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What effect does VOC sampling time have on derived OH reactivity?

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Abstract. State-of-the-art techniques allow for rapid measurements of total OH reactivity. Unknown sinks of OH and oxidation processes in the atmosphere have been attributed to what has been termed "missing" OH reactivity. Often overlooked are the differences in timescales over which the diverse measurement techniques operate. Volatile organic compounds (VOCs) acting as sinks of OH are often measured by gas chromatography (GC) methods which provide lowfrequency measurements on a timescale of hours, while sampling times are generally only a few minutes. Here, the effect of the sampling time and thus the contribution of unmeasured VOC variability on OH reactivity is investigated. Measurements of VOC mixing ratios by proton transfer reaction time-of-flight mass spectrometry (PTR-ToF-MS) conducted during two field campaigns (ClearfLo and PARADE) in an urban and a semi-rural environment were used to calculate OH reactivity. VOCs were selected to represent variability for different compound classes. Data were averaged over different time intervals to simulate lower time resolutions and were then compared to the mean hourly OH reactivity. The results show deviations in the range of 1 to 25 %. The observed impact of VOC variability is found to be greater for the semi-rural site.

The selected compounds were scaled by the contribution of their compound class to the total OH reactivity from VOCs based on concurrent gas chromatography measurements conducted during the ClearfLo campaign. Prior to being scaled, the variable signal of aromatic compounds results in larger deviations in OH reactivity for short sampling intervals compared to oxygenated VOCs (OVOCs). However, once scaled with their lower share during the ClearfLo campaign, this effect was reduced. No seasonal effect on the OH reactivity distribution across different VOCs was observed at the urban site.

1 Introduction

Atmospheric photochemistry produces a variety of radicals that exert a substantial influence on the ultimate composition of the atmosphere. The OH radical is the main oxidant in the atmosphere (Monks et al., 2009, and references therein). Its actual concentration is determined by the balance between its sources and sinks. While in many cases OH sources are well understood, OH sinks are manifold and not completely characterised. OH reactivity is a measure of the strength of the sinks for the OH radical. It can be derived from the reaction rates of the reactants k_{OH+X} and their concentrations [X] (Kovacs et al., 2003):

$$k_{\text{OH}} = \sum k_{\text{OH}+\text{VOC}_i} [\text{VOC}_i] + k_{\text{OH}+\text{CO}} [\text{CO}] + k_{\text{OH}+\text{NO}} [\text{NO}] + k_{\text{OH}+\text{NO}_2} [\text{NO}_2] + k_{\text{OH}+\text{SO}_2} [\text{SO}_2] + \dots$$
(1)

In situ measurements of OH reactivity have provided new insights into OH loss chemistry and the oxidative ability of the atmosphere (e.g. Di Carlo et al., 2004; Edwards et al., 2013; Hofzumahaus et al., 2009; Whalley et al., 2011;

Yoshino et al., 2006). There are a number of different techniques used for the direct measurement of OH reactivity. The total OH loss rate measurement technique (TOHLM) was one of the first techniques applied for determination of total OH reactivity based on a single measurement (Ingham et al., 2009; Ren et al., 2003a; Shirley et al., 2006). TOHLM is based on the measurement of the decay of artificially produced OH following the introduction of reactants into an ambient air sample within a flow tube. By varying the distance between the OH injection point and the detector, the reaction time changes and provides a series of relative decay rates (Kovacs et al., 2003; Kovacs and Brune, 2001). A similar approach is taken with the laser-induced pump and probe technique, whereby decay in OH is detected by timeresolved laser-induced fluorescence (Sadanaga et al., 2004). Another technique developed by Sinha et al. (2008), called the comparative reactivity method (CRM), is based on the measurement of a single reactant (most often pyrrole) which first reacts with OH under clean air conditions and then under competitive conditions with ambient air. The reaction takes place in a glass vessel and is most commonly probed by PTR-MS. Recently, Nölscher et al. (2012b) presented a GC-PID for the detection of pyrrole for CRM.

These techniques enable comparison of directly measured OH reactivity to calculated OH reactivity using Eq. (1) based on measurements of individual compounds. The difference between the two is referred to as missing OH reactivity. Reasons for an under-prediction of OH reactivity maybe due to incomplete or inaccurate measurements of individual compounds (Di Carlo et al., 2004; Kim et al., 2011; Kovacs and Brune, 2001). Therefore, direct measurements of total OH reactivity can help to evaluate the completeness of measured VOC budgets (Dolgorouky et al., 2012; Mao et al., 2009; Mogensen et al., 2011).

In urban environments good agreement between measured and calculated OH reactivity has been found. For example, no significant missing OH reactivity was found in New York during summer (Ren et al., 2003b) and for both Paris under clean marine air conditions (Dolgorouky et al., 2012) and Tokyo (Yoshino et al., 2006) in the winter. Larger missing OH reactivity of up to 30% was found for all other seasons in Tokyo by Yoshino et al. (2006), presumably owing to secondary reaction products, including semi-volatile oxygenated compounds, from atmospheric oxidation of VOCs. A similar amount of missing OH reactivity was reported by Kovacs et al. (2003) for urban measurements in Nashville. They suggest that non-measured short-lived VOCs accounted for the missing reactivity. In Paris, a missing OH reactivity of up to 75 % was found for continentally influenced air, which is also attributed to highly oxidised compounds from photochemical processes during transportation of these air masses (Dolgorouky et al., 2012). Similar reasons were reported by Lou et al. (2010) to account for missing OH reactivity measured in the highly populated Pearl River Delta.

Direct measurements of OH reactivity in rural areas generally tend to have larger missing OH reactivity. Using PTR-MS and the CRM method in a boreal forest in Finland during August 2008, Sinha et al. (2010) reported missing OH reactivity of approximately 50 %. This site was revisited in 2010, when missing OH reactivity of 58 to 89% was recorded (Nölscher et al., 2012a). Similar results in a mixed deciduous forest were obtained by Hansen et al. (2014), who reported missing OH reactivity of 46 to 65 %. Both studies concluded that unmeasured oxidation products were missing from the OH reactivity calculation. Undetected biogenic emissions and transport of reactive compounds are also cited as other reasons for missing OH reactivity. In contrast to those findings, Ren et al. (2006) found no significant missing OH reactivity on average during a summertime campaign in a deciduous forest in New York in 2002. They attributed this to differences in the composition of emitted biogenic VOCs (BVOCs). Rainforests are a large sink for OH as they emit a huge amount of VOCs. Measured OH reactivity in the rainforest of Borneo during April 2008 yielded a missing OH reactivity of 70% compared to calculated reactivity from measurements of single compounds (Edwards et al., 2013) and \sim 53 % compared to modelled reactivity (Whalley et al., 2011). Since isoprene makes up the biggest contribution to OH reactivity, the effect of oxidation products of isoprene was discussed (Edwards et al., 2013; Whalley et al., 2011).

While different possible explanations for missing OH reactivity have been given, the wide range in reported missing OH reactivity suggests that many reactants and processes remain unknown or cannot be measured at present. Measurements of total non-methane organic carbon in the western Los Angeles Basin (Chung et al., 2003) and results following the application of a double-column (orthogonal) GC for urban air measurements (Lewis et al., 2000) emphasise the large number of OH reactants that are not measured with standard field equipment.

Measurements of non methane hydrocarbons (NMHCs) used for calculation of OH reactivity are often performed with GC (Lou et al., 2010; Sadanaga et al., 2005; Shirley et al., 2006) and therefore the time resolution of the calculated OH reactivity is low due to long sampling and run times, e.g. canister samples once every hour (Kovacs et al., 2003; Sadanaga et al., 2005; Yoshino et al., 2006), when compared to measured total OH reactivity. However, the sampling time during one GC cycle is often shorter than the analysis time, and thus any high temporal variability in measured OH reactivity is not easily captured when it is derived from GC data (Nölscher et al., 2012a). When measured and calculated OH reactivity are compared, high-time-resolution data are often averaged over intervals that correspond to the GC cycle. The GC-FID systems for measurements of NMHCs used by Dolgorouky et al. (2012), for example, were sampling over 10 min and had an additional analysis run time of 20 min, resulting in a time resolution of 30 min. Their GC-MS for measurements of OVOCs used a sampling time of 30 min, resulting in an overall time resolution of 90 min, corresponding to the frequency they calculated OH reactivity.

This work addresses the question of how temporal VOC concentration variability is reflected with different sampling time resolutions. Furthermore, the effect of averaging VOC data on calculated OH reactivity is discussed alongside how this may affect the amount of "missing" OH reactivity.

Relatively high time-resolved VOC data collected by PTR-ToF-MS are used to calculate OH reactivity for selected compounds. Differing time resolutions are analysed to explore the effects. Data from an urban winter campaign are compared to measurements from a semi-rural summer campaign.

2 Experimental section

Two different sets of VOC mixing ratios measured with PTR-ToF-MS were used for analysis. One was collected during the ClearfLo (Clean Air for London, www.clearflo.ac.uk) (Bohnenstengel et al., 2015) winter campaign in 2012 at an urban background site in London, UK. The second was taken during the PARADE (PArticles and RAdicals: Diel observations of the impact of urban and biogenic Emissions, http://parade2011.mpich.de/) campaign in late summer 2011 at a semi-rural site located in the Taunus ridge, Germany.

2.1 Field data

2.1.1 ClearfLo

A PTR-ToF-MS (Series I; Kore Technology Ltd, UK) (see standard PTR-MS apparatus in Barber et al., 2012; Thalman et al., 2015) was deployed at Sion-Manning School (51°31'15" N, 0°12'51" W) near the North Kensington urban background station in London during the intensive observation periods of the ClearfLo project in 2012. A general overview of the ClearfLo project and the measurement site is given in Bohnenstengel et al. (2015). For background measurements a hydrocarbon trap (activated carbon filter by Grace Alltech) was employed once during the time period investigated here. Its efficiency was in the range of 87–96 %. Calibration measurements were performed before (acetone) and after (toluene and xylene) the campaign in the laboratory. For the calibration of toluene and xylene a permeation tube was used, and calibration of acetone was done by dilution of a gas standard with zero air. The stability of the instrument during the campaign was monitored with a bromobenzene internal standard. Based on these measurements no correction needed to be applied. Of the two intensive observation periods (IOPs) (i.e. winter: 6 January to 11 February; summer: 21 July to 23 August) data from 1 to 7 February 2012 were selected for analysis in this study. During this period the measurement site was influenced by local sources, as well as by air masses from other parts of the UK and the continent (Bohnenstengel et al., 2015).

A dual-channel GC with flame ionisation detector (DC-GC-FID; Hopkins et al., 2003) was deployed at the same site as the PTR-ToF-MS during the ClearfLo IOPs. A wide range of VOCs, including alkanes, alkenes, dienes, aromatic compounds and OVOCs, were measured (see Table 1). Stainless steel tubing heated to $80 \,^{\circ}$ C was used as a sampling line. This setup ensured the destruction of ozone present in the sample. The sampling time was 10 min, while the analysis runtime was around 50 min, resulting in approximately one measurement per hour.

2.1.2 PARADE

For comparison, data collected with a PTR-ToF-MS (Ionicon Analytik GmbH, Austria) (described in Jordan et al., 2009) during the PARADE field campaign were analysed. Measurements were taken between 15 August and 9 September 2011 at the Taunus observatory on the summit of Kleiner Feldberg (50°13'25" N, 8°26'56" E) under various meteorological conditions. A detailed description of the measurement site and measurements performed during PARADE can be found in Crowley et al. (2010) and Bonn et al. (2014). The PTR-ToF-MS was operated continuously with minor interruptions. Background measurements were conducted regularly with zero air throughout the campaign and calibration measurements were performed with a multicomponent gas standard before and after the campaign in the laboratory. For this study, 2 weeks of data (21 to 27 August 2011 – Period 1; 1 to 6 September 2011 - Period 2) were selected, each with approximately the same amount of data points as the ClearfLo data set. Period 1 was mainly influenced by continental air masses and only towards the end by air that travelled over the UK and the English Channel (UK-marine). Period 2 was dominated by UK-marine air, but was also influenced by air masses that travelled over the Atlantic (see Phillips et al., 2012).

2.1.3 Data

While the ClearfLo data presented here were collected at an urban background site with mainly anthropogenic emissions, the PARADE campaign took place at a semi-rural site. Biogenic emissions were expected from the direct vicinity, but some anthropogenic influence was apparent from the proximity of the highly populated Rhein-Main area and Frankfurt.

Three mass channels were selected for the analysis corresponding to acetone/propanal, toluene and ethylbenzene/xylene. For brevity, the combined signal of acetone and propanal is referred to as acetone, and that of ethylbenzene and xylene is referred to as xylene. The compounds used for analysis represent different sources of VOCs. Toluene and xylene are counted along anthropogenic VOCs, monoterpenes are of biogenic origin, and the OVOCs (acetone and methanol) are emitted directly or produced by photochemi-

Compound	VMR (ppbV)	Concentration (molecules cm^{-3})	k_{OH} (cm ³ molecules ⁻¹ s ⁻¹)	OH reactivity (s ⁻¹)
Alkanes				
Ethane ^a	12.91 ± 10.89	$(3.14 \pm 2.65) \times 10^{11}$	2.40×10^{-13}	0.075
Propane ^a	4.59 ± 3.35	$(1.12 \pm 0.81) \times 10^{11}$	1.10×10^{-12}	0.123
iso-Butane ^b	1.42 ± 1.00	$(3.45 \pm 2.43) \times 10^{10}$	2.12×10^{-12}	0.073
<i>n</i> -Butane ^b	2.35 ± 1.60	$(5.71 \pm 3.89) \times 10^{10}$	2.36×10^{-12}	0.135
Cyclopentaneb	0.10 ± 0.11	$(2.50 \pm 2.58) \times 10^9$	4.97×10^{-12}	0.012
iso-Pentane ^b	0.83 ± 0.62	$(2.03 \pm 1.50) \times 10^{10}$	3.60×10^{-12}	0.073
<i>n</i> -Pentane ^b	0.42 ± 0.26	$(1.02 \pm 0.64) \times 10^{10}$	3.80×10^{-12}	0.039
2,3-Methylpentaneb*	0.35 ± 0.29	$(8.56 \pm 6.93) \times 10^9$	3.10×10^{-11}	0.265
<i>n</i> -Hexane ^b	0.13 ± 0.09	$(3.16 \pm 2.29) \times 10^9$	5.20×10^{-12}	0.016
<i>n</i> -Heptane ^b	0.09 ± 0.07	$(2.18 \pm 1.58) \times 10^9$	6.76×10^{-12}	0.015
2,2,4-TMP ^b	0.04 ± 0.02	$(9.88 \pm 5.22) \times 10^8$	3.34×10^{-12}	0.003
<i>n</i> -Octane ^b	0.03 ± 0.02	$(6.75 \pm 3.75) \times 10^8$	8.11×10^{-12}	0.005
Alkenes				
Ethene ^a	1.93 ± 1.04	$(4.68 \pm 2.52) \times 10^{10}$	7.80×10^{-12}	0.365
Propene ^a	0.43 ± 0.30	$(1.05\pm 0.73)\times 10^{10}$	2.90×10^{-11}	0.306
trans-2-Buteneb	0.04 ± 0.03	$(1.03 \pm 0.81) \times 10^9$	6.40×10^{-11}	0.066
1-Butene ^b	0.08 ± 0.05	$(1.90 \pm 1.21) \times 10^9$	3.14×10^{-11}	0.060
iso-Butene ^a	0.11 ± 0.07	$(2.63 \pm 1.77) \times 10^9$	5.10×10^{-11}	0.134
cis-2-Butene ^b	0.03 ± 0.02	$(6.92 \pm 5.72) \times 10^8$	5.64×10^{-11}	0.039
trans-2-Penteneb	0.04 ± 0.03	$(9.13 \pm 7.37) \times 10^8$	6.70×10^{-11}	0.061
1-Pentene ^b	0.03 ± 0.02	$(7.32 \pm 5.27) \times 10^8$	3.14×10^{-11}	0.023
Acetylene ^a	1.43 ± 0.74	$(3.47 \pm 1.81) \times 10^{10}$	7.50×10^{-13}	0.026
Dienes				
Propadiene ^b	0.02 ± 0.01	$(4.40 \pm 2.61) \times 10^8$	9.82×10^{-12}	0.004
1,3-Butadiene ^b	0.05 ± 0.03	$(1.14 \pm 0.76) \times 10^9$	6.66×10^{-11}	0.076
Isoprene ^a	0.02 ± 0.02	$(5.37 \pm 4.07) \times 10^8$	1.00×10^{-10}	0.054
Aromatic compounds				
Benzene ^a	0.41 ± 0.17	$(9.88 \pm 4.06) \times 10^9$	1.20×10^{-12}	0.012
Toluene ^a	0.64 ± 0.48	$(1.56 \pm 1.17) \times 10^{10}$	5.60×10^{-12}	0.087
Ethylbenzene ^b	0.14 ± 0.11	$(3.48 \pm 2.57) \times 10^9$	7.00×10^{-12}	0.024
m+p-Xylene ^{b*}	0.18 ± 0.14	$(4.28 \pm 3.52) \times 10^9$	1.87×10^{-11}	0.080
o-Xylene ^b	0.17 ± 0.12	$(4.02 \pm 2.82) \times 10^9$	1.36×10^{-11}	0.055
Oxygenated VOCs				
Acetaldehyde ^a	2.37 ± 1.38	$(5.77 \pm 3.35) \times 10^{10}$	1.50×10^{-11}	0.866
MACR ^b	0.16 ± 0.12	$(3.89 \pm 2.97) \times 10^9$	2.90×10^{-11}	0.113
Methanol ^a	1.44 ± 0.81	$(3.50\pm 1.96)\times 10^{10}$	9.00×10^{-13}	0.031
Acetone ^a	1.11 ± 0.51	$(2.69 \pm 1.24) \times 10^{10}$	1.80×10^{-13}	0.005
MVK ^b	0.28 ± 0.15	$(6.72 \pm 3.61) \times 10^9$	2.00×10^{-11}	0.134
Ethanol ^a	5.48 ± 3.81	$(1.33\pm 0.93)\times 10^{11}$	3.20×10^{-12}	0.426
Propanol ^a	0.31 ± 0.21	$(7.41 \pm 5.15) \times 10^9$	5.80×10^{-12}	0.043
Butanol ^a	0.59 ± 0.33	$(1.45 \pm 0.80) \times 10^{10}$	8.50×10^{-12}	0.123

Table 1. Mean mixing ratios and concentrations with standard deviation, rate coefficient and OH reactivity of the VOCs measured with DC-GC-FID during ClearfLo from 1 to 7 February 2012.

^a IUPAC preferred value; ^b Atkinson and Arey (2003); * average of both.

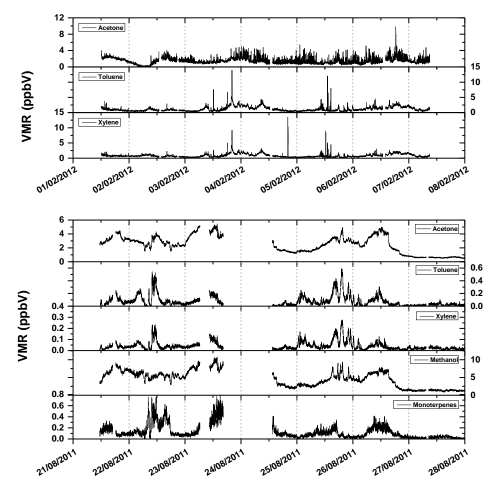


Figure 1. Time series of VOCs during ClearfLo (top) and PARADE Period 1 (bottom). The time resolution is 1 min.

cal oxidation in the atmosphere (Monks et al., 2009, and references therein). They were selected because their volume mixing ratios could be determined with low uncertainty for both instruments. Aromatic compounds such as toluene and xylene are well suited for this investigation because they often show short-term high variability. The analysis of the PA-RADE data also includes methanol and the sum of monoterpenes. The characteristic parameters of the measurements during ClearfLo and PARADE are given in Table 2.

Figure 1 shows the time series of the VOCs for ClearfLo (top) and PARADE Period 1 (bottom). The range of mixing ratios for ClearfLo is much wider and higher mixing ratios are reached. Acetone shows values up to a factor of 1.8 higher in ClearfLo compared to PARADE, while the aromatic compounds are 2 orders of magnitude higher. This emphasises the diversity of the two field sites. In the box plots, presented in Fig. 2, some interesting patterns are apparent. For ClearfLo all three compounds exhibit a similar interquartile range (0.60 to 0.86 ppbV) but also very high maximum values. For PARADE a different distribution is depicted. Acetone has a wider interquartile range of 1.83 ppbV and has a higher mean value than toluene and xylene. The aromatic compounds have a much smaller range compared to ClearfLo (0.03 to 0.08 ppbV). Methanol has a wider range than acetone and the monoterpenes look similar to the aromatic compounds. Both periods of PARADE show the same pattern. The ranges of the mixing ratios during the campaigns are summarised in Table 3. Values below the limit of detection (LOD) as well as negative values are not disregarded in this analysis to preserve the full range of the data in order that they can be compared to a randomly generated data set.

OH reactivity relating to the VOCs under study is calculated from the first term of Eq. (1). Reaction acetone $(1.8 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}),$ for rates $(5.6 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}),$ toluene methanol $(9.0 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$ and α -pinene $(5.3 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$ are taken from http://iupac.pole-ether.fr/index.html. The exact composition of the monoterpene signal is not known; thus, only the reaction rate of α -pinene is used. For xylene the average of the reaction rates of ethylbenzene and o-, m- and p-xylene $(14.5 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})$ (Atkinson and Arey, **Table 2.** Characteristics of the two different PTR-ToF-MS deployed during ClearfLo and PARADE. Given are the sensitivities based on normalised counts per second (ncps), accuracy as error for the measurements, and the limit of detection (LOD), which was calculated as 1σ for ClearfLo and 2.6 σ for PARADE based on 1 min data.

	Compound	Sensitivity (ncps ppbV ⁻¹)	Accuracy (%)	$\begin{array}{c} \text{LOD} (1\sigma) \\ (\text{ppbV}) \end{array}$
	Acetone	9.89	18*	0.56
ClearfLo	Toluene	6.36	18* (22)	0.38
	Xylene	9.00	18* (20)	0.41
	Compound	Sensitivity	Accuracy	LOD (2.6σ)
		$(ncps ppbV^{-1})$	(%)	(ppbV)
	Acetone	37.0	16	0.08
	Toluene	26.9	8	0.04
PARADE	Xylene	33.4	13	0.01
	Methanol	12.7	17	0.24
	Monoterpenes	14.1	10	0.02

Effects of isobaric overlap from fragmentation taken into account. *First column does not include effect of isobaric overlap from aromatic fragmentation; second column includes estimation of isobaric overlap.

Table 3. Overview of the range of VOC mixing ratios in ppbV during ClearfLo and PARADE (PAR).

		Minimum	Maximum	Mean	Interquart. range	Max–Min
ClearfLo	Acetone	-0.294	9.816	1.459	0.864	10.110
	Toluene	0.058	13.982	1.162	0.862	13.924
	Xylene	0.038	13.519	0.861	0.601	13.482
PAR 1	Acetone	0.426	5.447	2.400	1.833	5.021
	Toluene	-0.030	0.592	0.076	0.078	0.622
	Xylene	-0.008	0.277	0.041	0.030	0.285
	Methanol	0.851	10.775	4.438	3.858	9.923
	Monoterp.	-0.008	0.801	0.124	0.116	0.809
PAR 2	Acetone	0.544	4.873	1.987	1.797	4.329
	Toluene	-0.017	0.646	0.078	0.073	0.663
	Xylene	-0.004	0.358	0.046	0.039	0.362
	Methanol	0.781	10.649	3.776	2.739	9.869
	Monoterp.	-0.009	0.692	0.075	0.086	0.701

2003) was applied. Table 4 summarises the minimum, maximum and mean reactivity calculated from these VOCs as described.

The time resolution of PTR-ToF-MS is only limited by the signal-to-noise ratio and resulting detection limit. Both instruments were operated with a 1 min time resolution. Volume mixing ratios of VOCs were averaged over different intervals and standard deviations were derived. An average was only included for analysis if its data recovery was at least 50%. The OH reactivity for each VOC was calculated and summed as required. Only the standard deviations were propagated as errors of reactivity, as the focus of this work is on investigating VOC variability.

For clarity throughout this paper the notation $R = R_{OH}$ for reactivity regarding the OH radical replaces k_{OH} (see Eq. 1; see also Nölscher et al., 2012a). Indices denote the origin of the data (PTR (PTR-ToF-MS) or GC (DC-GC-FID) and CL (ClearfLo) or PAR (PARADE)). Numbers indicate the averaging time in minutes. If only some VOCs are taken into account for calculating the reactivity, this will be indicated, e.g. $R_{\text{PTR,CL}}^{\text{OVOC,5}}$ is the OH reactivity calculated from the 5 min mean concentration of acetone, measured with the PTR-ToF-MS during ClearfLo.

2.2 Distribution of OH reactivity of measured VOCs

For a more general view of the factors that drive variation in OH reactivity of VOCs, its frequency distribution was investigated. GC data from the winter (9 January–9 February 2012) and summer IOP (18 July–19 August 2012) during ClearfLo were applied. The OH reactivity, $R_{GC,CL}^{VOC_i}$, was calculated for each measured VOC_i and ranged from 0.003

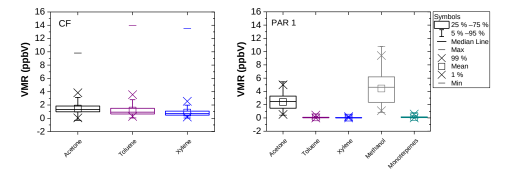


Figure 2. Box plots for ClearfLo (left) and PARADE Period 1 (right) showing the minimum, maximum, mean (), median, interquartile range (box) and percentiles at 1 and 99 % (×).

Table 4. Minimum, maximum and mean OH reactivity and standard deviation calculated from the VOCs under study for ClearfLo and PARADE.

OH reactivity (s^{-1})		Minimum	Maximum	Mean	SD
CF	ATX	0.036	4.864	0.463	0.289
PAR1	ATX	0.000	0.191	0.035	0.028
	ATX+M+MT	0.026	1.296	0.292	0.205
PAR2	ATX	0.001	0.222	0.036	0.024
	ATX+M+MT	0.044	0.987	0.215	0.119

ATX: acetone, toluene and xylene. ATX+M+MT: acetone, toluene, xylene, methanol and monoterpenes .

to 0.822 s^{-1} in winter and from 0.001 to 1.568 s^{-1} in summer with a total OH reactivity $R_{GC,CL}^{TVOC}$ of 4.010 s^{-1} and 3.862 s^{-1} , respectively. The majority of $R_{GC,CL}^{VOC_i}$ values lies below 0.1 s^{-1} , as can be seen from the frequency distribution plotted in Fig. 3, where more than 70% of the winter and 80% of the summer data are in the first interval from 0 to 0.1 s^{-1} . Seasonal differences in OH reactivity emission rates have previously been described by Nölscher et al. (2013) for measurements at a Norway spruce between spring and early autumn. Although the composition of VOCs during ClearfLo changed from winter to summer, no seasonal dependency could be found in the shape of the frequency distribution of $R_{GC,CL}^{VOC_i}$. In both cases $R_{GC,CL}^{TVOC}$ is dominated by the sum of low-reactivity contributions and less by single compounds with high reactivity.

2.3 Generation of a randomised data set

To differentiate between pure statistical effects and measurement related characteristics, a randomised data set was produced and analysed in the same way as the PTR-ToF-MS data. The distribution of OH reactivity is skewed towards smaller values and only positive values of OH reactivity are expected; hence, it is better described by a log-normal distribution compared to a normal distribution (Limpert et al., 2001). The data set of random numbers was generated by

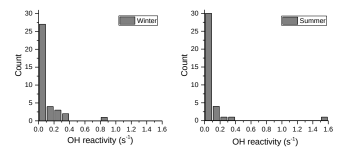


Figure 3. OH reactivity, $R_{GC,CL}^{VOC_i}$, frequency distribution for the ClearfLo campaign in winter (left) and summer (right). Bin size is 0.1 s^{-1} for both plots.

simulating an OH reactivity distribution comparable to the ClearfLo data set. The sample mean $m = 0.463 \text{ s}^{-1}$ and standard deviation SD = 0.289 s^{-1} from the ClearfLo 1 min data set were used to define the parameters μ (Eq. 2) and σ (Eq. 3) for the log-normal distribution of random numbers.

$$\mu = \log\left(\frac{m}{\sqrt{1 + \frac{\mathrm{SD}^2}{m^2}}}\right) \tag{2}$$

$$\sigma = \sqrt{\log\left(1 + \frac{\mathrm{SD}^2}{m^2}\right)} \tag{3}$$

A log-normal distribution of a total of 8040 random numbers was generated using the dlnorm (#, μ , σ) function in *R*. This provides a set of data comparable to 134 h of OH reactivity measurements with a time resolution of 1 min. Figure 4 shows the random data set as a time series together with the hourly mean containing 60 data points. On inspection of Fig. 4, it becomes obvious that the range of the hourly average is very small, with a standard deviation of 0.034 s⁻¹.

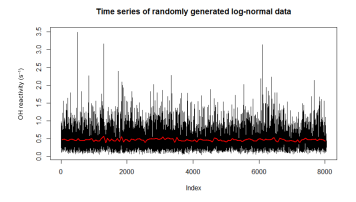


Figure 4. Time series of randomly generated log-normal data set containing 8040 numbers. The "1 min" data are shown in black. The average over 60 data points is plotted in red.

3 Results and discussion

Initially the OH reactivity R_{PTR}^{VOC} was calculated from the PTR data for ClearfLo and PARADE. For both campaigns the signals of acetone, toluene and xylene (referred to generally as VOCs) were used. The effects of differing sampling intervals on the derived reactivity were explored. For each campaign and VOC data set a correlation of the average values of $R_{PTR}^{VOC,t}$ for different intervals (t = 5, 10, 20 and 30 min) against the 60 min average $R_{PTR}^{VOC,60}$ was calculated. The intervals are chosen to be the first t minutes of each hour to simulate the initiation of a GC sequence; thus, the 10 min average also covers the 5 min averaging period and so on.

Figure 5 shows the linear correlation of the 5 min average $R_{\text{PTR,CL}}^{\text{VOC,5}}$ versus the 60 min value $R_{\text{PTR,CL}}^{\text{VOC,60}}$ for the ClearfLo winter campaign. Data were fitted with a bivariate regression line with an intercept (bvf) and forced through the origin (bvfo). The deviation from the slope of the linear regression to a unity gradient $m_{\text{res}} = (m_{R^{<60}/R^{60}} - 1)$ is taken as a measure of how well the value of hourly OH reactivity is represented by the shorter interval average and is further referred to as the residual slope.

3.1 Effects of different sampling intervals

The slopes of both fits in Fig. 5 are below 1.0, indicating an under-prediction of the reactivity during ClearfLo by the value calculated from the first 5 min of each hour. In this case there is only a small deviation (1.3%) from a unity gradient (see Fig. 6). For all averaging intervals the slope is equal to 1 in the range of the uncertainties of the fit.

For the different averaging intervals the difference to the hourly average ($\Delta R = R_{\text{PTR,CL}}^{\text{VOC},t<60} - R_{\text{PTR,CL}}^{\text{VOC},t=60}$) was calculated, and their standard deviations are given in Table 5 as a measure of variance. ΔR generally decreases with increasing averaging time. Also presented in Table 5 are the results from fitted Gaussian functions to the frequency distribution of the **Table 5.** Standard deviation of ΔR and results from Gaussian fits of the ratio of OH reactivity $R_{\text{PTR,CL}}^{\text{VOC},t<60} / R_{\text{PTR,CL}}^{\text{VOC},t=60}$ calculated from shorter interval averages to 60 min average for ClearfLo.

		ΔR	Gaussian fit	
Notation	Time interval (min)	SD (s ⁻¹)	Centre	FWHM
R _{PTR,CL}	5	0.12	0.998 ± 0.011	0.337 ± 0.025
$R_{\rm PTR,CL}^{\rm VOC,10}$	10	0.12	0.997 ± 0.008	0.244 ± 0.020
$R_{\rm PTR,CL}^{\rm VOC,20}$	20	0.10	0.988 ± 0.006	0.246 ± 0.013
$R_{\rm PTR,CL}^{\rm VOC,30}$	30	0.06	0.992 ± 0.004	0.198 ± 0.009

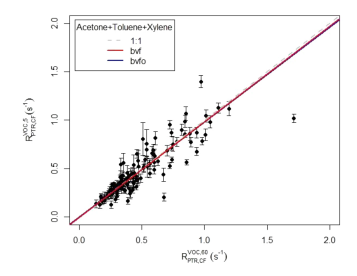


Figure 5. Linear correlation with bivariate fit (bvf: fit with intercept; bvfo: fit forced through the origin) of the OH reactivity calculated from the signals of acetone, toluene and xylene for average intervals of 5 min and 60 min for ClearfLo. The standard deviations of the 5 min means are plotted as error bars.

ratio of the shorter interval averages to the 60 min average. Bins of 0.1 were chosen for the frequency distributions. Ideally, the centre of the Gaussian fit is 1, while the full width at half maximum (FWHM) describes the spread of the distribution around its centre. Both the standard deviation of ΔR and the FWHM decrease when averages are calculated for longer intervals. The centres of all Gaussian fits achieve 0.99.

For Period 1 of the PARADE data (PAR1) the results show a slope greater than 1 (Fig. 7). The high variability in the data is reflected by a higher divergence of the slopes of 1.13 for bvf fit and 1.05 for the bvfo fit based on 5 min averaged data. The small standard deviations of ΔR given in Table 6 highlight the narrow range of calculated OH reactivity $R_{\text{PTR,PAR1}}^{\text{VOC}}$ However, the high variability in the data is reflected by the FWHM of the frequency distributions of the ratios, which is higher for each interval when compared to ClearfLo.

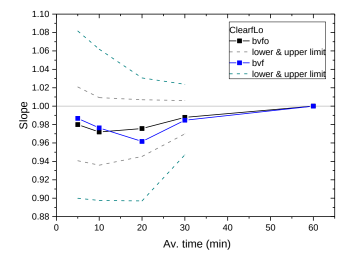


Figure 6. Development of the slope of the correlation of OH reactivity $R_{\text{PTR,CL}}^{\text{VOC},t}$ depending on the sampling interval for ClearfLo. Slopes for bvfo (black) and bvf (blue) with their lower and upper limits are shown.

Table 6. Standard deviation of ΔR and results from Gaussian fits of the ratio of OH reactivity $\frac{R_{\text{PTR,PAR1}}^{\text{VOC},t}}{R_{\text{PTR,PAR1}}^{\text{VOC},t}}$ calculated from shorter interval averages to 60 min average for PARADE – Period 1.

	ΔR	Gaussian fit	
Time interval (min)	SD (s ⁻¹)	Centre	FWHM
5	0.016	0.980 ± 0.011	0.379 ± 0.027
10	0.015	0.976 ± 0.012	0.353 ± 0.026
20	0.012	0.997 ± 0.013	0.310 ± 0.030
30	0.008	0.995 ± 0.009	0.273 ± 0.020
	(min) 5 10 20	Time interval (min) SD (s^{-1}) 5 0.016 10 0.015 20 0.012	Time interval (min) SD (s^{-1}) Centre 5 0.016 0.980 ± 0.011 10 0.015 0.976 ± 0.012 20 0.012 0.997 ± 0.013

For Period 2 (PAR2), an over-prediction of the OH reactivity $R_{\text{PTR,PAR2}}^{\text{VOC}}$ can be observed again (Fig. 8), but with an even greater slope of 1.26. In both periods of PARADE the slope approaches a value of 1 as increasing averaging time takes more of the variability within 1 h into account. Standard deviations of ΔR and FWHM values are similar to Period 1 of the PARADE data, while the centres of the Gaussians are closer to 1 (Table 7).

When OH reactivity is calculated from GC measurements of VOCs, some of the variability in the data is not captured, because air sampling alternates with the GC run itself (Hopkins et al., 2003). In this manner, the analytes are collected for a short duration which is then used to represent the whole measurement cycle. This work suggests that a discrepancy between 60 min averages and shorter intervals can be caused owing to the variable nature of atmospheric VOCs. A sampling time of only 5 min can cause a deviation of more than 25%. Accordingly, this would then artificially contribute to a deviation in OH reactivity, causing a positive or negative

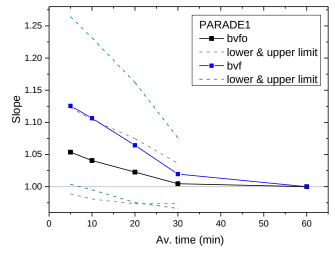


Figure 7. Development of the slope of the correlation of OH reactivity $R_{\text{PTR,PAR1}}^{\text{VOC},t}$ depending on the averaging time for PARADE – Period 1.

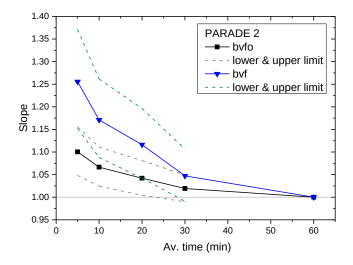


Figure 8. Development of the slope of the correlation of OH reactivity $R_{\text{PTR,PAR2}}^{\text{VOC},t}$ depending on the averaging time for PARADE – Period 2.

bias. Thereby, it is an additional error source when comparing measured total OH reactivity to OH reactivity calculated from GC data.

The deviation is greater for the semi-rural measurements in the Taunus during PARADE compared to the urban measurements in London. Although the range of the analysed OH reactivity of VOCs is smaller during PARADE, the highly frequent fluctuations cause a greater variability in OH reactivity for the investigated intervals.

Table 7. Standard deviation of ΔR and results from Gaussian fits of the ratio of OH reactivity $R_{\text{PTR,PAR2}}^{\text{VOC},t < 60} / R_{\text{PTR,PAR2}}^{\text{VOC},t = 60}$ calculated from shorter interval averages to 60 min average for PARADE – Period 2.

		ΔR	Gaussian fit	
Notation	Time interval (min)	SD (s ⁻¹)	Centre	FWHM
R _{PTR,PAR2}	5	0.013	0.996 ± 0.014	0.352 ± 0.034
$R_{\rm PTR,PAR2}^{ m VOC,10}$	10	0.009	0.994 ± 0.013	0.296 ± 0.031
R _{PTR,PAR2}	20	0.008	0.992 ± 0.008	0.238 ± 0.019
R _{PTR,PAR2}	30	0.006	1.010 ± 0.004	0.238 ± 0.010

3.2 The distribution of residual slopes across consecutive 5 min intervals

In the previous section, only reactivity calculated from the average of the first 5, 10, 20 and 30 min was compared to the hourly mean. Naturally, these averages have different values, depending on the point at which they are selected from the hour under study. They may over- or under-predict the hourly mean as can be seen from Fig. 9, where residual bvf slopes between $R_{\text{PTR}}^{\text{VOC},5}$ and $R_{\text{PTR}}^{\text{VOC},60}$ (see Fig. 5) are plotted for consecutive 5 min averaging periods within the hour. Depending on the selected 5 min interval the bvf resulted in a divergence of -0.1 to 44% for ClearfLo, 1 to 13 % for PAR1, -3 to 26 % for PAR2 and -2 to 10 % for the randomised data. A tendency towards an over-prediction of OH reactivity was observed for both campaigns (ClearfLo - top left; PARADE - bottom) and also for the randomised data set (top right). For the randomised data set byfo was used - a bvf has a much higher slope as the data are clustered together within a small range. On average the residuals are nearly 10% with a standard deviation of 0.1% or less (8.6 $\% \pm 0.1$ % for ClearfLo; 8.85 % ± 0.03 % for PA-RADE 1; $9.5\% \pm 0.1\%$ for PARADE 2; $4\% \pm 4\%$ for the randomised data).

For linear regression the standard least-squares fit is widely used. This method is less adequate when errors in both y and x are assumed or when the assignment of the independent variable is not clear (Isobe et al., 1990). Other methods for bivariate fitting in natural sciences have been discussed in the literature (Cantrell, 2008; Isobe et al., 1990; Warton et al., 2006). Cantrell (2008) found that a bivariate fit is less sensitive to outliers compared to an ordinary least-squares (ols) fit. Warton et al. (2006) described the major axis (ma) and standard or reduced major axis regression (sma/rma). These methods are preferred when the agreement between two measurement techniques is investigated. For equally important deviations from the regression line in the x and y directions ma is used, while sma can be used when the scales in x and y are not comparable. These two functions are implemented in the smatr package in R. The

Table 8. Summary of the statistics of the residual slopes and ratios from the comparisons of the 5 min means to their 60 min means for the ClearfLo data.

Method	Min	Max	Range	Mean	SD
ols	-0.129	0.250	0.379	-0.008	0.112
ols-bis	-0.119	-0.024	0.094	-0.080	0.036
bvfo	-0.033	0.116	0.149	0.014	0.043
bvf	-0.019	0.443	0.462	0.861	0.130
Ratio	-0.030	0.031	0.061	-0.006	0.017

ma function is used to produce the bivariate regression line (bvf) and the bivariate regression forced through the origin (bvfo) in this work. In the work of Isobe et al. (1990) the ordinary least-squares regression, major axis and reduced major axis regression, and additionally ols bisector (ols-bis) regression, are compared. They point out that different slopes are to be expected for all the bivariate fits (ma, sma, ols-bis). For ma they find large uncertainties for the slope. To carry out a symmetrical analysis they recommend using the ols-bis regression.

Figure 10 shows the residual slopes between $R_{\text{PTR,CL}}^{\text{VOC,5}}$ and $R_{\rm PTR,CL}^{\rm VOC,60}$ for consecutive 5 min intervals of the ClearfLo data using the different regression methods (ols, ols-bis, bvfo, bvf). The mean of the residual ratios (i.e. the average ratio minus 1) of $R_{\text{PTR,CL}}^{\text{VOC},5}$ to $R_{\text{PTR,CL}}^{\text{VOC},60}$ is also shown in Fig. 10. The byfo puts more weight onto low OH reactivity values compared to byf and produces a line that much better matches the majority of the data. Therefore smaller residuals are observed compared to the bvf. The very small residual of the average ratio also emphasise that deviation from the ideal slope of 1 is mainly driven by outliers. The ols-bis regression shows a negative residual for all 5 min intervals. Mean deviations and ranges for all regression methods based on consecutive 5 min averaging periods are summarised in Table 8, where it can clearly be seen that ols and the ratio have a negligible deviation once averaged across 12 intervals. On average the ols shows the smallest deviation from the ideal slope of 1, but in terms of stability across all 5 min intervals the ols-bis performs better. This analysis shows that the extent of underor over-predicting OH reactivity by short sampling intervals is a matter of how the data are compared to each other.

The same analysis was performed with an extended data set that included 10 times the number of data points of a randomised log-normal distribution to test for any artefacts relating to the limited sample size of the PTR-ToF-MS data. No appreciable difference was obtained when compared to the smaller data set. Hence, we conclude that the observed bias to an overestimation for the bivariate fits and an underestimation for the ols-bis regression on average is real and not an artefact caused by computing a shorter time series.

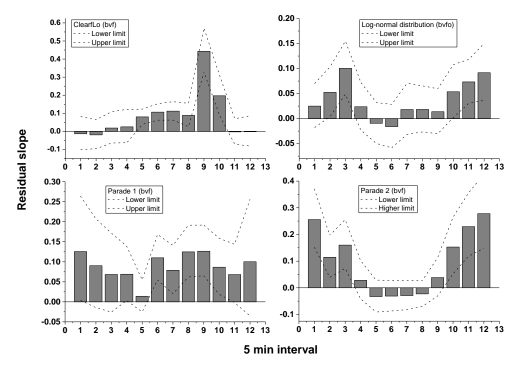


Figure 9. Residual slopes of the correlation of all 5 min means to the hourly mean.

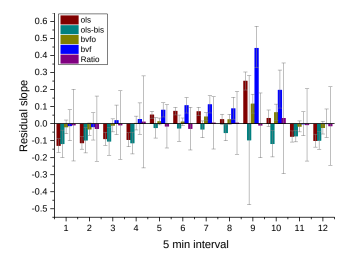


Figure 10. Residual slopes from different linear regression methods and the mean residual ratio for all 5 min intervals. Error bars depict the standard error of the slope for the ols fit, the square root of the variance for the ols-bis, and the lower and upper limit for the bvfo and bvf fits and the standard deviation of the ratios.

3.3 At what sampling interval can the hourly mean be represented with a smaller subsample?

The question being further investigated here is: how many data points are needed to calculate an average value that represents the hourly mean within its standard deviation? The ClearfLo data set of OH reactivity, based on acetone, toluene and xylene, was used to calculate 60 min means of consecutive 1 min data. Small gaps in the time series were skipped such that 60 contiguous data points were computed. However, data was discarded if it included larger gaps, e.g. 1h or more. The set of 60 data points was further subdivided into smaller intervals to calculate means of OH reactivity $R_{\mathrm{PTR,CL}}^{\mathrm{VOC},t\,<60}$ of 2, 3, 5, 10, 15, 20 and 30 min. Residual reactivities for these averages were calculated by subtracting the hourly mean $R_{\text{PTR,CL}}^{\text{VOC,60}}$ before being plotted against the number of data points n, which in this case corresponds to minutes (Fig. 11). Corresponding standard deviations were calculated for each 60 min mean, but only the minimum and maximum values are plotted in Fig. 11 (dashed and solid grey lines, respectively). Additionally, two models are plotted, describing the course of the functions $f_1(1/n)$ (light blue) and f_2 $(1/\sqrt{n})$ (dark blue) starting at the maximum and minimum value (both marked as red dots). The positive range of residual OH reactivity is much wider than the negative range and is capped by the $1/\sqrt{n}$ function. The negative values show a slower approach to the mean. The 20 min averages all lie within the maximum standard deviation, but even when averaging over 30 min, the range is much wider than the minimum standard deviation of OH reactivity.

The 2, 3, 5, 10, 20 and 30 min averages are now compared directly to their hourly mean and standard deviation to summarise the findings from Fig. 11. As can be seen in Table 9 at 20 min, 2.78 % of the ClearfLo data still exceed their hourly mean. At 30 min all data lie within the range of the standard deviation. Therefore, a sampling time greater than 20 min

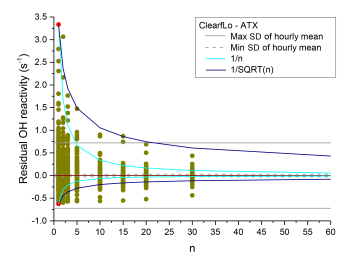


Figure 11. Dependency of the deviation in OH reactivity from the hourly mean on the number of data points for the entire ClearfLo data set.

would be required to represent the hourly mean. The random data reach a comparable level of data exceeding the hourly mean by 2.80 % for averaging over 5 min only. Here, sampling for only 10 min would be sufficient for representing an hour's worth of data. The required sampling times mentioned here correspond to the VOC variability in the analysed data sets. Likewise, longer sampling times could be necessary for representing hourly OH reactivity in other environments such as measurements closer to industrial sources. For example, Gilman et al. (2009) have shown that a much broader range of OH reactivity of VOCs with a high degree of variability can be found in the proximity of heavily industrialised areas like the Houston and Galveston Bay area in Texas, USA.

3.4 Effect of different VOC classes on OH reactivity

Many different atmospheric VOCs have been identified (Goldstein and Galbally, 2007), all of which contribute to OH reactivity. Based on their chemical characteristics they are often divided into different classes. In order to identify how the variation in individual components contributes to the observed deviation of $R_{\rm PTR}^{\rm VOC,5}$ from $R_{\rm PTR}^{\rm VOC,60}$, correlations between 5 min and hourly mean reactivities were analysed for different VOC classes separately. The results are shown in Fig. 12 for ClearfLo (blue area) and PARADE (grey and green areas), where OVOC contains the data from acetone for ClearfLo $(R_{\text{PTR,CL}}^{\text{OVOC},5})$ and acetone and methanol for PARADE $(R_{\text{PTR,PAR}}^{\text{OVOC,5}})$. The aromatics are calculated from toluene and xylene and BVOC refers to the sum of the monoterpenes, which were only available for PARADE. Again, a greater deviation from 1 is observed for the PARADE data. The OVOCs show no significant deviation from 1 for both campaigns, and while the aromatics are close to 1 for ClearfLo, they show a significantly different value for PARADE with a

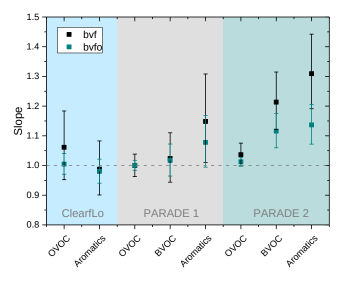


Figure 12. Bivariate fit results between 5 min averaged to 60 min averaged reactivity. Slopes are plotted for ClearfLo (blue shaded area, left) and PARADE (Period 1 – grey shaded; Period 2 – green shaded, right). Correlations were analysed separately for OVOCs (acetone for ClearfLo and acetone and methanol for PARADE), BVOCs (monoterpenes) and aromatic compounds (toluene and xy-lene). The error bars indicate lower and upper limits of the fitted slopes.

deviation of up to 31 %. Finally, BVOCs deviate from a perfect correlation by 21 % for the second period of PARADE.

These results are in line with observations from Williams et al. (2000), who investigated the variability–lifetime relationship of VOCs measured in an unpolluted region of Surinam based on the standard deviation of the natural logarithm of their concentration. They found a higher variability for toluene compared to acetone and methanol. Compounds with a lifetime below 2 days did not seem to fit into this relationship.

3.5 Scaling the effect to the share of OH reactivity of VOCs during ClearfLo

The observed deviations of the slopes from the ideal slope of 1 in Fig. 12 for ClearfLo were scaled by their share to determine the overall effect on total OH reactivity of VOC R_{CL}^{TVOC} . Data from the same week as the PTR-ToF-MS data were used to calculate the influence of VOC speciation on OH reactivity. Over the period of 1 to 7 February 2012 the total OH reactivity of these compounds is $R_{GC,CL}^{TVOC} = 4.05 \text{ s}^{-1}$. Based on Table 1, OVOCs contribute most to reactivity at 43 %, followed by alkenes at 26 % and alkanes at 21 % of $R_{GC,CL}^{TVOC}$. The aromatic compounds have a share of 6 %, and dienes, including isoprene, account for 3 %. Finally, the contribution of the only measured alkyne is less than 1 %.

The extent of which different VOC classes' variability effects $R_{GC,CL}^{TVOC}$ was calculated by weighting the deviation de-

Table 9. Comparison to the hourly averages and their standard deviation for ClearfLo and a data set of log-normal distributed randomised numbers. Listed are the number and the percentage of data that exceed the SD of the hourly mean for different *n*. *n* refers to the number of minutes that were averaged in each case.

		ClearfLo		Randomi		
n	No. data	No. data > SD	% of data > SD	No. data	No. data > SD	% of data > SD
2	3960	838	21.16	4020	534	13.28
3	2640	457	17.31	2680	199	7.43
5	1584	225	14.20	1608	45	2.80
10	792	80	10.10	804	0	0
15	528	38	7.20	536	0	0
20	396	11	2.78	402	0	0
30	264	0	0	268	0	0

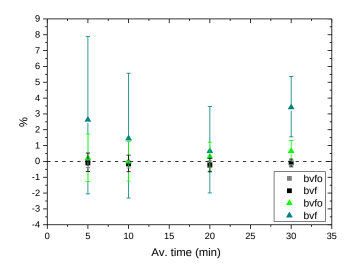


Figure 13. Percentage deviation in OH reactivity for different sampling intervals owing to VOC variability for the ClearfLo data. Results are plotted separately for aromatic compounds (black and grey squares) and OVOCs (green and teal triangles) for both bivariate fits without (bvfo) and with an intercept (bvf). The deviations are based on the share of the VOC's class to total OH reactivity for each investigated averaging interval. The error bars indicate lower and upper limits of the fitted slopes.

rived from the correlations for the different classes (i.e. the deviation of the slope between $R_{\rm PTR,CL}^{\rm class,5}$ and $R_{\rm PTR,CL}^{\rm class,60}$ from 1) by the proportion that each class contributes to the total reactivity (calculated from Table 1). Here, it is assumed that deviations derived from measurements of only a few compounds is representative of each class of VOCs under study.

Based on Fig. 13, 5 min averages over-predict OH reactivity by up to 2.6 % due to variability in OVOC concentrations. This value decreases for increasing averaging time but shows a maximum of 3.4 % for the 30 min mean. There is no significant contribution of the aromatic compounds to a deviation from the hourly mean OH reactivity for any averaging interval. A similar behaviour could be expected for other classes of VOCs such as the alkenes and alkanes, the second and third most important classes in Table 1. However, this could not be tested in the present study using PTR-ToF-MS data. However, this study shows how the effect of using short sampling intervals could account for a missing or over-predicted OH reactivity in the range of 10 % or more.

Lidster et al. (2014) investigated the potential increase in OH reactivity owing to higher substituted aromatic compounds, which are normally not measured in field campaigns. They state that they can contribute to up to 0.9 s^{-1} in OH reactivity. This would increase the share of aromatic compounds by more than a factor of 3; however, based on the results in Fig. 13, the effect on OH reactivity would still be in the range of less than 1 %, while the contribution of the OVOCs would only be altered slightly.

4 Conclusions

The effect of using short sampling intervals for VOC measurements on resulting OH reactivity was investigated using two different monitoring campaigns as case studies. OH reactivity was found to be both under- and over-predicted due to missing variability in VOC data. The divergence between OH reactivity calculated from 5 min sampling intervals and hourly values was found to be around 2-26% and 0-44% for the PARADE and CleafLo campaigns, respectively, owing to the variability in the VOC concentrations. These discrepancies may contribute to missing OH reactivity when compared to direct measurements. Results from the urban and the semirural site show on average similar effects when comparing reactivity averaged over 5 min intervals to the hourly mean.

Comparison to a randomised data set with a similar distribution as the CleafLo data showed that the variability in the VOC concentrations with time is the main reason for deviating results from shorter sampling intervals. For the randomised data a sampling time of less then 10 min is sufficient so that all data points are within the range of the hourly standard deviation, while for the ClearfLo data it takes more than 20 min.

The effect of short sampling times of VOC concentrations on calculated OH reactivity is differently pronounced for each VOC class. When comparing OH reactivity calculated from VOCs sampled over a 5 min period to the hourly mean, a larger divergence was found for the OVOCs than aromatic compounds during ClearfLo. The opposite trend was observed for the PARADE campaign, while the effect of OVOCs is almost negligible. Biogenic VOCs, with the monoterpenes as representatives, were added for analysis. They show a similar behaviour to the OVOCs, but with a slightly greater divergence.

The bigger proportion of measured OVOCs, compared to the aromatic compounds, at the urban site during ClearfLo contributes to a higher deviation in calculated OH reactivity when using short sampling intervals. Taking the results from Lidster et al. (2014) into account, the effect of aromatic VOCs increases but is still small.

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