified then the forest floor as an important but poorly characterized source of OH reactivity and Bourtsoukidis et al. (2018) recently identified strong sesquiterpene emissions from soil micro-organisms at the same site.
- 40 Also in the boreal forest, which represents approximately one third of the Earth's forested surface (Keenan et al., 2015), a large discrepancy was observed between the total measured OH reactivity and the reactivity calculated from individual compounds present in the forest air (Sinha et al., 2010; Nölscher et al., 2012). Up to 89 % of the measured total OH reactivity could not be explained for periods in which the forest experienced stress conditions (elevated temperature).
- The two main assumptions for the missing reactivity are 1) missing primary emissions and 2) missing oxidation products
 from the emissions. Several studies have been conducted to investigate these hypotheses. Nölscher et al. (2013), for instance, found an increasing missing fraction of Norway spruce (*Picea abies*) emissions from about 15–27 % in spring and early summer and up to 70–84 % in late summer and autumn. In contrast, Kim et al. (2011) found no significant unknown primary BVOC contributing to OH reactivity (for red oak, white pine, beech, and red maple) during their study period in July 2009 in a forest in Michigan. They also found that the missing reactivity from ambient measurement at this site could be explained
 by oxidation products from isoprene. Kaiser et al. (2016) found in an isoprene-dominated forest in Alabama that emissions and their modelled oxidation products reduced the unexplained reactivity to 5–20 % during the day and 20–32 % at night and attribute the missing reactivity to unmeasured primary emissions. Mao et al. (2012) also demonstrated that including modelled oxidation products in OH reactivity calculations reduce the difference with measurements significantly.
- Sinha et al. (2010) and Nölscher et al. (2012) conducted their studies at the second Station for Measuring Ecosystem-55 Atmosphere Relation (SMEAR II; Hari and Kulmala, 2005) in Hyytiälä, Finland, for about three weeks in August 2008 and for about three and a half weeks in July-August 2010, respectively, with the Comparative Reactivity Method (CRM, Sinha et al., 2008). Mogensen et al. (2011) modelled the full year of OH reactivity at SMEAR II for 2008, based on modelled emissions, known chemistry, and environmental conditions. A comparison with results from Sinha et al. (2010) showed that

compounds other than monoterpenes, isoprene, and methane contribute to only about 8 % of the measured OH reactivity. Taking

- 60 all compounds into account, about 61 % of the OH reactivity remained unexplained on average during that period. Mogensen et al. (2015) also compared modelled reactivity at SMEAR II with OH reactivity measurements from Nölscher et al. (2012), using measured trace gases as input but found on average about 65 % of unexplained reactivity, similarly to the previous study. In order to investigate OH reactivity at SMEAR II in more details, in particular its missing fraction and the seasonal variations which are often neglected for summer intensive campaigns, a new implementation of the CRM was developed at the Finnish
- 65 Meteorological Institute (Praplan et al., 2017). It was installed at SMEAR II along with instrumentation to measure VOCs in spring and summer 2016.

2 Methods

2.1 Measurement site

Measurements were conducted at the boreal forest site SMEAR II (Hari and Kulmala, 2005; Ilvesniemi et al., 2009) in Hyytiälä,
Finland (61°51' N, 24°17' E, 181 m above sea level). The site is located in a ca. 60-year old managed conifer forest with modest height variation of the terrain. The stand is dominated by Scots pine (*Pinus sylvestris* L.) homogeneously for about 200 m in all directions, extending to the north for about 1.2 km. Tampere is the largest city near the station about 60 km S-SW.

The instruments were located inside a container in an opening about 115 m from the site mast, from which meteorological data as well as ozone (O_3) , nitrogen oxides (NO_x) , methane (CH_4) , carbon monoxide (CO) and sulfur dioxide (SO_2) concen-

75 trations were retrieved to be included in the analysis. Proton-transfer-reaction mass spectrometer (PTR-MS) measurements of VOCs usually operated at the station mast were not operational during the measurement period and could not be used in this study.

In situ measurements of the total OH reactivity (section ??) and of VOC concentrations (section 2.2) were done at the container, sampling outside air at a height of about 1.5 m (Fig. 1). Station data (from the mast, measurement towers and soil)

80 are open data under Creative Commons 4.0 Attribution licence (CC BY 4.0) and were retrieved from the online SmartSMEAR interface (https://avaa.tdata.fi/web/smart/smear, Junninen et al., 2009).

Temperature and relative humidity (RH) are taken at 4.2 m above ground on the mast; soil properties are an average of five locations throughout the site; and radiation and precipitation data are collected at 18 m height on a nearby tower.

2.2 In-situ measurements of volatile organic compounds

VOCs were measured with two in situ GC-MS. The first GC-MS was used for the measurements of mono- and sesquiterpenes, isoprene, 2-methyl-3-butenol (MBO) and C_{5-10} aldehydes. With this GC-MS air was drawn at the flow rate of 11 min^{-1} through a meter-long fluorinated ethylene propylene (FEP) inlet (i.d. 1/8 inch) and for O₃ removal (Hellén et al., 2012) through a meter-long heated (120°C) stainless steel tube (o.d. 1/8 inch). VOCs were collected from a 40 ml min⁻¹ subsample flow in the cold trap (Carbopack B/Tenax TA) of the thermal desorption unit (TurboMatrix, 650, Perkin-Elmer) connected to a gas



Figure 1. Orthophotograph of the SMEAR II station in Hyytiälä and its surroundings with the marked location of the station mast and the container where the measurements were performed. (Source: Land Survey of Finland Topographic Database 09/2018, CC BY 4.0).

- 90 chromatograph (Clarus 680, Perkin-Elmer) coupled to a mass spectrometer (Clarus SQ 8 T, Perkin-Elmer). A HP-5 column (60m, i.d. 0.25 mm, film thickness 1 µm) was used for separation. The second GC-MS was used for the measurements of C₄₋₈ alcohols and C₂₋₇ volatile organic acids (VOAs). Samples were taken every other hour. The sampling time was 60 min. Samples were analysed in situ with a thermal desorption unit (Unity 2 + Air Server 2, Markes International LTD, Llantrisant, UK) connected to a gas chromatograph (Agilent 7890A, Agilent Technologies, Santa Clara, CA, USA) and a mass spectrom95 eter (Agilent 5975C, Agilent Technologies, Santa Clara, CA, USA). A polyethylene glycol column DB-WAXetr (30-m, i.d. 0.25 mm, a film thickness 0.25 µm) was used for the separation. These methods and measurements have been described in more

detail by Hellén et al. (2017, 2018).

2.3 Offline measurements of volatile organic compounds

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Additional sampling took place between 27 April to 3 May in canisters and through adsorption cartridges (24-hour time resolution) to be analysed by GC-FID (C_{2-6} hydrocarbons) and LC-UV (carbonyls), respectively. During this period, Tenax tube samples were also taken (4-hour time resolution) and analysed later in the laboratory with GC-MS. These results were used as backup to fill in data during interruptions of the online GC-MS measurements. Between 20 and 29 July, additional sampling through adsorption cartridges for offline analysis with LC-UV was performed.

2.4 Mixing Layer Height measurements

- 105 The Mixing Layer Height (MLH) was estimated from measurements with a 1.5 m pulsed Doppler lidar (Halo Photonics Stream Line; Pearson et al., 2009) similar to Hellén et al. (2018). MLH was determined from a combination of turbulent kinetic energy dissipation rate profiles and conical scanning at 30° elevation angle according to the method described in Vakkari et al. (2015). With this method MLH could be determined from 60 m above ground level (a.g.l.) to more than 2000 m a.g.l. at SMEAR II. Periods when MLH was <60 m a.g.l. could be identified although the actual MLH was not determined due to minimum range</p>
- 110 limitations. MLH was not determined for rainy periods. For more detailed specifications of the lidar system and the applied MLH determination method see Hellén et al. (2018).

2.5 Total OH reactivity measurements: the Comparative Reactivity Method (CRM)

The OH reactivity, R_{OH} , is defined as the sum of the concentration of individual compounds X_i multiplied by their respective reaction rate coefficient with respect to OH ($k_{\text{OH}+X_i}$). This can be summarised by the following equation:

$$115 \quad R_{OH} = \sum_{i} [X_i] k_{OH+X_i} \tag{1}$$

The OH reactivity of a compound is the inverse of the OH chemical lifetime due to its reaction with that compound. High OH reactivity values correspond to short lifetimes and long-lived species (such as methane) have a low reactivity.

Our analysis includes up to over 100 individual species from two GC-MS, GC-FID and LC-UV measurements (see sections 2.2 and 2.3). However, not all compounds have been measured at all times (see Fig. 7c). In addition NO_x, O₃, SO₂ and, and CO concentrations were retrieved from the mast of the SMEAR II station, about 115 m away from the sampling position of total OH reactivity and VOCs.

Measurements of total OH reactivity (R_{exp}) have been conducted using the Comparative Reactivity Method (CRM, Sinha et al., 2008; Michoud et al., 2015). Our particular implementation of the method is described in Praplan et al. (2017).

The CRM is based on the monitoring of pyrrole (C₄H₅N) mixed in a 100 ml-reactor with zero air and ambient air, alternatively. The total flow through the reactor is about 465 ml min⁻¹ and the residence time in the reactor estimated about 12–15 s. Pyrrole detection is performed with a gas chromatograph (GC) equipped with a photon ionization detector (PID) every two minutes (Synthec Spectras GC955, Synspec BV, Groningen, The Netherlands). The sensitivity of this detector is independent from the RH of the sample (Fig. 2), but decreased from 1645 a.u./ppbv (data from April to June) to 1290 a.u./ppbv (July data). OH is produced by the photolysis of water (H₂O) in a nitrogen flow (99.9999% N₂) using ultraviolet (UV) radiation and introduced into the CRM instrument reactor. Note that hyroperoxyl radicals (HO₂) are concurrently produced from the reaction of hydrogen (H) with molecular oxygen (O₂). In the zero air mixture, all OH are consumed by pyrrole (C₂ level), while ambient air contains other reactive compounds that compete for OH leading to a higher pyrrole concentration (C₃ level). The instrument switches between measurement of zero air and ambient air every 8 minutes. Stabilization of the conditions takes a couple of minutes and the first data point after each switch is discarded. From the difference between C₂ and C₃ pyrrole levels and taking



Figure 2. Left: Sensitivity of the GC-PID for pyrrole used in the present study. Right: Same day (30 June 2015) sensitivity test for sensitivity of GC-PID for pyrrole in humid and dry conditions.

into account the amount of pyrrole in the reactor in the absence of OH (C₁, typically between 26 and 43 ppb_v), the total OH reactivity R_{eqn} can be derived from the following equation:

$$R_{\rm eqn} = \frac{C_3 - C_2}{C_1 - C_3} \cdot k_p \cdot C_1 \tag{2}$$

with k_p the reaction rate of pyrrole with OH (1.2 · 10⁻¹⁰ cm³ s⁻¹, Atkinson et al., 1985). C₁ is measured by introducing a large concentration of 0.6 % propane (C₃H₈) in nitrogen (N₂) to act as an OH scavenger (Zannoni et al., 2015). Therefore, C₁
takes into account the photolysis of pyrrole due to the UV radiation entering the reactor (8–13 %), which decreases the pyrrole concentration from the total amount of pyrrole injected in the reactor (C₀ level).

Equation (2) assumes that OH levels are identical during C_2 and C_3 measurements. Therefore, variations of RH within the reactor, but also the presence of NO_x and O₃ needs to be taken into account. Therefore C_3 in Eq. (2) results from the following:

$$C_{3} = C_{3,exp} + \Delta C_{3,H_{2}O} + \Delta C_{3,NO_{2}} + \Delta C_{3,O_{3}}$$
(3)

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with $C_{3,exp}$ the measured level of pyrrole in C_3 mode, $\Delta C_{3,H_2O}$ the correction due to different RH in C_2 and C_3 (the difference in RH is 4 % or less 99 % of the time, which corresponds to a change of no more than 5 % for R_{eqn}), and $\Delta C_{3,NO_2}$ and $\Delta C_{3,O_3}$ the corrections due to the presence in the reactor of nitrogen dioxide (NO₂) and O₃, respectively.

The corrections due to the presence of NO₂ and O₃ are discussed in detail in sections 2.5.1 and 2.5.2 Due to the low NO levels at the measurement site, this correction described in earlier publications (e.g. Michoud et al., 2015; Praplan et al., 2017) 150 is not applied nor discussed in the present study.

In addition, because of the dilution of the sampled air with humid nitrogen, the experimental total OH reactivity (R_{exp}) is derived from the following equation:

$$R_{\rm exp} = D \cdot R_{\rm measured} \tag{4}$$

with D the dilution factor (ratio of sampling flow over total flow through the reactor) and R_{measured} the reactivity inside the 155 reactor after applying corrections to $R_{\rm CRM}$ (see section 2.5.3).

Finally, the missing fraction of the total OH reactivity is obtained by comparing R_{exp} with R_{OH} :

$$R_{\rm missing,faction} = \frac{R_{\rm exp} - R_{\rm OH}}{R_{\rm exp}}$$
(5)

2.5.1 Nitrogen dioxide correction factor

Praplan et al. (2017) describe the derivation of this correction in more details. Briefly, the introduction from NO₂ from ambient 160 air in the reactor causes an increase of OH in C_3 mode compared to C_2 (there NO_x is removed by the catalytic converter). This is possibly due to the photolysis of NO_2 to NO and the reaction of NO with HO_2 yielding NO_2 and OH. To derive this correction factor, the change in reactivity due to the injection of NO₂ has been taken into account.

The correction for C_3 ($\Delta C_{3,NO_2}$) for the presence of NO₂ derived experimentally for this study is plotted in Fig. 3 along with the correction from Praplan et al. (2017) for comparison. The correction $\Delta C_{3,NO_2}$ has been applied when it is larger than the standard deviation of C_3 .

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Ozone correction factor 2.5.2

As discussed in Praplan et al. (2017) and by Fuchs et al. (2017) for the CRM system of the Max Planck Institute, the pyrrole signal obtained during analysis of ambient air must be corrected for the presence of O_3 . In the reactor O_3 most probably gets photolysed producing $O(^{1}D)$, which reacts further with H₂O, yielding two OH.

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The corrections $\Delta C_{3,O_3}$ were derived experimentally for various pyr:OH by injecting a known amount of O₃ in the CRM's reactor (Fig. 4, left panel) and then the slope of the linear fit (through the origin) for each pyr:OH (F_{O_3}) was plotted against pyr:OH (Fig. 4, right panel). Based on these data, a linear fit has been derived to calculate F_{O_3} according to pyr:OH and the uncertainty of this correction $(U_{F_{O_3}})$ is 33.7%. When correcting ambient data in this study, the correction for a pyr:OH of 3

 $(F_{O_3} = 0.079)$ has been applied when pyr:OH was higher than 3 due to the lack of experimental data at higher pyr:OH values. Note that a couple of experiments (with pyr:OH 1.27 and 1.05) were performed with additional injection of propane (C_3H_8) as the pyrrole signal would have decreased to zero otherwise and no $\Delta C_{3,O_3}$ could have been determined.

Praplan et al. (2017) used a correction ($\Delta C_{3,O_3}$) independent of pyr:OH as the experimental pyr:OH for the measurements was in a narrow range close to 2. However as pyr:OH varied from 1.0 to 5.3 in this study, a pyr:OH-dependent correction has been derived.



Figure 3. Correction of C_3 ($\Delta C_{3,NO_2}$) as a function of nitrogen dioxide in the reactor (NO_{2,reactor}). Error bars represent standard deviations and the black line the overall fit to derive the correction factor $\Delta C_{3,NO_2}$. For comparison, the correction from Praplan et al. (2017) is depicted as a gray dashed line.

180 The correction $\Delta C_{3,O_3}$ is then derived from the following equation:

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$$\Delta C_{3,O_3} = F_{O_3}[O_3] = (0.022 \cdot (\text{pyr}:\text{OH}) + 0.013) \cdot [O_3]$$
(6)

As observed in Praplan et al. (2017), inhomogeneity of the air composition at the sampling site can affect the comparison between experimental total OH reactivity and calculated reactivity from known composition. It can for instance be directly affected by meteorology or changes in concentrations between the various sampling locations due to local emissions during low mixing periods (see Liebmann et al., 2018). As VOCs in this study were sampled at the same location than the total OH reactivity, the effect of inhomogeneity of the air composition is minimized. However, the ozone mixing ratio used to derive the ozone correction (described in section 2.5.2) is retrieved from the station mast (115 m away) and at a height of 4.2 m. It is very likely that emissions from soil and understorey vegetation (or from standing water close to the OH reactivity sampling location) would further deplete the ozone close to the ground, leading to an overestimation of the correction. Under some circumstances, such as when there is a strong O_3 gradient below canopy (Chen et al., 2018), the correction might be overestimated.

For instance, on 29 and 30 April total OH reactivity peaks close to 100 s^{-1} in the afternoon are followed by O₃ concentration drops below canopy (Fig. 5, see also Chen et al. (2018)). While the high reactivity peaks themselves are likely not affected by an overestimation of the correction, the period following them (night-time) might be slightly overestimated due to the sampling of O₃ further away and higher above ground. This effect is difficult to take into account in retrospect. The concentration of O₃



Figure 4. Left: Correction of C_3 ($\Delta C_{3,O_3}$) as a function of ozone in the reactor ($O_{3,reactor}$). Right: O_3 correction factor (F_{O_3}) as a function of pyr:OH.

195 should have been measured immediately next to the CRM system. Similar conditions were observed during nights between 11 and 16 May and to some extent in July (without reaching such high total OH reactivity values as in spring). This effect on the inhomogeneity of the forest air composition might affect total OH reactivity measurements and in turn partly explain some of the missing fraction.



Figure 5. One-hour averages of total measured OH reactivity, R_{exp} and ozone mixing ratios at 4.2 and 125.0 m above ground. Mixing Layer Height (MLH) is shown as a gray shadow. Note that the detection limit for MLH is 60 m and values below this limit are displayed at 30 m (and zeros denote gaps in the data).

2.5.3 First order correction factor

200 Sinha et al. (2008) used a two-equation model to correct for the deviation from pseudo-first-order kinetics ([Pyr]»[OH]). Michoud et al. (2015) used more detailed modelling taking into account OH recycling reactions, but could not match the model results with their experimental data. For this reason, Michoud et al. (2015) favoured the experimental approach to correct the reactivity data. Nevertheless, the experimental approach also has drawbacks. For instance, impurities from standards and changes over time (ageing) might alter its reactivity. Also it is based on calibrations using one compound at the time, which do not represent complex ambient mixtures of reactive gases.

Nevertheless, reactivity calibrations were performed for the present study with a 10 ppm_v C_3H_8 standard as well as with an in-house made gas mixture containing α -pinene with small impurities from aromatic compounds. The concentrations of the C_3H_8 and in-house α -pinene standards were checked periodically by taking adsorbent tube samples and analysing them by GC-MS. At the same time impurities (4.7–17% of the reactivity) could be measured and taken into account.

- 210 The comparison between OH reactivity expected from the standard (R_{std}) and the measured OH reactivity $(R_{measured})$ is presented in Fig. 6. Note that the data has been corrected for deviation from pseudo-first-order kinetics similarly to the work of Sinha et al. (2008) by using numerical simulations and fitting the relationship between R_{eqn} and R_{true} for various pyr:OH ratios with equations of the form $R_{true} = F_1 \cdot R_{eqn}^{F_2} + F_3$, so that F_1 , F_2 , and F_3 are pyr:OH-dependent coefficients (see Appendix A for additional details).
- 215 The calibration for C_3H_8 is consistent with the one from Sinha et al. (2008). Due to the high reactivity of α -pinene, the calibration consistently underestimate the expected reactivity and because monoterpenes constitute the most important class of compounds in the boreal forest, this needs to be taken into account by applying the overall correction for α -pinene to the ambient data in this study. The reactivity measured in the reactor $R_{measured}$ then is derived from the following equation:

 $R_{\text{measured}} = (R_{\text{CRM}} + 0.449)/0.497$

(7)



Figure 6. Comparison between measured OH reactivity for C_3H_8 and α -pinene standards with the expected OH reactivity.

220 2.6 The model to Simulate the concentrations of Organic vapours, Sulphuric Acid and Aerosols (SOSAA)

In this study we applied the model to Simulate the concentrations of Organic vapours, Sulphuric Acid and Aerosols (SOSAA) to simulate the OH reactivity at the SMEAR II station for selected days in April, May, and July 2016. SOSAA is a one-dimensional chemical transport model comprised of boundary layer meteorology, biogenic emission of VOCs, gas-phase chemistry, aerosol dynamics and gas dry deposition (e.g. Boy et al., 2011; Zhou et al., 2014) and has been previously used to simulate OH reactivity at this site (Mogensen et al., 2011, 2015).

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The boundary layer meteorology was derived from SCAlar DIStribution (SCADIS; Sogachev et al., 2002), as described in Boy et al. (2011). The biogenic emission module was deactivated because in situ measurements were used to provide input concentrations. Biogenic compounds were set to the measured values up to 18 m (canopy height), while aromatic compounds were set to the measured values at all heights. Measured inorganic gas concentrations at SMEAR II were used as input. The gas-phase chemistry was created using the Kinetic PreProcessor (KPP; Damian et al., 2002). The chemical reaction equations used in this study were selected from the Master Chemical Mechanism v3.3.1 (MCMv3.3.1 Jenkin et al., 1997; Saunders et al., 2003; Bloss et al., 2005; Jenkin et al., 2012, 2015). The chemistry scheme included more than 15000 reactions, and a total of 3525 chemical species representing the complete reaction paths for isoprene, α -pinene, β -pinene, limonene, β -caryophyllene, methane, 2-methyl-3-buten-2-ol (MBO), benzene, toluene, styrene, ethylbenzene, 1,2-dimethylbenzene, 1,3-dimethylbenzene,

- 235 1,4-dimethylbenzene, 1,2,3-trimethylbenzene, 1,2,4-trimethylbenzene, 1,3,5-trimethylbenzene, 1-ethyl-2-methylbenzene, 1ethyl-3-methylbenzene, 1-ethyl-4-methylbenzene, heptane, octane, nonane, butanal, pentanal, methacrolein and relevant inorganic reactions. First order reactions between OH, O₃, and NO₃ with the following monoterpenes were also included in the chemistry: Δ^3 -carene, myrcene, camphene and 1,8-cineole. Likewise, first order reactions between OH, O₃, NO₃ and β farnesene were included. The photochemistry has been improved by calculating the photodissociation constants more precisely
- 240 using data from Atkinson et al. (1992), as described in Mogensen et al. (2011). The OH reactivity has been calculated similarly as in Mogensen et al. (2011, 2015). The condensation sinks for sulfuric acid and nitric acid, based on Differential Mobility Particle Sizer (DMPS) and Aerodynamic Particle Sizer (APS) data from SMEAR II, were included (Boy et al., 2003). Since sulfuric acid and nitric acid make up most of the condensation sinks, sinks of VOCs into the particle phase are not taken into account, thereby the aerosol module is turned off.
- The model runs in the present study include the dry deposition module implemented in SOSAA by Zhou et al. (2017a) and extended in Zhou et al. (2017b). The latter describes the explicit simulation of the loss of every compound in the model by dry deposition inside the canopy for all height levels and shows that the sink by dry deposition inside the canopy is comparable to the chemical production for several oxidised VOCs (e.g. pinic acid or BCSOZOH, a reaction product of β -caryophyllene).

3 Results and discussion

250 3.1 Overview

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An overview of the measured total OH reactivity together with the calculated OH reactivity from up to 104 compounds, depending on data availability, as well as selected ancillary data, such as environmental conditions (air and surface soil temperatures as well as surface soil water content), and contributions from different compounds and groups of compounds are presented in Fig. 7. The following sections are discussing in details various aspects of the results such as a) seasonality, b) diurnal variations, and c) missing reactivity. Nevertheless, from this overview, the following observations can be made:

- The range of measured total OH reactivity values is similar to previous studies at the same site in August 2008 and July-August 2010 (Sinha et al., 2010; Nölscher et al., 2012), with notably higher values in the spring.
- These high total OH reactivity peaks in the spring (with values higher than at the end of July) seems to be associated with changes in the soil water content resulting from soil thawing.
- The calculated OH reactivity from measured compounds is in general lower than the measured total OH reactivity (also for periods with a large number of compounds included in the analysis), leading to a large fraction of *missing* reactivity (see section 3.4).

- Inorganic compounds (CH₄, CO, O₃, and NO₂) form an important fraction of the calculated OH reactivity.



as well as surface soil water content), c) Pyr:OH in the CRM reactor, d) data availability from the different instrumentation/sources, e) fraction of experimental total OH reactivity, and f) fraction of calculated OH reactivity. The periods shaded in gray in panels (a) to (d) represent the periods investigated with SOSAA (see Figure 7. a) Experimental total OH reactivity $R_{\rm exp}$ (1-h average) and calculated OH reactivity $R_{\rm OH}$, b) environmental conditions (air and surface soil temperatures, sect 3.4).

3.2 Total OH reactivity

Table 1. Monthly means and standard deviations (std.) of experimental total OH reactivity (R_{exp}), the missing OH reactivity fraction ($R_{missing,fraction}$), monoterpene and sesquiterpene mixing ratios ([MT] and [SQT], respectively), Photosynthetically Active Radiation (PAR), precipitation (Precip), relative humidity (RH), air temperature (T), surface soil temperature ($T_{soil,humus}$), surface soil water content ($w_{soil,humus}$), and Mixing Layer Height (MLH). Coefficients a and b from linear regressions between the weekly means of these variables and weekly averaged R_{exp} and the corresponding coefficients of determination (r^2). n_{days} indicates the number of days with measurements. n denotes the amount of R_{exp} observations. Note that all other means (except MLH) have been derived for the same measurement period as R_{exp} . n_{MLH} indicates the amount of observations with overlaping R_{exp} and MLH measurements.

	April	May	June	July	Linear re	egressions ($(aR_{\exp} + b)$
	mean (std.)	mean (std.)	mean (std.)	mean (std.)	a	b	r^2
$n_{ m days}$	17	26	16	12			
n	1032	1854	1416	952			
$R_{\mathrm{exp}} (\mathrm{s}^{-1})$	17.1 (20.4)	17.5 (12.2)	7.4 (2.6)	12.3 (5.9)			
$R_{\rm missing, fraction}$	0.80 (0.15)	0.85 (0.15)	0.85 (0.05)	0.74 (0.16)	0.004	0.768	0.16
[MT] (ppt _v)	99.6 (186.5)	233.8 (493.4)	83.6 (408.2)	564.5 (508.6)	8.5	86.5	0.10
$[SQT] (ppt_v)$	0.092 (0.319)	2.15 (2.92)	1.12 (3.78)	23.1 (23.7)	-0.04	5.31	0.001
PAR (μ mol m ⁻² s ⁻¹)	177.3 (277.1)	313.0 (420.0)	490.9 (521.2)	359.0 (422.1)	-4.8	445.7	0.10
Precip (mm)	0.13 (0.10)	0.12 (0.14)	0.12 (0.18)	0.10 (0.00)	0.0002	0.1158	0.004
RH (%)	83.4 (17.8)	67.8 (22.4)	58.0 (21.1)	79.0 (16.0)	0.6	60.2	0.10
<i>T</i> (°C)	3.7 (3.9)	11.8 (4.6)	12.2 (5.6)	18.0 (3.5)	-0.02	11.7	0.0005
$T_{ m soil,humus}$ (°C)	1.6 (1.2)	8.2 (2.4)	9.9 (2.1)	15.2 (1.4)	-0.06	9.08	0.009
$w_{\rm soil,humus}~({ m m}^3~{ m m}^{-3})$	0.38 (0.05)	0.32 (0.03)	0.24 (0.03)	0.28 (0.03)	0.004	0.247	0.33
$n_{ m MLH}$	1016	1833	1291	945			
MLH (m)	196.5 (296.0)	335.9 (553.6)	573.0 (679.3)	309.3 (443.5)	-8.0	538.3	0.13

- Keeping in mind that the experimental data have not always been acquired continuously, the total experimental OH reactivity (R_{exp}) monthly mean was high in April and May (about 17 s^{-1}) compared to June (7.4 s⁻¹) and July (12.3 s⁻¹), due to few very high values at night time (Table 1). Consequently, no strong correlation could be found between R_{exp} and other variables looking at weekly means. The highest coefficient of determination (r^2) was obtained for the correlation with $w_{soil,humus}$ (0.33), which indicates that soil moisture might be an important driver for the high reactivity values measured in spring. The
- 270 highest reactivity peaks happened when the surface soil water content was the highest as the surface soil temperature started to increase above 1.5 °C, indicating thawing of the soil, a possible source of OH reactive compounds. Forest floor emissions of monoterpenes are known to be high in spring after snow has melted (Hellén et al., 2006; Aaltonen et al., 2011; Mäki et al., 2017) and VOC emission bursts have been observed after wetting events (e.g. Rossabi et al., 2018). There has also been some indication that thawing snow/soil could be a source of volatile organic amines (Hemmilä et al., 2018). In the present study, the

- soil was snow-free already on 8 April, but a short snowfall episode happened later with 5 cm of snow measured on the morning 275 of 25 April (which were gone on the next day). This episode happens just before the first OH reaction peak (at about $47 \, \text{s}^{-1}$). but this single occurrence is too little information to conclude of the role of snow in the large OH reactivity values observed and it might well be due to a combination of factors (including snowfall and immediate melting). These results deviate however from the conclusions of Nölscher et al. (2016), which suggested that a wet (and cold) soil in the Amazon rainforest acts as 280 sinks for reactive compounds.

The data for July cover days that were cloudier and more humid (both air and soil) but warmer than the period covered by the data in June leading to higher total OH reactivity. Monthly means of ambient concentrations of locally emitted terpenoids had a weak correlation ($r^2 = 0.43$) with temperature (see also Hellén et al., 2018), which is not reflected in the correlation of total measured OH reactivity with temperature, as observed earlier (e.g. Nakashima et al., 2014; Ramasamy et al., 2016). 285 However, these studies were performed during summer, which highlights the different regimes governing OH reactivity in various seasons and how most likely other (unknown) compounds in addition to terpenes contribute to OH reactivity during spring. In other words, while conditions that favour high OH reactivity values seem to favor BVOC (terpene) emissions in the summer as well, OH reactivity is driven by other parameters in spring.

It should be noted, though, that the use of a correction factor based on α -pinene throughout the measurement period even though the air composition varied might lead to an overestimation of the measured total OH reactivity. However, average mixing 290 ratios of monoterpenes were similar in April and June (99.6 pptv_y and 83.6 pptv_y, respectively), so that relative differences in measured total OH reactivity cannot be explained this way. This further indicates that non-terpene compounds that were not measured in the spring might have contributed to the total OH reactivity.

3.3 **Diurnal variations**

The calculated OH reactivity of various groups of compounds shows different diurnal patterns, which vary with the season as 295 well. Their normalized values are depicted in Fig. 8 (second to fourth row), separated by month (April to July in columns), together with the normalized diurnal patterns of R_{exp} and its missing fraction and temperature difference between measurements at 4.2 m and 125.0 m above ground as a proxy for mixing layer height (top row). Compounds that had a 24-hour sampling time were removed from this analysis. Sinha et al. (2010) did not measure a clear OH reactivity diurnal pattern during their two-week measurement period and the modelling of the OH reactivity also showed no diurnal pattern (Mogensen et al., 2011). 300 However, Mogensen et al. (2015) modelled a weak diurnal pattern with a maximum at night, mostly due to improvements in the meteorological scheme. The observations in the present study, even though at higher OH reactivity levels show this pattern from May to July. Nölscher et al. (2012), for measurements roughly at the same time of the year, identified a similar diurnal pattern with maximum at night during the identified stress period. For normal boreal forest conditions, they measured large 305 variations in the afternoon reactivity, sometimes leading to a maximum, which they associated with long-range transport. In the present study, afternoon reactivity maxima were dominating April's diurnal pattern.

When the total measured OH reactivity hourly average is at a minimum during the day and a maximum at night (May to July), it follows the pattern of BVOCs concentrations (and calculated OH reactivity) due to the low mixing layer height and



Figure 8. Normalized monthly averaged diurnal variations of experimental OH reactivity R_{exp} and the missing fraction as well as temperature gradient between 4.2 and 125.0 m above ground as a proxy for mixing layer height (top row), and calculated OH reactivity separated by group of compounds (second to fourth row).

despite slightly lower emissions due to the lower temperatures at night (Hellén et al., 2018). The hourly average of missing
reactivity fraction remained consistently high (between 65 and 92 %), similar to values from Nölscher et al. (2012) and despite the inclusion of more compounds in our analysis (see section 3.4 for a detailed discussion).

While the OH reactivity daily patterns from monoterpenoids and MBO had a minimum during the day for all months, other groups of compounds showed this reactivity pattern only for some periods. Isoprene showed this pattern except in July, where the light-induced emissions during the day were dominating. Sesquiterpenes, other carbonyls and NO₂ showed a similar pattern with daytime minima from May to July, while C₇₋₁₀ alkenes, aromatics, C₃₋₁₀ aldehydes, and methacrolein showed a pattern with daytime minimum only in May and June. Alcohols exhibit an OH reactivity pattern with a maximum in the morning (9-11 a.m.). The absolute OH reactivity of alcohols is low and dominated by 1-butanol, which is used in aerosol measuring devices at the site. It is not clear what causes the diurnal pattern, but SO₂ reactivity had a similar pattern in April and May, and NO_x had such a pattern in April, when the photochemistry is not yet very strong.

320 Overall, from May to July the total OH reactivity exhibits a minimum during the day and a maximum at night, following the OH reactivity pattern for biogenic compounds (except for isoprene in July, which is present in low concentrations in this pine forest, and has a maximum in the afternoon then). In April, the total OH reactivity has a maximum in the afternoon, and sesquiterpenes, even though present in low concentrations show a similar reactivity pattern. Mäki et al. (2019) found high levels of sesquiterpenes from soil emissions at the same site in spring. This is an additional indication that unknown primary emissions (in particular from soil) could drive the reactivity during that time of the year.

3.4 Missing OH reactivity

The comparison between the calculated and measured OH reactivity is challenging as the calculated values are derived from a number of compounds that varies because of the availability of the measurements (Fig. 7d). Some periods include only a few inorganic compounds from the station mast while other periods include a large amount of (O)VOCs analysed by the GC-MSs.
The contribution to the known reactivity is shown in Fig. 7f. It is also good to keep in mind that part of the missing reactivity can be explained by measurement uncertainties and potential overestimation due to applied correction factors. As the data in this study has been uniformly corrected based on *α*-pinene calibrations, it is likely that the obtained values are an upper limit for the reactivity considering that *α*-pinene (and monoterpenes in general) are not dominating the air composition and reactivity at the site for the whole measurement period. It should also be remembered that because of technical problems, PTR-MS data (VOC data) from the station mast is unavailable for our measurement period. Some compounds such as acetaldehyde were measured during two short periods with offline 24-hour sampling methods. In late July, acetaldehyde contributed on average 0.13 s⁻¹ to the OH reactivity, which can be a small but significant fraction for low OH reactivity values. This is likely the case for other compounds that were not measured at all in the present study such as formaldehyde, acetonitrile, or methanol, to name a few.

340 in the present work, the impossibility to include aforementioned compounds in the analysis explain partly why missing OH reactivity fractions remain high. Therefore, even with the maximum number of compounds used to calculate OH reactivity (late April/early May) a large fraction of the measured total OH reactivity remain unexplained (*missing* reactivity, Fig. 7e).

However, this period coincided also with high reactivity peaks observed likely due to soil thawing as mentioned previously. Only sesquiterpenes peaked at the same time as the total OH reactivity, but their concentrations are still low, which is why we

In addition, this also makes the comparison with previous studies difficult. Despite the higher number of compounds included

- 345 mentioned amines and non-terpene BVOCs as potential classes of compounds contributing to the observed total OH reactivity. Kumar et al. (2018) identified various non-hydrocarbon classes of compounds associated with biomass burning that potentially contribute to OH reactivity. However, even if long-range transported biomass burning emissions are observed occasionally at the measurement site of this study Leino et al. (2014), no significant increase of CO concentrations (above 250 ppb_v as in Leino et al. (2014)) were observed during the measurement period. Only between 23 and 26 July, concentrations of 150 ppb_v (slightly
- above the average background levels of 100 ppb_v) were detected. Nevertheless, these classes of compounds could potentially be emitted by local sources of a different kind.

As it has been shown for forests dominated by isoprene emitters (Kim et al., 2011; Kaiser et al., 2016), oxidation products from BVOCs might contribute significantly to the missing OH reactivity. As oxidation products of monoterpenes and sesquiterpenes are neither measured routinely nor were they monitored for this study, the SOSAA model was used (see Sec-

355 tion 2.6) using measured trace gases and meteorological conditions as inputs in oder to calculate the potential contribution of terpenes oxidation products to missing OH reactivity. Three periods of two to three days for the months of April, May and July were simulated. The results for the inclusion of modelled oxidation compounds in the analysis are presented in Fig. 9. These compounds labelled modelled OVOCs are mostly peroxides, alcohols, and carbonyl compounds due to the generally low NO_x levels at the site. Modelled inorganics, whose contributions is negligible, regroup molecular hydrogen (H₂), hydrogen peroxide (H₂O₂), nitrous acid (HONO), peroxynitric acid (HO₂NO₂), nitric acid (HNO₃), and the nitrate radical (NO₃).

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While the trend of $R_{\text{OH,model}}$ follows qualitatively the general trend of R_{exp} , $R_{\text{OH,model}}$ underestimates R_{exp} , especially at night. Total OH reactivity values are in general lower during the day and they are closer to $R_{\text{OH,model}}$ values. In April, the high peaks in the late afternoon of 29 and 30 April indicate missing primary emissions, which also contribute (or their oxidation products) to the missing reactivity in the following nights.

Retrieving the additional reactivity from these modelled compounds that were not included in R_{OH} reduced the missing reactivity by only a small fraction (about 7.3 % for the studied period in July and less for the other periods) as seen in Fig. 10. A detailed breakdown of the individual compounds contributing to the reactivity and their mixing ratios can be found in the Appendix B.

Most of the missing reactivity could be then due to oxidation products that are not included in the model from measured precursors such as Δ^3 -carene, myrcene, camphene, 1,8-cineol, β -farnesene, or unidentified sesquiterpenes (in contrast with the well-studied isoprene chemistry scheme), but the contribution to the OH reactivity from these precursors is small due to their low atmospheric concentrations, so that the contribution from their oxidation products is also expected to be small (Hellén et al., 2018). As mentioned earlier, missing primary emissions also contribute to the missing reactivity, more so in spring than in summer.

Amines released from soil, as mentioned previously, are a potential class of compounds that could contribute to OH reactivity. Kumar et al. (2018) identified various non-hydrocarbon classes of compounds associated with biomass burning that potentially contribute to OH reactivity. However, even if long-range transported biomass burning emissions are observed occasionally at the measurement site of this study Leino et al. (2014), no significant increase of CO concentrations (above 250 ppb_v as in Leino et al. (2014)) were observed during the measurement period. Only between 23 and 26 July, concentrations of 150 ppb_v (slightly above the average background levels of 100 ppb_v) were detected.

It is also good to keep in mind that part of the missing reactivity can be explained by measurement uncertainties and potential overestimation due to applied correction factors. As the data in this study has been uniformly corrected based on α -pinene calibrations, it is likely that the obtained values are an upper limit for the reactivity considering that α -pinene (and monoterpenes in general) are not dominating the air composition and reactivity at the site all the time.

- A previous study by Mogensen et al. (2011) modelled the OH reactivity at the SMEAR II site for the year 2008, using modelled emissions, and estimated the OH reactivity to be about $2-3 \text{ s}^{-1}$ between April and July. This is lower than the measured averages from the present and previous studies and lower than the night-time modelled values in July from the present study. Mogensen et al. (2011) report that secondary organics, β -caryophyllene, farnesene, and MBO represent 8% of the total OH reactivity, which represent the same magnitude as the results from this study. Mogensen et al. (2015) modelled the
- 390 OH reactivity at the same site for July and August 2010 with the same methodology (including minor model improvements) and obtained values between 2.7 and 3.2 s^{-1} . The higher modelled values in our study indicates that modelled emissions lead to lower monoterpene concentrations than measured concentrations.



Figure 9. Measured total OH reactivity (R_{exp}), calculated OH reactivity from measured compounds (R_{OH}), calculated OH reactivity including measured and modelled compounds ($R_{\rm OH,model}$) (1 hour averages, top panels) and normalized contributions to $R_{\rm exp}$ for various compounds and group of compounds (bottom panels) for the three periods investigated with SOSAA (see main text for details).

Our results are not entirely in line with other studies that showed reductions of the missing reactivity by constraining VOC concentrations to model their oxidation products (e.g. Mao et al., 2012; Edwards et al., 2013; Kaiser et al., 2016), as the reduction observed remains small in this study. This approach still leaves a large unexplained fraction of OH reactivity. This is a strong indication that on one hand non-terpenoid compounds contribute to the total OH reactivity and that on the other hand more compounds have to be included in the chemical model.

Finally, heterogeneous loss of OH to particles might be a contribution to missing OH reactivity, but this process is poorly quantified (Donahue et al., 2012). Due to the low sampling flow and long FEP sampling line to the CRM instrument, it is unlikely that particles will reach the reactor. Additionally, we could not find any correlation between ambient particle numbers 400 and either total measured OH reactivity or its missing fraction.

As a side note, total OH reactivity measurements were unfortunately not available in the autumn, but Liebmann et al. (2018) who measured nitrate radical (NO₃) reactivity at the same site made found about 30 % of unexplained NO₃ reactivity at night and about 60 % during daytime. Mogensen et al. (2015) modelled NO₃ reactivity at the site and found a maximum in the early morning, while the measurements from Liebmann et al. (2018) showed a maximum at night. The modelled NO_3 reactivity values were similar to the measured ones without strong temperature inversion at night, while higher measured values were

Hellén et al. (2018) showed that the balance between the emissions of VOCs and the production of oxidation compounds and the sinks vary with the season, leading to different diurnal profiles for compounds such as isoprene, C_{7-10} aldehydes, and nopinone. This can also be observed in terms of OH reactivity in the present study (see section 3.3). 410

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recorded for nights with strong temperature inversion.



Figure 10. Contributions of various compounds and groups of compounds to the measured total OH reactivity (R_{exp}). For clarity, labels for fractions smaller than 2.0 % have been omitted.

4 Conclusions

Total OH reactivity is not a simple function of a few variables. It includes many complex processes involving sources and sinks that can change dramatically depending on the environmental conditions and the time of the year. Measurement uncertainties and data availability for comparison between measured total OH reactivity values and calculated values also represent

415 a challenge when interpreting results.

In the present study total OH reactivity measurements were performed at a Finnish boreal forest research site (SMEAR II). The monthly averaged experimental total OH reactivity were high in April and May (about 17 s^{-1}) due to some very large afternoon reactivity peaks captured when the soil was thawing. The low sampling height and the peaking of sesquiterpene emissions at the same time than OH reactivity in April, which are known to be emitted from soil, indicate that the forest floor is a potential important but overlooked source of reactive compounds. The total OH reactivity diurnal pattern from May to July follows the one of biogenic compounds with high values during the night due to the low mixing height, even though emissions are lower at night.

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A suite of online and offline (O)VOCs measurements was used to calculate the known fraction of OH reactivity to compare it to the total OH reactivity measured. The missing fraction of the OH reactivity remained high for the measurement period. This can be due to various reasons. As the data availability of (O)VOCs varies, the comparison between experimental and calculated OH reactivity is difficult but three different explanations can lead to high missing (unexplained) OH reactivity: 1) simply the lack of measurements, 2) not measuring oxidation products (only their precursors), and 3) not measuring the right class of compounds. We showed that compounds not included (or only partially included) in the analysis due to the unavailability

of measurements (e.g. due to technical problems), such as acetaldehyde, might contribute a small but significant fraction to

the total OH reactivity, in particular for low reactivity values. Using one-dimensional transport model to estimate oxidation products concentrations from measured precursor concentrations for three short periods of two to three days in various months (with most (O)VOC data availability) it is demonstrated that only a small fraction (up to ca. 7.3%) of the missing reactivity can be explained by these oxidation products. On one hand, this is due to the absence in the model of degradation scheme for detected compounds in the ambient air (e.g. Δ³-carene, β-farnesene), but on the other hand it is also possible that non-hydrocarbon compounds contribute to the OH reactivity as well.

More measurements of oxidised compounds and identification of non-terpene reactive compounds from emissions also from other sources than vegetation (e.g. soil) are required to better understand the reactivity and local atmospheric chemistry in the forest air in general, in particular during winter, spring, and autumn, when the forest air chemistry is not dominated by emissions from the vegetation.

440 Author contributions.

A. P. Praplan conducted total OH reactivity measurements, offline sampling, LC-UV analysis, performed data analysis, and led the writing of the manuscript. H. Hellén designed the measurement campaign, conducted GC-MS measurements and data analysis, and commented the manuscript. T. Tykkä assisted GC-MS measurements and data analysis and commented on the manuscript. V. Vakkari provided mixing layer height data, their description in the method part and commented the manuscript. D. Chen, M. Boy and P. Zhou performed model runs with the help of D. Taipale and all commented the manuscript.

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Figure A1. Numerical simulations of R_{true} as a function of R_{eqn} for various pyr:OH values and their corresponding fit curves of the form $R_{\text{true}} = F_1 \cdot R_{\text{eqn}}^{F_2} + F_3$ (upper plot). Fit coefficients F_1 , F_2 , and F_3 as function of pyr:OH and corresponding exponential fit curves (lower panels).

Appendix B

Table B1: Averages of individual compounds mixing ratios [ppt_v] and calculated OH reactivity, $R_{\rm OH}$ [s⁻¹], and group $R_{\rm OH}$ for the three periods studied with SOSAA. 'n.d.' means 'not detected' and 'n.m.' means 'not measured'.

		29.	– 30 April			22	– 25 May			24	– 26 July	
	Mixing ra	tio [pptv]	$R_{\rm OH}$	[s ⁻¹]	Mixing ra	ttio [pptv]	$R_{\rm OH}$	[s ⁻¹]	Mixing 1	atio [pptv]	$R_{\rm OH}$	[s ⁻¹]
	mean	(std)	mean	(std)	mean	(std)	mean	(std)	mean	(std)	mean	(std)
Alkanes			0.065	(0.013)			0.00035	(0.00024)			0.00035	(0.00022)
ethane	2774	(78)	0.0115	(0.0003)	n.m.	-		(-)	n.m.	(-)		(-)
propane	575	(78)	0.014	(0.002)	n.m.	(-)		(-)	n.m.	-		(-)
<i>n</i> -butane	138	(48)	0.0077	(0.0026)	n.m.	(-)	,	(-)	n.m.	-)	'	(-)
2-methylpropane	88	(31)	0.0044	(0.0015)	n.m.	(-)		(-)	n.m.	-)		(-)
<i>n</i> -pentane	61	(13)	0.0055	(0.0012)	n.m.	(-)		(-)	n.m.	-)		(-)
2-methylbutane	112	(16)	0.010	(0.002)	n.m.	(-)		(-)	n.m.	-)		(-)
<i>n</i> -hexane	23	(2)	0.0028	(0.000)	n.m.	(-)		(-)	n.m.	-)		(-)
2-methylpentane	25	6	0.0034	(0.0010)	n.m.	(-)		(-)	n.m.	(-)		(-)
<i>n</i> -heptane	5.6	(1.9)	0.0020	(0.0007)	0.54	(0.35)	0.000058	(0.000109)	0.25	(0.21)	0.000011	(0.000038)
<i>n</i> -octane	7.3	(2.5)	0.0015	(0.0005)	1.0	(0.4)	0.00019	(0.0000)	1.9	(0.4)	0.00032	(0.00016)
<i>n</i> -nonane	3.3	(1.2)	0.00081	(0.00030)	0.43	(0.14)	0.00010	(0.00004)	0.13	(0.07)	0.000024	(0.00019)
<i>n</i> -decane	2.3	(1.1)	0.00064	(0.00031)	n.d.	(-)	ı	(-)	n.d.	(-)	ı	(-)
Alkenes			0.38	(0.04)				(-)				(-)
ethene	354	(26)	0.076	(0000)	n.m.	-		(-)	n.m.	(-)		(-)
propene	135	(9)	0.11	(0.01)	n.m.	(-)		(-)	n.m.	(-)		(-)
1-butene	47	(4)	0.042	(0.004)	n.m.	(-)		(-)	n.m.	-)		(-)
trans-2-butene	46	(8)	0.086	(0.016)	n.m.	(-)		(-)	n.m.	-)		(-)
cis-2-butene	27	(4)	0.043	(0.007)	n.m.	(-)	,	(-)	n.m.	-		(-)
1,3-butadiene	n.d.	-	ı	(-)	n.m.	(-)	'	-	n.m.	(-)	'	(-)
1-pentene	35	(2)	0.025	(0.005)	n.m.	(-)		(-)	n.m.	-		(-)
trans-2-pentene	n.d.	(-)		(-)	n.m.	(-)		(-)	n.m.	(-)		(-)
ethyne	259	(21)	0.0051	(0.0004)	n.m.	-		(-)	n.m.	(-)		(-)
Aromatics			0.038	(0.014)			0.012	(0.009)			0.021	(0.017)
benzene	93	(16)	0.0028	(0.0005)	12	(3)	0.00035	(0.00010)	15	(4)	0.00036	(0.00020)
toluene	37	(6)	0.0058	(0.0014)	32	(10)	0.0046	(0.0017)	22	(9)	0.0026	(0.0014)
ethylbenzene	10	(2)	0.0018	(0.0004)	2.7	(0.8)	0.00047	(0.00015)	6.5	(1.7)	0.00093	(0.00050)
<i>p/m</i> -xylene	14	(2)	0.0067	(0.0033)	3.1	(2.2)	0.0014	(0.0011)	11	(2)	0.0041	(0.0020)
o-xylene	5.8	(1.9)	0.0020	(0.0007)	1.0	(0.0)	0.00017	(0.00029)	2.3	(1.0)	0.00063	(0.00041)
styrene	7.6	(3.2)	0.012	(0.005)	1.8	(1.4)	0.0025	(0.0020)	8.8	(8.2)	0.010	(0.011)
2-ethyltoluene	1.2	(0.3)	0.00036	(0.00010)	0.42	(0.23)	0.000053	(0.000077)	0.61	(0.23)	0.00015	(0.0000)
3-ethyltoluene	n.d.	(-)	,	:	0.80	(1.53)	0.00031	(0.0065)	0.28	(0.16)	0.00011	(0.00008)
4-ethyltoluene	0.12	(0.05)	0.000036	(0.000016)	0.27	(0.12)	0.000032	(0.000045)	0.35	(0.29)	0.000032	(0.000067)

Table B1 (continued).												
		- 29 -	- 30 April			22 -	- 25 May			- 74 -	- 26 July	
	Mixing n	atio [ppt _v]	$R_{\rm OH}$	[s ⁻¹]	Mixing ra	tio [pptv]	$R_{\rm OH}$	$[s^{-1}]$	Mixing ra	tio [pptv]	$R_{\rm OH}$	s ⁻¹]
	mean	(std)	mean	(std)	mean	(std)	mean	(std)	mean	(std)	mean	(std)
1,2,3-trimethylbenzene	2.2	(0.8)	0.0019	(0.0007)	2.4	(2.4)	0.0019	(0.0020)	1.4	(0.5)	0.000	(0.0006)
1,2,4-trimethylbenzene	3.3	(1.0)	0.0028	(0.000)	0.56	(0.52)	0.00028	(0.00040)	0.44	(0.21)	0.00029	(0.00019)
1,3,5-trimethylbenzene	1.3	(0.7)	0.0021	(0.0011)	0.37	(0.21)	0.00018	(0.00031)	0.24	(0.10)	0.00024	(0.00019)
isoprene	3.9	(3.0)	0.010	(600.0)	8.0	(6.4)	0.020	(0.017)	29	(14)	090.0	(0.041)
Monoterpenoids			0.53	(0.39)			0.47	(0.53)			1.7	(1.4)
α -pinene	223	(142)	0.33	(0.22)	120	(134)	0.16	(0.19)	635	(318)	0.70	(0.51)
eta-pinene	28	(26)	090.0	(0.058)	24	(27)	0.047	(0.053)	105	(74)	0.17	(0.15)
camphene	22	(16)	0.030	(0.022)	29	(29)	0.037	(0.038)	52	(33)	0.057	(0.047)
Δ^3 -carene	4	(35)	0.10	(0.08)	72	(82)	0.16	(0.18)	224	(141)	0.40	(0.33)
<i>p</i> -cymene	5.5	(2.3)	0.0021	(0.000)	23	(24)	0.0080	(0.0087)	11	(5)	0.0033	(0.0022)
limonene	1.8	(1.4)	0.0037	(0.0061)	12	(14)	0.050	(0.058)	76	(65)	0.33	(0.28)
terpinolene	n.d.	-		(-)	0.53	(0.37)	0.00068	(0.00160)	11	(2)	0.051	(0.041)
myrcene	0.25	(0.26)	1.6e-12	(1.9e-12)	4.2	(3.0)	1.8e-11	(2.2e-11)	14	(8)	8.0e-11	(6.6e-11)
1,8-cineol	2.6	(2.4)	0.00076	(0.00069)	12	(6)	0.0033	(0.0024)	22	(6)	0.0050	(0.0032)
bornylacetate	0.31	(0.20)	0.00011	(0.00007)	1.7	(0.0)	0.00033	(0.00038)	2.7	(1.3)	0.00075	(0.00052)
Sequiterpenes			0.0015	(0.0030)			0.022	(0.024)			0.17	(0.14)
longicyclene	0.32	(0.27)	0.000079	(0.000066)	0.81	(0.27)	0.0000	(0.00010)	0.78	(0.37)	0.00015	(0.00010)
iso-longifolene	0.0600	(0.0003)	0.000042	(0.000068)	0.28	(0.13)	0.000036	(0.000168)	n.d.	-		(-)
β -farnesene	n.d.	-		(-)	n.d.	(-)		(-)	4.0	(1.4)	0.014	(0.008)
eta-cary ophyllene	0.94	(0.61)	0.0013	(0.0027)	7.3	(3.7)	0.020	(0.023)	28	(16)	0.11	(0.0)
lpha-humulene	0.0514	(0.0001)	0.000071	(0.000149)	0.21	(0.15)	0.0014	(0.0011)	n.d.	-)		(-)
SQT1*	n.d.	-)		(-)	n.d.	(-)		(-)	2.7	(1.5)	0.0055	(0.0042)
SQT2*	n.d.	(-)		(-)	n.d.	(-)		(-)	5.4	(3.3)	0.011	(0.00)
SQT3*	n.d.	-)		(-)	n.d.	(-)		(-)	4.5	(2.5)	0.0072	(0.0072)
SQT4*	n.d.	(-)		(-)	n.d.	(-)		(-)	12	(9)	0.017	(0.018)
GLVs				(-)			0.0021	(0.0020)			0.013	(0.020)
1-hexanol	n.d.	(-)		-	n.d.	-		(-)	7.8	(4.2)	0.0010	(0.0017)
cis-2-hexen-1-ol	n.d.	-)		(-)	n.d.	(-)		(-)	n.d.	-)		(-)
trans-2-hexen-1-ol	n.d.	-		(-)	n.d.	(-)		(-)	n.d.	(-)	,	(-)
cis-3-hexen-1-ol	n.d.	(-)		(-)	n.d.	(-)		(-)	4.3	(2.4)	0.0026	(0.0056)
trans-3-hexen-1-ol	n.d.	(-)		(-)	n.d.	(-)		(-)	6.7	(1.5)	0.0063	(0.0096)
trans-2-hexenal	n.d.	-		(-)	2.4	(1.8)	0.0021	(0.0020)	3.3	(2.1)	0.0027	(0.0025)
hexylacetate	n.d.	-	ı	-	n.d.	-	ı	-	n.d.	-		(-)
cis-3-hexenylacetate	n.d.	-)		(-)	n.d.	-)		(-)				
n.d.	(-)		(-)									
trans-2-hexenyl-acetate	n.d.	-		•	n.d.	-)		(-)	n.d.	-	ı	(-)

Table B1 (continued).												
		29 -	- 30 April			- 22	- 25 May			24	– 26 July	
	Mixing ra	tio [pptv]	$R_{ m OH}$	[s ⁻¹]	Mixing rat	tio [pptv]	$R_{\rm OH}$	[s ⁻¹]	Mixing ra	atio [pptv]	$R_{\rm OH}$	[s ⁻¹]
	mean	(std)	mean	(std)	mean	(std)	mean	(std)	mean	(std)	mean	(std)
Aldehydes			0.10	(0.07)			0.075	(0.051)			0.35	(0.12)
formaldehyde	121	(110)	0.028	(0.025)	n.m.	(-)		-	620	(06)	0.13	(0.02)
acetaldehyde	16.5	(0.1)	0.0019	(0.0030)	n.m.	-)	,	(-)	342	(62)	0.13	(0.02)
propanal	86	(32)	0.046	(0.017)	93	(49)	0.040	(0.027)	112	(36)	0.017	(0.027)
butanal	n.d.	-		(-)	4.7	(1.5)	0.00039	(0.00104)	17	(26)	0.0085	(0.0148)
pentanal	19	(9)	0.015	(0.005)	24	(20)	0.011	(0.015)	41	(16)	0.028	(0.011)
hexanal	8.03	(0.04)	0.0018	(0.0029)	7.3	(3.3)	0.0052	(0.0025)	17	(8)	0.012	(0.005)
heptanal	n.d.	(-)	,	(-)	5.9	(1.5)	0.0043	(0.0013)	0.10	(0.08)	0.0000017	(0.0000143)
octanal	n.d.	(-)		(-)	4.2	(1.0)	0.0032	(0.000)	6.1	(1.7)	0.0040	(0.0021)
nonanal	n.d.	(-)		(-)	2.8	(1.0)	0.0024	(0.000)	9.4	(4.1)	0.0069	(0.0045)
decanal	n.d.	(-)		(-)	3.1	(0.8)	0.0026	(0.0008)	9.9	(3.0)	0.0070	(0.0039)
methacrolein	8.1	(1.4)	0.0030	(0.0033)	8.0	(3.3)	0.0058	(0.0025)	7.1	(6.5)	0.0043	(0.0045)
crotonaldehyde	1.6	(0.1)	0.00080	(0.00071)	n.m.	-		-	n.d.	-)	,	(-)
benzaldehyde	26	(2)	0.0016	(0.0035)	n.m.	-)	·	(-)	n.d.	-	ı	(-)
tolualdehyde	75	(2)	0.0058	(0.0128)	n.m.	(-)	,	(-)	n.d.	(-)		(-)
Alcohols			0.087	(0.080)			0.21	(0.56)			0.050	(0.123)
isopropanol	26	(9)	0.0035	(0.008)	37	(29)	0.0041	(0.0039)	171	(95)	0.0069	(0.0121)
1-butanol	367	(350)	0.083	(0.079)	1122	(2704)	0.21	(0.55)	614	(745)	0.041	(0.105)
1-pentanol	n.d.	-		(-)	3.7	(1.4)	0.000065	(0.000285)	8.6	(3.3)	0.00076	(0.0129)
1-penten-3-ol	n.d.	(-)	,	(-)	1.9	(0.7)	0.00041	(0.00120)	3.6	(2.3)	0.0015	(0.0034)
3-methyl-2-buten-1-ol	n.d.	-		(-)	n.d.	(-)		(-)	n.d.	-)		-
1-octen-3-ol	n.d.	(-)	'	(-)	n.d.	(-)	·	(-)	1.5	(0.4)	0.00026	(0.00059)
2-methyl-3-buten-2-ol (MBO)	5.4	(4.6)	0.021	(0.018)	15	(16)	0.054	(0.058)	47	(28)	0.14	(0.11)
Other carbonyls			0.014	(0.018)			0.0012	(0.0019)			0.050	(0.022)
acetone (and acrolein)	3073	(4151)	0.013	(0.017)	n.m.	(-)		(-)	9161	(1632)	0.038	(0.007)
6-methyl-2-hepten-3-one	n.d.	(-)		(-)	n.d.	(-)		(-)	1.5	(0.6)	0.00015	(0.00063)
methyl ethyl ketone (MEK)	n.d.	(-)		(-)	n.d.	(-)		(-)	9.0	(0.3)	0.00023	(0.00005)
butylacetate	2.9	(1.3)	0.000052	(0.000143)	n.d.	(-)		(-)	n.d.	-)		-
4-acetyl-1-methylcyclohexene	n.d.	-		(-)	1.3	(0.6)	0.00022	(0.00105)	4.5	(4.0)	0.0082	(0.0120)
nopinone	4.8	(3.2)	0.0018	(0.0012)	2.9	(2.5)	0.0010	(0.009)	10	(4)	0.0030	(0.0019)
Organic acids			0.071	(0.013)			0.024	(0.018)			0.029	(0.044)
acetic acid	2800	(447)	090.0	(0.008)	1507	(430)	0.020	(0.014)	3007	(283)	0.017	(0.026)
propanoic acid	142	(25)	0.0044	(0.0008)	84	(26)	0.0018	(0.0013)	160	(53)	0.0015	(0.0024)
butanoic acid	98	(38)	0.0046	(0.0018)	58	(33)	0.0018	(0.0017)	153	(26)	0.0021	(0.0032)
isobutanoic acid	n.d.	-)		-	n.d.	-)		•	32	(14)	0.00028	(0.00069)
pentanoic acid	20	(13)	0.0016	(0.0015)	21	(11)	0.00055	(0.00109)	176	(36)	0.0057	(0.0086)

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	[s ⁻¹] (std)	(0.00046)	(-)	(0.0024)	(0.00095)	(0.2)	(0.010)	(0.027)	(0.016)	(0.0021)	(0.11)	(0.01)	(0.36)	(0.005)	(9)	(5.7)
- 26 July	$R_{ m OH}$ mean	0.00022	ı	0.0015	0.00050	1.1	0.010	0.033	0.046	0.0012	0.74	0.28	0.78	0.077	12	16.2
- 24 -	ratio [ppt _v] (std)	(5.2)	(-)	(11)	(3)		(42)	(94)	(8.9e3)	(114)	(2.1e4)	(2.2e-16)				
	Mixing	8.5	n.d.	35	13		69	149	2.7e4	74	1.4e5	1.9e6				
	[s ⁻¹] (std)	(0.0000234)	(-)	(0.000275)	(-)	(0.2)	(0.013)	(0.08)	(0.014)	(0.00054)	(0.03)	(0.02)	(0.12)	(0.007)	(2)	(7.2)
– 25 May	$R_{ m OH}$ mean	0.0000025	ı	0.000055		1.0	0.013	0.11	0.69	0.00058	0.58	0.26	0.25	0.070	10	12.7
22	ratio [pptv] (std)	(0.5)	(-)	(2.8)	(-)		(52)	(295)	(7.4e3)	(25)	(6.1e3)	(5.1e3)				
	Mixing	2.0	n.d.	9.7	n.d.		89	418	4.2e4	37	1.1e5	1.9e6				
	[s ⁻¹] (std)	(0.0000122)	(-)	(0.000217)	(-)	(0.2)	(0.012)	(0.12)	(0.015)	(0.00132)	(0.06)	(0.01)	(0.06)	(0.004)	(29)	(29.3)
– 30 April	$R_{ m OH}$ mean	0.0000014	,	0.000052		1.2	0.013	0.14	0.066	0.00091	0.72	0.23	0.12	0.057	46	49.0
29	io [ppt _v] (std)	(0.2)	(-)	(2.1)	(-)		(41)	(377)	(9.0e3)	(09)	(9.4e3)	(2.2e3)				
	Mixing rat mean	1.0	n.d.	5.9	n.d.		LL	450	4.28e4	53	1.29e5	1.9e6				
		isopentanoic acid	4-methylpentanoic acid	hexanoic acid	heptanoic acid	Inorganics	NO	NO_2	0_3	SO_2	CO	CH_4	Model OVOCs	Model inorganics	Missing	Total

* quantified with β -caryophyllene calibration and an estimated reaction coefficient (1e-10 cm³ s⁻¹)