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Ambient measurements of aromatic and oxidized VOCs by PTR-MS and GC-MS: intercomparison between four instruments in a boreal forest in Finland

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Abstract. Proton transfer reaction mass spectrometry (PTR-MS) and gas chromatography mass spectrometry GC-MS are commonly used methods for automated in situ measurements of various volatile organic compounds (VOCs) in the atmosphere. In order to investigate the reliability of such measurements, we operated four automated analyzers using their normal field measurement protocol side by side at a boreal forest site. We measured methanol, acetaldehyde, acetone, benzene and toluene by two PTR-MS and two GC-MS instruments. The measurements were conducted in southern Finland between 13 April and 14 May 2012. This paper presents correlations and biases between the concentrations measured using the four instruments. A very good correlation was found for benzene and acetone measurements between all instruments (the mean $R$ value was 0.88 for both compounds), while for acetaldehyde and toluene the correlation was weaker (with a mean $R$ value of 0.50 and 0.62, respectively). For some compounds, notably for methanol, there were considerable systematic differences in the mixing ratios measured by the different instruments, despite the very good correlation between the instruments (mean $R = 0.90$). The systematic difference manifests as a difference in the linear regression slope between measurements conducted between instruments, rather than as an offset. This mismatch indicates that the systematic uncertainty in the sensitivity of a given instrument can lead to an uncertainty of 50–100 % in the methanol emissions measured by commonly used methods.

1 Introduction

Volatile organic compounds (VOCs) play a crucial role in atmospheric chemistry (Goldstein and Galbally, 2007; Helmig et al., 2014). They participate in tropospheric ozone production (Atkinson and Arey, 1998, 2003), contribute to aerosol particle formation and growth (e.g. Kulmala et al., 2001; Birmili et al., 2003; Tunved et al., 2006; Paasonen et al., 2013; Riipinen et al., 2012; Patoulias et al., 2014), and also affect the oxidation capacity of the atmosphere (e.g. Lelieveld et al., 2008; Rohrer et al., 2014). The large majority of the atmospheric VOCs originate from biogenic sources, but anthropogenic emissions also contribute significantly (Picco et al., 1992; Guenther et al., 1995, 2012).

In remote and rural locations, biogenic compounds such as isoprene or monoterpenes dominate the concentrations among reactive VOCs. Oxygenated VOCs (OVOCs) are also significant (Guenther et al., 1995, 2012). In urban air, aromatic and oxygenated VOCs make a notable contribution to the total VOC concentrations (e.g. Hellén et al., 2003, 2006; Filella and Peñuelas, 2006; Patokoski et al., 2014). Many of the oxygenated and aromatic VOCs emitted by anthropo-
pogenic sources have long atmospheric lifetimes (from a few days to several weeks) and can be transported thousands of kilometres, making them capable of affecting atmospheric concentrations also in remote locations. These compounds may also have local anthropogenic sources, such as wood combustion or traffic (Hellén et al., 2008; Patokoski et al., 2014).

A variety of models are used to study the atmospheric processes of VOCs. Some simulate the VOC emissions from vegetation (e.g. Grote and Niinemets, 2008; Smolander et al., 2014), others simulate the degradation of VOCs due to their chemical reactions with e.g. atmospheric oxidants (e.g. Jenkin et al., 1997; Apel et al., 2010), and others model their role in new particle formation and other boundary layer and tropospheric processes (e.g. Fast et al., 2006; Holzinger et al., 2007; Makkonen et al., 2012). Many of these models involve dozens of chemical species (including VOCs and trace gases) and complicated chemical and physical processes. Evaluation of these models requires accurate measurements of atmospheric properties, including the VOC concentrations.

Often, VOC concentrations have been measured by collecting samples into canisters or onto adsorbents with subsequent off-line analysis with gas chromatography mass spectrometry (GC-MS) or gas chromatography connected to a flame ionization detector (e.g. Grosjean et al., 1998; Na and Kim, 2001; Hakola et al., 2009; Sauvage et al., 2009). In addition, automated measurements based on both GC techniques and chemical ionization techniques have been developed and utilized (e.g. Lewis et al., 1997; Lindinger et al., 1998; Hakola et al., 2012). Compared to the off-line adsorbent methods, the in situ GC techniques often require only one concentration step, therefore lowering both the background and the detection limits. In addition to lower detection limits, interferences from transport and storage of samples can be avoided.

For automated VOC concentration measurements, in situ GC-MS, proton transfer reaction mass spectrometer (PTR-MS) and other chemical ionization-mass spectrometers (CIMS) have been used (e.g. Lindinger et al., 1998; Munsan, 2000; de Gouw and Warneke, 2007; Moller et al., 2008; Lopez-Hilfiker et al., 2014). Of these methods, the PTR-MS provides very high time resolution (up to a few Hz) and is capable of measuring a wide range of compounds. However, it cannot separate isobaric compounds. GC-MS, on the other hand, can be highly specific for compound identification, but it has lower time resolution (typically 30 min or more). Both of these methods have been used for measurements in different environments, ranging from highly polluted urban areas to remote locations with low VOC concentrations (e.g. Karl et al., 2003; Rinne et al., 2005; Jordan et al., 2009; Holst et al., 2010; Molina et al., 2010; Hellén et al., 2012b; Hakola et al., 2012).

Typically, a long-term measurement set-up consists of a single analyzer, which is regularly calibrated. Occasionally these instruments are compared with each other either in the laboratory or in the field. The laboratory comparisons are usually conducted by measuring VOC concentrations of a known standard mixture (see e.g. Apel et al., 1994, 1999, 2003, 2008; Slmer et al., 2002; Plass-Dülmer et al., 2006; Rappenglück et al., 2006; Hoerger et al., 2014). However, certain problems in the field measurements cannot be easily identified when only one instrument is used. Therefore, it is important to compare the performance of different real-time instruments in field conditions as well. So far, few studies on field comparison of in situ VOC measurements with PTR-MS and GC-MS have been published (de Gouw et al., 2003b, 2004; Kaser et al., 2013; Warneke et al., 2015). de Gouw et al. (2004) studied the correlation between two PTR-MS instruments, Kaser et al. (2013) and Warneke et al. (2015) the correlation between a quadrupole PTR-MS and a PTR-ToF-MS (PTR-MS with a time-of-flight mass spectrometer) and de Gouw et al. (2003b) the correlation between PTR-MS and GC-MS. Such comparison experiments have not been conducted before in high latitude boreal forest, where the anthropogenic influence on the concentrations is rather small.

The main aim of this study was to evaluate how reliable the in situ measurements of aromatic and oxygenated VOCs are when a single stand-alone instrument is used. This was achieved by comparing VOC concentration measurements of four in situ instruments: two PTR-MSs and two GC-MSs. In order to have a comparison as realistic as possible, no data were rejected from the analysis based on bad correlation of the data between the instruments. Thus, only external reasons have been used to filter the data prior the analysis. This study was part of ACTRIS (Aerosols, Clouds and Tracer gases research InfraStructure Network, www.actris.eu, sited on 20 October 2015), which aims to harmonize the European trace gas measurements and to establish a reliable network of continuous long-term measurements. The concentration measurements of three oxygenated VOCs (methanol, acetaldehyde and acetone) and two aromatic VOCs (benzene and toluene) were compared in this study.

2 Methods

2.1 Measurement site

The measurements were conducted between 13 April and 14 May 2012 at the SMEAR II site (Station for Measuring Forest Ecosystem-Atmosphere Relations, 61°51′ N, 24°17′ E, 181 m a.s.l.) in Hyytiälä, southern Finland. The site is a well-characterized measurement station located in a rural boreal forest dominated by Scots pine (Pinus sylvestris) (for details see Hari and Kulmala, 2005; Ilvesniemi et al., 2010). In addition to Scots pine, there are some Norway spruce (Picea abies) and broadleaved trees such as European aspen (Populus tremula) and birches (Betula sp.) at the site. The annual mean temperature at the site is 3°C, with the coldest month being January (mean −9°C) and the warmest July
The annual mean precipitation is 700 mm. The nearest village (Korkeakoski) is about 6 km away and the nearest big city (Tampere, ca. 200,000 inhabitants) is about 50 km from the site.

The concentrations and sources of oxidized and aromatic VOCs at the site have previously been characterized by Rinne et al. (2005, 2007); Patokoski et al. (2014) and Rantala et al. (2014). Oxidized and aromatic VOCs arrive at the SMEAR II station from both long range and local anthropogenic sources (Liao et al., 2011; Patokoski et al., 2014). OVOCs are also emitted by the vegetation at the site and formed in the oxidation of, for example, monoterpenes (Rinne et al., 2005, 2007; Aaltonen et al., 2013; Aalto et al., 2014; Rantala et al., 2014).

2.2 The measurement set-up

The concentrations were measured with two different gas chromatography mass spectrometers (GC-MS1 and GC-MS2), and two similar proton transfer reaction quadrupole mass spectrometers, which are hereafter called PTR-MS1 and PTR-MS2. Both PTR-MS instruments were operated by the University of Helsinki, the GC-MS1 was operated by Empa (Switzerland) and the GC-MS2 was operated by the Finnish Meteorological Institute. The two GC-MS instruments and the PTR-MS1 used the same ca. 20 m long inlet line (Teflon PTFE, 8 mm id), which sampled 10 m above the ground with a sample air flow of 20 L min$^{-1}$ (Fig. 1).

The PTR-MS2 is part of the permanent instrumentation of the site and sampled air from a tower about 30 m away from the common inlet of the other instruments. It measured the ambient atmospheric concentrations during every third hour, as the instrument was used for other measurements during the other 2 h (Aalto et al., 2014; Rantala et al., 2014). The measurements were obtained at six heights (4.2, 8.4, 16.7, 33.6, 50.4 and 67.2 m) for 1 min at a time at each height. One sampling cycle (MID cycle) lasted for 55 s, during which 27 different compounds were sampled sequentially each with a 2 s dwell time. Additionally, the primary ion, the first water cluster, oxygen (which is used as a marker for impurities inside the instrument) and mass $m/z$ 25 (indicator for detector noise; dark counts) were sampled with dwell times of 0.2 s. The remaining 5 s were used for switching to the next height. Thus each compound was effectively sampled for $9 \times 2$ s during 1 h. The switching was performed close to the PTR-MS with a solenoid valve (chamber volume of 11 µL, FKM diaphragm). From each measurement height, the sample air was drawn into the PTR-MS2 via a heated (few degrees above the ambient temperature) 100 m long inlet line with a constant flow of 45 L min$^{-1}$ (Teflon PTFE, 14 mm id). In this comparison study, only the measurements taken at 8.4 m were used. Previously, Kolari et al. (2009) have tested the VOC line losses with a 50 m long PTFE inlet tubing. The losses were found to be only few percentages, although the inlet flow was only 2 L min$^{-1}$. Therefore, the inlet losses of the PTR-MS2 were assumed to be negligible.

Each instrument measured more than 20 different compounds. However, only methanol, acetaldehyde, acetone, benzene and toluene were measured with both PTR-MS instruments and at least with one GC-MS. As such, they were selected for the comparison study.

2.3 PTR-MS

The proton transfer reaction is a chemical ionization technique in which VOCs are ionized by proton transfer from hydronium ions ($\text{H}_3\text{O}^+$). In the PTR-MS (Ionicon Analytik
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GmbH, Austria), the sample air is drawn continuously, without any pretreatment, through the drift tube, where the VOCs of the sample air collide with the H$_3$O$^+$ ions and get ionized if their proton affinity is higher than that of water. From the drift tube the ions are guided to a quadrupole mass spectrometer for mass selection and are then detected by a secondary electron multiplier.

The VOCs gain one proton (H$^+$) in the proton transfer reaction, thus their mass increases by one atom mass unit (amu) and they are singly charged. As PTR-MS has a mass resolution of one Thomson (Th, i.e. mass-to-charge-ratio), different compounds with the same nominal mass cannot be distinguished. Therefore it is not used for exact identification of the measured compounds (for more details about the instrument, see Lindinger et al., 1998; de Gouw et al., 2003a; de Gouw and Warneke, 2007). However, the selectivity of the instrument can be enhanced by varying the E/N (Maleknia et al., 2007; Misztal et al., 2012).

Hydronium primary ions may become hydrated and thus form H$_3$O$^{+\,\cdot}$ (H$_2$O)$_n$ cluster ions in the drift tube. Cluster formation and fragmentation of the molecules are minimized by applying a suitable electric field $E$, or rather $E/N$ (where $N$ is the density of the buffer gas i.e. air), over the length of the drift tube. As a compromise between minimizing the formation of water clusters and inhibiting the fragmentation of the product ions, $E/N$ values from 110 to 140 Townsend (Td) are typically used. Even with an optimized $E/N$ ratio, a considerable amount of H$_3$O$^{+\,\cdot}$H$_2$O clusters are always present in the drift tube when measuring ambient air. Therefore, the measured signals were normalized to 10$^6$ primary ion counts per second using both H$_3$O$^+$ and H$_3$O$^+$H$_2$O signals. As benzene and toluene react slowly with the H$_3$O$^+$H$_2$O, only H$_3$O$^+$ ions were considered for the normalization of these two compounds. Heavier water clusters were not taken into account (Warneke et al., 1996, 2001; Tani et al., 2003; de Gouw and Warneke, 2007; Taipale et al., 2008).

The drift tube pressures and voltages of the two PTR-MS instruments were not the same during this experiment, as the instrument parameters were optimized individually. PTR-MS1, which is the newest of the two PTR-MS instruments, was set to have a drift tube pressure of 2.2 mbar and voltage of 600 V, while PTR-MS2 was run with a drift pressure and voltage of 2.0 mbar and of 510 V, respectively. Therefore the $E/N$ of PTR-MS1 was 136 Td, and that of PTR-MS2 was 120 Td. Possible changes in the drift tube pressures were taken into account by normalizing the drift pressures to 2 mbar. Exactly the same procedures were applied for the calibration and volume mixing ratio (VMR) calculation for both PTR-MS instruments. Each measured compound was sampled with a 2 s dwell time, while the dwell time of the H$_3$O$^+$ and H$_3$O$^+$H$_2$O was 0.2 s in case of both PTR-MS instruments.

A thorough description of the VMR calculation procedure is given by Taipale et al. (2008). Thus, only a brief summary is given here. To convert the measured counts per second (cps) signals to concentrations (ppb, parts per billion, 10$^{-9}$), one needs to know (a) the signals of the primary ions H$_3$O$^+$ and H$_3$O$^+$H$_2$O, (b) the instrumental background signals and (c) the sensitivities for all the measured compounds.

a. The signals of primary ions are measured during each measurement cycle, but in order to maximize the lifetime of the detector, the count rates of the primary ions were measured at the m/z 21 and m/z 39, corresponding to the isotopes, H$_3^1$O$^+$ and H$_3^{18}$O$^+$ (H$_2$O), respectively. The abundance of the primary ions is not constant, but can vary substantially over the course of a few days and sometimes even within hours. This variation is taken into account by normalizing the VOC count rates and sensitivities with pre-averaged (5 min running mean) primary ion signal/signals (Taipale et al., 2008).

b. Instrumental background signal can be caused by, for example, desorption of impurities inside the instrument or inside the inlet system, manifesting as a notable offset in the count rates of many of the VOCs (Steinbacher et al., 2004). These background signals are taken into account, by regular measurement of VOC free air (hereafter “zero air”). The background signals are then subtracted from the measured signals. During this campaign, zero air was measured every second hour with PTR-MS1 and every third hour with PTR-MS2. The zero air, was produced by pumping ambient air through a catalytic converter (Parker Balston zero air generator HPZA-3500, USA and Parker ChromGas Zero Air Generator 3501, USA).

c. In order to quantify the sensitivities for the measured compounds, calibrations were done using an automatic calibration method using mixing units. These mixing units dilute a standard gas flow of ca. 6 mL min$^{-1}$ to a zero air flow of ca. 1000 mL min$^{-1}$. Both instruments had their own identical mixing units, each of which consists of a 1 L (40 bar) standard gas cylinder and two mass flow controllers, which regulate the standard gas and the zero air flow (Bürkert 8710-10, and Bürkert 8710-03, Bürkert GmbH Germany, respectively) automatically. Previously Taipale et al. (2008) calibrated the instrument by diluting 50–120 mL min$^{-1}$ of standard gas to 1000–3000 mL min$^{-1}$ of zero air, which was done with a set-up that uses a 60 L standard gas bottle (with an initial pressure of 140 bar). The flow is regulated manually with a pressure regulator and is fine-tuned with a needle valve. Hereafter this calibration method is referred to as “manual calibration”. The comparability of the manual and the automatic calibration methods was studied in separate calibration method comparison tests, which were performed after the campaign for both PTR-MS instruments.

Both instruments were calibrated three times during the campaign, using the same gas standard mixture (Apel-
Riemer Environmental Inc., CO, USA), consisting of 13 different VOCs including methanol, acetaldehyde, acetone, benzene and toluene in the range of 0.84–1.14 ppb.

The detection limits of the PTR-MS instruments were calculated as 3 times the SD (3σ) of the background measurement. This background signal varies over time, leading to changes in the detection limits. Possible changes in background signals were taken into account by calculating detection limits separately for all calibration periods of the PTR-MS instruments.

2.4 GC-MS instruments

The analysis of VOCs with gas chromatographic techniques relies on the separation of the VOC species in a chromatographic column. Traditionally, the samples have been collected into a canister or adsorbent tubes and analyzed off-line in the laboratory. With more recent in situ GC-MS systems, the samples are collected directly into adsorbent traps at the measurements site, from which they are desorbed by heating the trap in the gas chromatograph. After separating the compounds by their retention times in the chromatographic columns, they are ionized by electron ionization and detected individually with a quadrupole mass spectrometer.

2.4.1 GC-MS1

The instrumental set-up of the adsorption–desorption system coupled to a gas chromatograph–mass spectrometer (GC-MS1) is described in detail by Legreid et al. (2007, 2008). Briefly, every 46 min a 12 min air sample of 350 mL was collected. VOCs were collected into a two-stage adsorbent system connected to a GC-MS (Agilent 6890-5973N, Agilent Technologies, CA, USA). The water removal was performed on the sampling trap (0.6 g of Hayesep D, Supelco, Switzerland) at room temperature. The hydrophobic nature of the adsorbent material allowed most of the water to pass through the trap, and the remaining humidity was removed by dry helium flushing. Thereafter, the compounds were refocused on a microtrap (127 µL), which was flushed with helium into the adsorbent trap.

Methanol was only recovered at 45%, and this was corrected for the measurement campaign. The detection limit for each compound was calculated as 3 times the SD of five zero air samples.

2.4.2 GC-MS2

Measurements of GC-MS2 were conducted using an in situ thermal desorption unit (Markes’ Unity, Markes International Ltd, UK) with a gas chromatograph (Agilent 7890A, Agilent Technologies, CA, USA) and a mass spectrometer (Agilent 5975N, Agilent Technologies, CA, USA). The column used was the 60 m long DB-5 with an inner diameter of 0.235 mm and a film thickness of 1 µm. One 60 min sample was taken every other hour. Ozone was removed by a heated stainless steel inlet (temp. 150°C, length 1 m, o.d. 1.8 inch, flow 0.8 L min⁻¹; Hellén et al., 2012a). Due to the gradually decreasing ozone removal capacity, the ozone removal inlet was changed every second week. VOCs in 30 mL min⁻¹ subsample were collected directly into the microtrap (U-T15ATA-2S, Markes International Ltd, UK) of the thermal desorption unit. Water was removed by keeping the hydrophobic cold trap at 15°C. For calibration, a gaseous standard mixture of 17 VOCs each at a concentration of 2 ppb (National Physical Laboratory, UK) was run after every 50th sample. The limits of detection, calculated as 3 times the SD of the zero levels, were 215 ppt for acetone, 34 ppt for benzene and 9 ppt for toluene.

2.5 Uncertainties of the instruments

The uncertainties of the VOC concentrations measured by a PTR-MS or a GC-MS are affected by several factors. The total uncertainty (ΔU) can be estimated by using the Gaussian propagation of uncertainty if the uncertainties of different steps of the data processing are known. In the following chapters, the total uncertainty calculations of PTR-MS instruments and the GC-MS instruments are described separately. One should keep in mind that, in addition to the total (random) uncertainty described in this chapter, the measurements may still have additional constant error (bias) of unknown magnitude, which can bias the measured concentrations.

2.5.1 Uncertainty of the PTR-MS measurement

The total measurement uncertainty of PTR-MS consists of two parts; the uncertainty of the signal (ΔU_signal) and the uncertainty of the calibration (ΔU_calibration):

\[ ΔU^2 = ΔU_{\text{signal}}^2 + ΔU_{\text{calibration}}^2 \]  

(1)

The measured signal and the background signal are normalized with the primary ion signal for the VMR calculation.
The normalized background signal ($I_{\text{zero}}$) is subtracted from the normalized measured signal ($I_{\text{meas}}$) and this background-corrected normalized signal is divided by the normalized sensitivity, $S$, which is obtained from the calibration. Thus,

$$\text{VMR} = \frac{I_{\text{meas}} - I_{\text{zero}}}{S}. \quad (2)$$

The uncertainty of the signal in Eq. (1) contains the uncertainties of the measured signal ($\Delta U_{\text{meas}}$) and the background signal ($\Delta U_{\text{zero}}$).

$$\Delta U_{\text{signal}}^2 = \Delta U_{\text{meas}}^2 + \Delta U_{\text{zero}}^2. \quad (3)$$

The measured count rates (cps, counts per second) and the count rates of the zero measurement were converted to counts ($I_{\text{counts}}$ and $I_{\text{counts, zero}}$) by multiplying with the dwell time (2 s for each molecule). As the PTR-MS statistics follow the Poisson distribution, the uncertainty of a single measurement point ($\Delta I_{\text{meas}}$) is simply the square root of the counts ($\sqrt{I_{\text{counts}}}$). One background measurement consisted of 11 measurement points, from which the average background signal was derived. The nearest background value was subtracted from each individual ambient measurement point. The uncertainty of one background measurement ($\Delta I_{\text{zero}}$) was calculated as the SD of the 11 measurement points.

In order to normalize $I_{\text{counts}}$ and $I_{\text{counts, zero}}$, they both need to be divided by the primary ion (H$_3$O$^+$ and H$_2$O$^+$H$_2$O) counts, which are obtained by multiplying the count rates of the primary ions by their dwell times. However, the primary ion signal is much higher than the measured signals and the zero signals. In addition, it remained approximately constant during the time when the $I_{\text{counts}}$ and the nearest $I_{\text{counts, zero}}$ were measured. Thus, the primary ion signal uncertainty is less than 1 % and it was thus neglected.

The uncertainty of the calibration ($\Delta U_{\text{cal}}$) is due to the uncertainty of the sensitivity ($\Delta S$) and the uncertainty of the calibration gas standard ($\Delta U_{\text{stdgas}}$) due to uncertainty of the concentrations in the calibration gas standard ($\Delta \chi_{\text{cal}}$):

$$\Delta U_{\text{cal}}^2 = \Delta U_{\text{cal}}^2 + \Delta U_{\text{stdgas}}^2. \quad (4)$$

PTR-MS sensitivity for a certain compound is determined by calibrating the instrument with a known concentration of that compound. When the ratio of the sensitivity and its uncertainty is assumed to be constant, the sensitivity’s uncertainty can be determined from the SD of a series of calibrations, performed using the same instrument settings. The laboratory tests for the similarity of the two calibration methods were done under the same instrument conditions, making the relative sensitivity uncertainty ($\Delta S$) obtainable from those measurements. The manufacturer of the calibration gas standard reports relative uncertainty ($\Delta \chi_{\text{cal}}$), of $\pm 5 \%$ for the concentration of each VOC compound in the calibration gas mixture.

By combining Eqs. (1)–(4) and using the Gaussian propagation of error, the total uncertainty of PTR-MS for one measurement point is

$$\Delta U = \sqrt{\left(\frac{\Delta I_{\text{meas}}}{I_{\text{meas}} - I_{\text{zero}}} \times \text{VMR}\right)^2 + \left(\frac{\Delta I_{\text{zero}}}{I_{\text{meas}} - I_{\text{zero}}} \times \text{VMR}\right)^2 + \left(\Delta S \times \text{VMR}\right)^2 + \left(\Delta \chi_{\text{cal}} \times \text{VMR}\right)^2}. \quad (5)$$

For $N$ measurement points, the total relative uncertainty can be calculated as

$$\Delta U_{\text{rel}} = \frac{1}{N \times \text{VMR}} \sqrt{\sum_{i=1}^{N} \left(\frac{\Delta I_{i, \text{meas}}}{I_{i, \text{meas}} - I_{i, \text{zero}}} \times \text{VMR}\right)^2 + \left(\sum_{i=1}^{N} \Delta S \times \text{VMR}\right)^2} + \left(\sum_{i=1}^{N} \Delta \chi_{\text{cal}} \times \text{VMR}\right)^2}, \quad (6)$$

where VMR is the average volume mixing ratio of $N$ measurements. Because different measurement points are independent, the total precision can be calculated using the Gaussian propagation of error. However, as $\Delta S$ and $\Delta \chi_{\text{cal}}$ are constant, the total systematic error is calculated as a linear sum of the errors of single measurement points.

Total uncertainties of 1 h were calculated for PTR-MS1 and PTR-MS2, as the data comparison was mostly done using 1-hour averages.

### 2.5.2 Uncertainty of the GC-MS1 measurement

The total uncertainty is divided into two components: precision ($\Delta U_{\text{precision}}$) and systematic error ($\Delta U_{\text{systematic}}$):

$$\Delta U^2 = \Delta U_{\text{precision}}^2 + \Delta U_{\text{systematic}}^2. \quad (7)$$

The precision is calculated as

$$\Delta U_{\text{prec}} = \frac{1}{3} \times \text{DL} + \chi \times \sigma_{\text{sample, rel}}, \quad (8)$$

where DL is the detection limit, $\chi$ is the mole fraction (peak area) of the considered peak and $\sigma_{\text{sample, rel}}$ is the relative SD of the sample. The first term of Eq. (8) considers the resolution of the instrument (e.g. background noise) and the second term considers the repeatability of the measurements. For low mole fractions the first term dominates, while for high mole fractions the second term dominates.

The systematic error of GC-MS1 includes the following: the error due to uncertainty of the calibration standard’s mole fractions ($\Delta \chi_{\text{cal}}$), systematic integration errors due to peak overlay or poor baseline separation ($\Delta \chi_{\text{int}}$), systematic errors due to blank correction ($\Delta \chi_{\text{blank}}$), and potential further instrument problems ($\Delta \chi_{\text{instrument}}$) caused by, for example,
sampling line artefacts, possible non-linearity of the detector or changes of split flow rates. Hence, the systematic error is

$$\Delta U_{\text{systematic}}^2 = \Delta \chi^2_{\text{cal}} + \Delta \chi^2_{\text{int}} + \Delta \chi^2_{\text{blank}} + \Delta \chi^2_{\text{instrument}}. \quad (9)$$

The systematic error due to the calibration gas uncertainty ($\Delta \chi_{\text{cal}}$) is calculated as

$$\Delta \chi_{\text{cal}} = \frac{A_{\text{sample}} V_{\text{cal}}}{A_{\text{cal}} V_{\text{sample}}} \delta \chi_{\text{cal}}, \quad (10)$$

where $A_{\text{sample}}$ is the peak area of the sample measurement, $A_{\text{cal}}$ the peak area of the calibration standard measurement, $V_{\text{sample}}$ the volume of sample, $V_{\text{cal}}$ the sample volume of the calibration standard, and $\delta \chi_{\text{cal}}$ certified standard uncertainty of calibration standard and potential drift of the calibration standard.

The systematic integration error ($\Delta \chi_{\text{int}}$) is

$$\Delta \chi^2_{\text{int}} = \left( \frac{f_{\text{cal}}}{V_{\text{sample}}} \delta A_{\text{sample}} \right)^2 + \left( \frac{A_{\text{sample}} V_{\text{cal}} \chi_{\text{cal}}}{V_{\text{sample}} A_{\text{cal}}^2} \delta A_{\text{cal}} \right)^2, \quad (11)$$

with $f_{\text{cal}} = \frac{V_{\text{cal}} \chi_{\text{cal}}}{A_{\text{cal}}}$, where $\delta A_{\text{cal}}$ is the relative error in peak area due to integration of the calibration measurement, $\delta A_{\text{sample}}$ is the integration error of the sample measurement and $\chi_{\text{cal}}$ is the mole fraction of the calibration standard peak. If a blank correction has to be applied, the error of this correction is described as the deviation from the mean blank value:

$$\Delta \chi_{\text{blank}} = \sigma_{\text{blank}} \frac{1}{\sqrt{N - 1}}, \quad (12)$$

where $\sigma_{\text{blank}}$ is the SD of the zero gas measurements and $N$ is the number of those zero-gas measurements. For more details on the uncertainty calculation of GC-MS1 see Hoerger et al. (2014).

The precision of acetone, acetaldehyde, benzene, and toluene was around 5%, whereas the precision for methanol was 10%. The total expanded uncertainty was around 15% for acetone, benzene, and toluene, 23% for acetaldehyde, and 28% for methanol (Table 1). These values are in good agreement with previous studies (Apel et al., 2008).

### 2.5.3 Uncertainty of the GC-MS2 measurement

Total uncertainty ($\Delta U$) of the GC-MS2 is calculated as

$$\Delta U^2 = \Delta \chi^2_{\text{cal}} + \Delta \chi^2_{\text{blank}} + \Delta \chi^2_{\text{analysis}} + \Delta \chi^2_{\text{flow}}, \quad (13)$$

where $\Delta \chi_{\text{cal}}$ is the uncertainty of the standard preparation, $\Delta \chi_{\text{blank}}$ is the uncertainty of the blank level, $\Delta \chi_{\text{analysis}}$ is the uncertainty of the analysis and $\Delta \chi_{\text{flow}}$ is the uncertainty of the sample flow. Uncertainties of the standard preparation and sample flow were given, respectively, by the manufacturers of the calibration gas standard and the mass flow controller. The blank level uncertainty was calculated as the SD of all blank values measured during the campaign. The uncertainty of the analysis was obtained from the relative SD of the analysis of 15 identical calibration standards. Analytical uncertainties calculated from partial uncertainties at a concentration level of 2 ppb were 17% for acetone, 4% for benzene and 5% for toluene.

### 2.6 Data processing

The concentrations measured with different instruments had temporal discrepancies, as all of the instruments had different sampling intervals and data integration times. PTR-MS1 measured several compounds sequentially, each with a dwell time of 2 s, which lead to a 1 min time-resolution. The ambient concentrations were measured 43 times during each hour, after which the background was sampled 11 times. PTR-MS2 measured ambient concentrations every third hour, during which each of the six measurement heights were sampled every sixth minute. Also, PTR-MS2 measured background during the same hour as the ambient concentrations were measured. Each measurement height was sampled eight times within the 1 h averaging period, followed by 11 background samples. In this analysis, the concentrations measured at 8.4 m height were used. GC-MS1 collected a sample for 12 min, after which the sample was analysed for 34 min. Thus, a concentration value was recorded every 46 min. GC-MS2 sampled for 60 min during every second hour and analysed the sample in between the sampling times.

In order to make the instrument comparison as consistent as possible, the measurements by the two PTR-MS instruments were averaged for the same time periods with each other and with the GC-MS instrument whenever possible. For the comparison between the two PTR-MS instruments and PTR-MS1 and GC-MS2, hourly averages were used. For the comparison between PTR-MS1 and GC-MS1, PTR-MS1 data were averaged for the same 12 min time periods when GC-MS1 samples were taken.

The detection limits of all the instruments were determined as 3 times the SD of the instrument noise (i.e. the zero air sample concentration). Values below the detection limits were removed from the GC-MS data. When hourly or 12 min average values were calculated from PTR-MS data, the averages were calculated for all data points. If an average
The sensitivity uncertainties of both calibration methods were lower for PTR-MS2. Regarding the manual calibration method, the pump used to generate zero air for the calibration of PTR-MS1 caused some fluctuation to the zero air flow and thus increased the sensitivity variation (i.e. the SD) between different calibrations. The uncertainty of sensitivity for methanol obtained with the automatic calibration system was clearly higher than the uncertainties for all other compounds, 63% for PTR-MS1 and 25% for PTR-MS2.

Methanol calibration is difficult due to its strong interaction with metal surfaces, as in mass flow controllers (de Gouw et al., 2003a). Higher apparent methanol sensitivities and sensitivity uncertainties were obtained with the manual calibration method, which contains fewer metal surfaces than the automatic calibration system. It had also been used for a longer time, and the surfaces of the pressure regulator and needle valve were evidently more saturated with methanol than the metal surfaces of the mixing units that were used for the automatic calibration.

In the case of PTR-MS1, the sensitivity uncertainties were higher than the uncertainties of the signal statistics or the concentration uncertainty of the calibration gas standard (Table 2). The signal uncertainty was 1% or less for all compounds for PTR-MS1, while for PTR-MS2 the signal uncertainties were higher, and contributed to the total uncertainty. The higher signal uncertainties of PTR-MS2 were due to the rather low sampling frequency (eight samples per hour) of the PTR-MS2. The signal uncertainty of toluene was particularly high (65%).

### 3.2 Total uncertainties of the concentration measurements

The total uncertainties of all instruments were below 30%, with the exception of the methanol uncertainty of PTR-MS1 and the toluene uncertainty of PTR-MS2 (Table 1). GC-MS2 had low total uncertainties for benzene and toluene concentrations. However, uncertainties of GC-MS2 were determined at a concentration of 2 ppb, which is higher than the concentrations measured for benzene and toluene during
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Figure 2. Concentrations of methanol, acetaldehyde, acetone, benzene and toluene measured with PTR-MS1, PTR-MS2, GC-MS1 and GC-MS2 and ambient temperature and relative humidity during the measurement campaign. Hourly averages were calculated for the PTR-MS instruments. For the GC-MS instruments, all data are shown.

3.3 General features of the ambient data

The time series of methanol, acetaldehyde, acetone, benzene and toluene concentrations measured with all instruments are presented in Fig. 2. All instruments measured similar concentration patterns for methanol, acetone and benzene. Figure 3 also reveals similar patterns in daily median concentrations. However, both figures show systematic differences between the instruments.

The highest concentrations of methanol, acetone and benzene were measured with PTR-MS2, while GC-MS2 measured systematically lower concentrations of acetone and benzene than the other three instruments. In the case of acetone, the lower concentrations measured by GC-MS2 were probably due to the 60 min sampling time. This may have been too long, leading to break through of acetone in the microtrap. Consequently, the absolute values of the GC-MS2 were underestimations, but the concentrations trends were still observed.

The measured acetaldehyde concentrations had fairly small temporal variation. Additionally, the concentration trends measured with the three instruments are divergent until the beginning of May. After the 1 May, PTR-MS1 and GC-MS1 measurements agree rather well.

Toluene concentrations were mostly below the detection limits of the PTR-MS instruments. This is clearly visible in Fig. 2, where the toluene time series of PTR-MS1 and PTR-MS2 consists of fewer points than the time series of the other compounds. The daily median concentrations measured by PTR-MS1, however, show a trend similar to GC-MS1 and GC-MS2.

The different location of the PTR-MS2 inlet could partly explain the higher concentrations observed for methanol, acetaldehyde and acetone. Acetaldehyde and acetone are formed in the oxidation of, for example, monoterpenes and methylbutenol (Kesselmeier et al., 1997; Goldstein and Schade, 2000; Villanueva-Fierro et al., 2004; Millet et al., 2010) and acetaldehyde, acetone and methanol are emitted by the vegetation at the site (Rinne et al., 2007; Aalto et al., 2014; Rantala et al., 2014). The relative contribution of local biogenic sources to the total atmospheric concentrations of methanol and acetone is likely to be low as these compounds have relatively long atmospheric lifetimes (4, 16 and 33 days, respectively, in the spring) and high background concentrations originating from distant sources (Patokoski et al., 2014). As such, their concentrations are likely to have relatively low small-scale spatial variability at the site.
Benzene and toluene observed at the SMEAR II site originate from local traffic and small-scale wood combustion, as well as from distant anthropogenic sources (Hakola et al., 2003; Hellén et al., 2006; Patokoski et al., 2014). Occasional traffic near the measurement site may have caused short pollution events and concentration differences between the two inlets. However, these episodes were rare and their influence on the hourly average values was probably low. There should not be considerable spatial variation in benzene and toluene concentrations at the site, as a majority of both compounds originate from anthropogenic sources outside of the measurement site, and have relatively long atmospheric lifetimes (Hakola et al., 2003; Hellén et al., 2006; Patokoski et al., 2014).

Toluene concentrations were mostly below the detection limits of the PTR-MSs. This is clearly visible in Fig. 2, where the toluene time series of PTR-MS1 and PTR-MS2 consists of fewer points than the time series of the other compounds. The daily median concentrations measured by PTR-MS1, however, show a trend similar to GC-MS1 and GC-MS2.

### 3.4 Differences between the instruments by compound

In order to analyze in more detail how consistent the concentration measurements were, box plots representing the medians and quartiles were drawn for all compounds (Fig. 4). The concentration ranges of different instruments were determined from the box plots. Accordingly, concentration range is hereafter defined as the range between 25 and 75 percentile.

Correlations between different instruments were studied using scatter plots and by calculating Pearson’s correlation coefficients ($R$) between the instruments (Table 3). As PTR-MS2 used a different inlet than the other three instruments did, its measurements were compared only with PTR-MS1.

Additionally, the overall consistency of the concentration measurements of the four different instruments was investigated by calculating: (1) the mean of all correlation coefficients, (2) the root mean square (RMS) difference of the scatterplot slopes from 1 : 1 line, and (3) the RMS of the intercepts for each compound (Table 4). The RMS difference between the slopes and 1 : 1 line ($RMS_{slope}$) was calculated as

$$RMS_{slope} = \sqrt{\frac{1}{N} \sum_{i=1}^{N} (1 - \text{slope}_i)^2},$$

where slope$_i$ is the slope of a scatter plot and $N$ is the number of slopes used for the calculation. In an ideal case, the scatter plot slopes are close to unity, and the RMS$_{slope}$ is close to zero. The slope and intercept values of a scatter plot depend on the positioning of the two data sets on the $x$ and $y$ axes, thus the all slopes and intercepts were calculated for both axis configurations.

Generally, the measurements of PTR-MS2 were most scattered for all the compounds (Fig. 4). This was at least partly due to the discontinuous measurement cycle of the instrument, which meant that fewer data points (8 per hour) were available for calculating the hourly average than were available when using the PTR-MS1 (43 per hour). When fewer data points are used, individual divergent values have larger effects on the average value, as the SD is inversely proportional to the square root of data points. Data from the GC-
MS2, which had the longest sampling time, were least scattered.

In the following sections, the concentration distributions and correlations between different instruments are discussed separately for all five compounds.

3.4.1 Methanol

Methanol was measured with three out of four instruments: PTR-MS1, PTR-MS2 and GC-MS1. There were large differences in the concentration ranges of the methanol measurements (Fig. 4). PTR-MS2 measured the highest concentrations, varying from 2.6 to 5.5 ppb. The measurements of PTR-MS1 and GC-MS1 were less scattered and the ranges were more congruent: 1.7–3.3 and 1.2–2.6 ppb respectively. Also, the median methanol concentration of PTR-MS2 (3.6 ppb) was clearly higher than the median of PTR-MS1 (2.4 ppb), whereas the median concentration measured with GC-MS1 was the lowest (1.6 ppb). It is important to note that the measurement uncertainty of PTR-MS1 was very high for methanol (Table 1).

As Fig. 5 and Table 1 show, the correlation between the two PTR-MS instruments was very good ($R = 0.96$), but the linear regression slope was 1.80. Thus, concentrations measured with PTR-MS2 are almost twice as high as those measured with PTR-MS1. The correlation between PTR-MS1 and GC-MS1 was also good (0.84), but between these two instruments there was a constant offset and the slope was far from one (0.42). The mean correlations and RMS values of the slopes and intercepts are presented in Table 3, which shows that the measured methanol concentrations correlated well but the RMS slope of 0.87 was far from the ideal 1:1 slope.

For methanol the correlation coefficients of this study agreed with those found in prior research. de Gouw et al. (2003b, 2004) and Kaser et al. (2013) reported $R$ values above 0.92 and slope values between 1.03 and 1.16 for PTR-MS vs. GC-MS, PTR-MS vs. PTR-MS and PTR-MS vs. PTR-Tof-MS, respectively. In this study the slopes were clearly less robust than in previous studies, indicating that the time trends of methanol can be captured well with all instruments, but also suggesting that the quantitative concentration values of all three instruments should be regarded with suspicion.

Methanol measurements are known to encounter some challenges. Calibrating PTR-MS for methanol is difficult because methanol deposits on the metal surfaces of the calibration system (de Gouw et al., 2003a), reducing the apparent sensitivity and potentially making the concentrations seem higher than they actually are. However, in our study the systems used to calibrate the two PTR-MS instruments were identical, thus having the same materials. Furthermore, an oxygen isotope ($^{17}$O) is detected with the same mass (33 amu) as methanol in the PTR-MS. However, this is not a problem as it is taken into account in the VMR calculation (Taipale et al., 2008). Apart from the oxygen isotope, no significant interference of any other species has been reported in the literature (de Gouw and Warneke, 2007).

The solubility of methanol in water can introduce problems to the GC-MS measurements, because when water is removed from the sample, part of the methanol could be re-
Figure 5. Comparison of volume mixing ratios of methanol, acetaldehyde, acetone, benzene and toluene measured by four different instruments. PTR-MS1 was compared against all three other instruments and the two GC-MSs were compared to each other. Solid lines represent linear fits and the number in brackets goodness of the fits normalized to the number of data points.
moved as well. To correct for the methanol loss in water trap we used information from two previous intercomparison campaigns. A laboratory intercomparison campaign was conducted in 2005 in Germany, during which OVOCs were measured with several GC-MS instruments at the SAPHIR chamber at Forschungszentrum Jülich (see Apel et al., 2008, for details). During the campaign, the SAPHIR chamber was filled with ambient air and spiked with an unknown number of compounds. The results of the GC-MS1 showed overall good agreement with the other instruments, though a tendency to underestimate the mole fractions in the chamber was observed. For methanol, the loss was around 40 % and it was suspected to occur in the bulk trap during the water removal step. Since the intercomparison in the SAPHIR chamber, the material in the bulk trap of GC-MS1 has aged, and the loss of methanol has increased. During an ACTRIS OVOC intercomparison at Hohenpeissenberg (Germany) in October 2013, the methanol loss was 55 % (unpublished). The methanol concentrations measured during this campaign were corrected for the 55 % loss.

### 3.4.2 Acetaldehyde

Three instruments out of four, PTR-MS1, PTR-MS2 and GC-MS1, measured acetaldehyde. The concentration range was very similar for all the instruments, between 0.3 and 0.6 ppb (Fig. 4). Also, the median concentrations of 0.4, 0.4 and 0.5 ppb for PTR-MS1, PTR-MS2 and GC-MS1 respectively, are within a 25 % range of each other. Despite the similar concentration distributions, the correlation between methanol concentrations measured by PTR-MS1 and GC-MS1 seems to depend slightly on RH, (Fig. 6), while other meteorological parameters had no effect. The reason for the RH dependency is unknown. However, the methanol loss of the GC-MS1 due to water removal could be dependent on RH.

### Table 3. Statistical parameters of the correlation analysis for the measured compounds. $N$ is the number of data points considered in the correlation analysis and $R$ the correlation between two instruments for each compound. Parameters $a$ and $b$ are the slope and the offset of a linear fit, respectively.

<table>
<thead>
<tr>
<th>Compounds</th>
<th>$a$ [ppb]</th>
<th>$b$ [ppb]</th>
<th>$R$</th>
<th>$N$</th>
</tr>
</thead>
<tbody>
<tr>
<td>methanol</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTR-MS1 vs. PTR-MS2</td>
<td>1.80 ± 0.08</td>
<td>-0.20 ± 0.25</td>
<td>0.96</td>
<td>159</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS1</td>
<td>0.52 ± 0.03</td>
<td>0.46 ± 0.10</td>
<td>0.84</td>
<td>392</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GC-MS1 vs. GC-MS2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>acetaldehyde</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTR-MS1 vs. PTR-MS2</td>
<td>0.54 ± 0.21</td>
<td>0.16 ± 0.10</td>
<td>0.37</td>
<td>160</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS1</td>
<td>0.60 ± 0.07</td>
<td>0.25 ± 0.03</td>
<td>0.62</td>
<td>425</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GC-MS1 vs. GC-MS2</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>acetone</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTR-MS1 vs. PTR-MS2</td>
<td>1.25 ± 0.05</td>
<td>-0.04 ± 0.05</td>
<td>0.97</td>
<td>162</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS1</td>
<td>1.03 ± 0.05</td>
<td>0.16 ± 0.05</td>
<td>0.88</td>
<td>423</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS2</td>
<td>0.59 ± 0.04</td>
<td>-0.04 ± 0.04</td>
<td>0.91</td>
<td>206</td>
</tr>
<tr>
<td>GC-MS1 vs. GC-MS2</td>
<td>0.47 ± 0.03</td>
<td>-0.01 ± 0.01</td>
<td>0.77</td>
<td>237</td>
</tr>
<tr>
<td>benzene</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTR-MS1 vs. PTR-MS2</td>
<td>1.38 ± 0.11</td>
<td>-0.01 ± 0.01</td>
<td>0.88</td>
<td>168</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS1</td>
<td>0.99 ± 0.06</td>
<td>0.001 ± 0.005</td>
<td>0.84</td>
<td>449</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS2</td>
<td>0.88 ± 0.06</td>
<td>-0.01 ± 0.01</td>
<td>0.89</td>
<td>213</td>
</tr>
<tr>
<td>GC-MS1 vs. GC-MS2</td>
<td>0.74 ± 0.04</td>
<td>0.005 ± 0.003</td>
<td>0.92</td>
<td>280</td>
</tr>
<tr>
<td>toluene</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PTR-MS1 vs. PTR-MS2</td>
<td>1.36 ± 0.52</td>
<td>-0.04 ± 0.02</td>
<td>0.50</td>
<td>85</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS1</td>
<td>0.55 ± 0.11</td>
<td>-0.01 ± 0.01</td>
<td>0.53</td>
<td>232</td>
</tr>
<tr>
<td>PTR-MS1 vs. GC-MS2</td>
<td>0.92 ± 0.18</td>
<td>-0.01 ± 0.01</td>
<td>0.69</td>
<td>118</td>
</tr>
<tr>
<td>GC-MS1 vs. GC-MS2</td>
<td>0.60 ± 0.07</td>
<td>0.006 ± 0.004</td>
<td>0.77</td>
<td>182</td>
</tr>
</tbody>
</table>
The correlations of this study were weaker than the correlation reported by de Gouw et al. (2003b) (PTR-MS vs. GC-MS) and Kaser et al. (2013) (PTR-MS vs. PTR-ToF-MS). The slopes of both this study and the study by de Gouw et al. (2003b) are equally far from one, while Kaser et al. (2013) reported a slope close to one. However, in the study by Kaser et al. (2013) the concentration range was much higher than in our study, up to 3.5 ppb.

Temperature and wind properties did not explain any of the concentration divergences between different instruments. Instead, concentration difference of both instrument pairs was observed to depend on RH. This phenomenon was clearly stronger between PTR-MS1 and PTR-MS2 than between PTR-MS1 and GC-MS1 (Fig. 7). RH has a diurnal cycle but if there were diurnal cycle in the concentration differences this would be seen also as temperature dependence of the concentration difference. However, no such dependence was observed. The reason for the humidity dependence of the acetaldehyde concentration difference as measured by the two PTR-MS instruments remain unknown for now.

PTR-MS measures acetaldehyde with a mass of 45 amu, but in air masses that are strongly influenced by biogenic emissions, several other compounds with the same mass (isomers) exist (de Gouw et al., 2003a). However, in this study the wind direction did not have an effect on the correlations between the different instrument pairs. Furthermore, de Gouw et al. (2003a) have reported that the acetaldehyde concentration in the calibration gas may decrease over time, which again would lead to an overestimated concentration. The calibration gas standard used in this study was less than 1-year old during the measurement campaign, thus the acetaldehyde concentration in the calibration gas was probably not considerably decreased.

### 3.4.3 Acetone

Acetone concentrations were measured with all four instruments. GC-MS1 and PTR-MS2 measured similar acetone concentrations, ranging from 0.9 to 1.3 ppb, whereas the range of PTR-MS1 was slightly lower, between 0.8 and 1.1 ppb. The lowest concentrations were measured with GC-MS2, 0.4–0.6 ppb. The median concentrations of PTR-MS1 (0.9 ppb), PTR-MS2 (1.0 ppb) and GC-MS1 (1.1 ppb) were within 20 %, while the median for GC-MS2 was clearly lower (0.5 ppb).

As in previous comparison studies (de Gouw et al., 2003b, 2004; Kaser et al., 2013; Warneke et al., 2015), acetone measurements correlated well in this study. The best correlation coefficient was between the two PTR-MS instruments (0.97). PTR-MS1 also correlated well with both GC-MS1 (0.88) and GC-MS2 (0.91). The different sampling times of the two GC-MS instruments could cause at least part of their lower correlation (0.77), as acetone concentration can vary within 1 h. Furthermore, the slope for PTR-MS1 against GC-MS1 was very good (1.03). However, the intercept was 0.2 ppb, indicating a difference in the background levels of acetone for these two instruments. The slope between PTR-MS1 and PTR-MS2 was rather good (1.25). The slopes between GC-MS2 and both PTR-MS1 and GC-MS1 were far from unity, 0.56 and 0.47, respectively. This was probably due to the long sampling time, causing acetone to break through the micro trap. Consequently, even though GC-MS2 measured the time trends of acetone equally well as the other instruments, it underestimated the quantitative concentrations. The average correlation coefficient for acetone was good (0.88), but the low slope values of GC-MS2 plotted against both PTR-MS1 and GC-MS1 were far from unity (Fig. 5), also increased the RMS_slope (0.54). When the RMS_slope was calculated only for PTR-MS1 vs. PTR-MS2 and for PTR-MS1 vs. GC-MS1 pairs, it is very close to zero (0.02). Meteorological parameters could not explain the small differences between different instruments.
PTR-MS measurements of acetone can be affected by propanal, which is detected at the same mass (59 amu) as acetone. GC-MS1 measured propanal concentrations, and during the whole campaign its concentration was less than 5% of the acetone concentration. Hence in this campaign, it could be assumed that PTR-MS measurements at mass 59 amu were acetone.

3.4.4 Benzene

The measured benzene concentrations of all four instruments were in good agreement, as found in previous studies by de Gouw et al. (2003b); Kaser et al. (2013) and Warnke et al. (2015) as well. The concentration ranges of PTR-MS1, GC-MS1 and GC-MS2 were similar (0.07–0.11 and 0.05–0.09 ppb respectively) while the PTR-MS2 measurements fluctuated more, between 0.07 and 0.21 ppb. The median concentrations of PTR-MS1 and GC-MS1 were the same (0.08 ppb), while the median of GC-MS2 was 0.06 ppb and the median of PTR-MS2 was 0.11 ppb.

In general, the correlations between different instrument pairs were good. The two GC-MS instruments had the highest correlation (0.92), yet the slope was not close to unity (0.77). The low slope value could be due to different sampling times of these instruments. However, as benzene does not have local sources at SMEAR II, changes in benzene concentration are slow and different sampling times should not have a great effect on the measured concentrations. PTR-MS1 correlated equally well with both PTR-MS2 and GC-MS2 (0.88 and 0.89 respectively). The slope of PTR-MS1 vs. GC-MS2 was reasonably good (0.84), while the slope between the two PTR-MS instruments was rather high (1.38). Between PTR-MS1 and GC-MS1, the correlation was 0.84 and the slope was very good (0.99). The average correlation coefficient of benzene was the same as the mean $R$ ($\overline{R}$) of acetone (0.88), and the RMS $\overline{R}$ (0.23) was lower than it was for the other compounds. As in the case of acetone, no dependency between meteorological parameters and instrument discrepancies was found.

Good correlations were expected for benzene, as the temporal and spatial changes in benzene concentration are low at the site and there are no known problems concerning benzene measurements with either GC-MS or PTR-MS. PTR-MS measurements at mass 79 amu have been reported to be only benzene, thus benzene measurements of PTR-MS are not interfered with other VOCs.

3.4.5 Toluene

Toluene was measured with all four instruments. The concentration ranges of PTR-MS1, GC-MS1 and GC-MS2 were the same from 0.02 to 0.05 ppb, with a median of 0.03 ppb. Due to high detection limits for toluene, the toluene concentrations measured with PTR-MS2 were high (0.05–0.11 ppb) and the median value (0.07 ppb) was more than twice as high as when measured by the other instruments.

Although the concentrations of the three instruments agreed well, their correlation values were only moderate. $\overline{R}$ was 0.62, while the RMS $\overline{R}$ was rather far from zero, at 0.45. The best correlation was between the two GC-MS instruments (0.77). Similarly to benzene, toluene does not have significant local sources at the site. Even though some studies (e.g. Heiden et al., 1999) have shown that under stress, many plant species may emit toluene, a recent study by Rantala et al. (2015) reported only low toluene fluxes during April and May at the site. Thus the effect of the different sampling times of the two GC-MS instruments should not be considerable. Yet, the slope of the GC-MS1 vs. GC-MS2 was far from unity (0.60). Between PTR-MS1 and GC-MS2 the slope was good (0.92), and also the correlation coefficient of 0.69 was fairly good, but the slope had a rather high confidence interval ($\pm 0.18$). The correlation between PTR-MS1 and GC-MS1 was low, 0.53 and slope far from unity (0.55). The lowest correlation was between the two PTR-MS instruments (0.50). Their slope was 1.36, with a high confidence interval of $\pm 0.52$. The toluene concentration remained below the detection limits of the PTR-MS instruments for a large amount of the time during the campaign, biasing the concentrations towards higher values. The number of data points used for the correlation analysis of toluene was less than half of the number of data points used for the other compounds.

In the study by de Gouw et al. (2003b), the correlation between PTR-MS and GC-MS was stronger ($R > 0.98$ and slope $= 1.08$) than the correlations found in this study. Additionally, the correlation coefficients between PTR-MS and PTR-Tof-MS reported by Kaser et al. (2013) and Warnke et al. (2015) were stronger than the ones measured during this campaign ($R > 0.85$). However, toluene concentrations were higher during the studies by de Gouw et al. (2003b) and Kaser et al. (2013), 0.003–1 ppb and 0.01–0.25, respectively, than the measurements presented in this study.

Although, large measurement discrepancies were found between different instruments, the explanation for this was unknown. For example, meteorological parameters had no correlation with the concentration differences of different instruments. It has been suggested that a p-cymene fragment is detected at the same mass (93 amu) as toluene with PTR-MS (Ambrose et al., 2010). Kaser et al. (2013) reported that in correcting the toluene signals for p-cymene, the linear regression between PTR-Tof-MS and another mass spectrometer improved from 0.72 to 0.98. During this campaign, p-cymene parent ion concentrations were not measured with PTR-MS instruments. Earlier measurements at SMEAR II showed that between 12 April and 15 May 2011, the toluene concentration was on average 15 times higher than the p-cymene concentration. The mean p-cymene concentration was 8 ppt, while the maximum concentration was 107 ppt (Hakola et al., 2012). Consequently, p-cymene may occasionally have an effect on the toluene concentrations measured with PTR-MS.

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High p-cymene concentrations could be expected, for example, during the monoterpene pollution episodes (Liao et al., 2011) from the nearby saw mill.

4 Conclusions

Ambient concentrations of methanol, acetaldehyde, acetone, benzene and toluene were measured using two PTR-MS instruments and two GC-MS instruments at a rural boreal forest site in the spring of 2012. Additionally, two different calibration methods, automatic and manual, were tested for the PTR-MS instruments. The calibration tests showed that both calibration methods resulted in similar sensitivities for acetaldehyde, acetone, benzene and toluene. For methanol, sensitivities obtained with the automatic method resulted in lower apparent sensitivities than the manual calibration method did. Also the sensitivity uncertainties of both PTR-MS instruments were higher for methanol than for the other compounds.

Very good correlation was found for benzene and acetone measurements between all instrument pairs. The mean correlation coefficient was 0.88 for both compounds. In the case of acetone, the RMS difference from the 1:1 line was high. However, probably due to the long sampling time of the GC-MS2, acetone broke through the adsorbent trap, resulting in measured concentrations that were too low. When the acetone data of GC-MS2 was omitted from the calculation, the RMS difference from the 1:1 line was close to zero. To measure acetone or other very volatile OVOCs using GC-MS2 it is recommended to use: a shorter sampling time, a lower flow or a stronger or a cooled adsorbent trap.

The correlation coefficients of acetaldehyde and toluene were quite far from unity, with respective averages of 0.50 and 0.62. The cause of the bad correlation in the case of acetaldehyde remains unresolved. Toluene concentrations were below the detection limits of the PTR-MS instruments for a considerable amount of the time, which biased the concentrations towards higher values and also reduced the amount of data points used for analysis.

Methanol measurements showed a robust correlation between the instruments. However, the slope values were far from unity, with an RMS difference of 0.87 from the 1:1 line. Hence, all the instruments measured the same time trends of methanol, but the quantitative concentration values must be regarded with caution. It should be noted that the uncertainty in the sensitivity of the instruments, manifesting as the deviation of the correlation slopes from unity, leads directly to similar uncertainty in the emission measurements of these compounds. This applies to, for example, eddy covariance, surface layer gradient and chamber techniques. Thus, it can be easily estimated that, for example, any emission measurement of methanol has an uncertainty of 50–100 % due to the sensitivity of the instrument used.

This study raises a few open questions yet to be answered. These include the following:

- the reasons for major biases between instruments for many compounds. This may be due to the materials used in different set-ups, but the reason remains unknown for now.
- the reason for RH dependence of the differences in acetaldehyde measurements.
- the constancy of the methanol loss correction in the GC-MS1. It is assumed to be constant, which may not be the case.

Technical recommendations arising from this experiment include the following:

- GC-MS sampling time must be adjusted to prevent breakthrough of any compound of interest. Suitable sampling time must be determined individually for each compound and the shortest time used for the measurement protocol.
- for the PTR-MS, the time used for the calibration must be long enough to reach the stable regime. The length of the calibration period must be determined individually for all compounds in the calibration gas, and the actual calibration time is determined as the longest.

The results of this study show that when doing long-term measurements of ambient air, occasional comparison measurements are needed to evaluate the measured concentrations and to quantify the uncertainties, even if the instrument is calibrated regularly.
Appendix A

Table A1. Results of the PTR-MS calibration tests. “Manual calibration method” (MCM) refers to the system in which the calibration gas standard and zero air flows are controlled manually with a pressure regulator and needle valves (see Taipale et al., 2008, for details). “Automatic method” (ACM) refers to the calibration mixing units, in which the flows are controlled automatically with mass flow controllers. All parameters of the table are presented in the [ppb·ncps$^{-1}$] unit. $\bar{S}$ and $\sigma_S$ are the mean sensitivity and the standard deviation of the sensitivity, respectively.

<table>
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<th>PTR-MS1 MCM</th>
<th>ACM</th>
<th>PTR-MS2 MCM</th>
<th>ACM</th>
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<td>$\sigma_S$</td>
<td>$\bar{S}$</td>
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</table>

$^1$ MVK: methyl vinyl ketone; $^2$ MEK: methyl ethyl ketone.
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