This is a review of the REVISED MANUSCRIPT performed by Reviewer #3. All line numbers given below are for the revised version of the manuscript.

The authors have provided additional information that answers most of the comments from the 3 reviewers. However, there are still a few points that need to be addressed before publication in ACP:

We appreciate Reviewer 3's insightful and constructive comments, which significantly improve the manuscript. Please find our response as shown below.

1/ The authors used a catalytic converter to generate zero air from ambient air. The term "VOC-free air" is more appropriate since ambient NOx are not removed. This zero air is used to perform the "C2" measurement. NOx will be present in the CRM reactor during both the "C3" measurement, when ambient air is sampled, but also during the "C2" measurement. Since OH can react with NO and NO2 during both "C3" and "C2", the CRM should be blind to the OH reactivity generated by NOx. The reported measurements of OH reactivity may therefore be biased low. Interestingly, the authors indicated L185-186 that during the SOAS campaign measurements from the UCI CRM instrument were 16% lower on average than measurements from a LIF system. Could the authors comment on this?

We have discussed the possibility and accept the limitation of this study. Now it reads:

In conclusion, it is possible that our reported OH reactivity may systematically underestimate ambient total OH reactivity as much as ambient OH reactivity coming from NO₂. (Line 206 - 208, In the track change version)

2/ The authors provided more information about operating conditions for their CRM instrument and indicated that the pyrrole-to-OH ratio was kept constant at a value of 3. This ratio can depend on the amount of OH that is produced from the photolysis of ambient water-vapor in the reactor due to the leakage of 185-nm photons. As a consequence the ratio usually changes with ambient humidity. This ratio was found to vary significantly for other CRM instruments when operated continuously during field campaigns. How did the authors manage to keep the pyrrole-to-OH constant? Was the geometry of the CRM reactor optimized to avoid the photolysis of ambient water-vapor?

We've realized that even with As we are adding 150 cc per minute of humidified N2 (likely 100 % RH) to the reactor with flow rate of 240 cc per minute, therefore, the variation of ambient humidity change would be dampened quite a bit from the mixing effect. We have included the discussion and now it reads:

Even in the field environment with various relative humidity, we have not observed noticeable changes in this ratio as we flow bulk humidified nitrogen (150 sccm) to the reactor with the total flow of 240 cc, which result in dampening the temporal ambient relative humidity variations. (Line 210 - 213, in the track change version)

Minor comments:

L192-193: "An extensive intercomparison study was conducted by Fuchs et al. (2017) with various OH reactivity measurement techniques that highlighted potential analytical artifacts in the CRM technique. These artifacts have all been examined and preventive measures have been implemented in the UCI CIMS-CRM system deployed at TRF." - I would add some caveat here since these artefacts have not been fully investigated for this instrument. While some testing has been performed to check whether ambient NO and O3 could lead to measurement artifacts, the authors acknowledged in their responses to the first review that additional tests are needed to well characterize these artefacts.

We have clarified the limitation in the revised manuscript. Now, it reads:

Again, as the CRM method is relatively new technique, one should keep in mind that the future studies may find potential artifacts that we do not report in this study. (Line 223 - 225, in the track change version)

L200-201: "Our approach to this type of interference has been to determine the maximum NO level, noticeably interfering with the calibration regression line shown in Sanchez et al. (2018). Laboratory tests indicate that the statistical agreement started to veer off when the NO level is 5 ppb in 1 σ of the linear regression" – These tests are of interest for the CRM community and the reviewer recommends to show them in the supplementary material. This will also provide additional confidence in the dataset.

We have added the quantitative information. Now it reads.

Laboratory tests indicate that the statistical agreement started to veer off when the NO level is 5 ppb in 1 of the linear regression as the slope for the calibration curve has changed from 0.238 to 0.246. (Line 201 - 203, in the track change version)

L214-216: "In the 2015 field campaigns conducted in Seoul South Korea (Kim et al., 2016), we conducted a standard addition experiment for the propene standard for additional ~ 30 s-1 in two different ozone environment 65 ppb and 123 ppb. The outcome illustrates an agreement between two additions within the analytical uncertainty." – While a standard addition test could highlight an artefact impacting the linear response of CRM to OH reactivity, it cannot rule out an artefact leading to a positive or negative offset that would only depends on O3. For the later, a standard addition of 30s-1 of OH reactivity would always lead to the right change in the measured total OH reactivity. However, the total OH reactivity with (ambient reactivity + standard addition reactivity) or without (ambient reactivity) standard addition would be biased low by the same amount. This should be acknowledge in the manuscript.

We have clarified the limitation in the revised manuscript. Now, it reads:

The outcome illustrates an agreement between two additions within the analytical uncertainty although a systematic laboratory study will warrant an accurate uncertainty assessment from ozone. (Line 222 - 224, in the track change version)

L206-211: "Therefore, we performed multi-point calibrations with a propene mixture using a NIST traceable gas standard (AirLiquide LLC, 0.847 ppm) during the field campaign to avoid any circumstances where the pseudo first-order reaction regime is not established." – Please indicate the range of OH reactivity generated during the multipoint calibration.

We have included the information. Now, it reads:

we performed multi-point calibrations (5 s⁻¹ to 30 s⁻¹) with a propene mixture using a NIST traceable gas standard (AirLiquide LLC, 0.847 ppm) during the field campaign to avoid any circumstances where the pseudo first-order reaction regime is not established. (Line 214 - 217, in the track change version)

L244-245: The writing of Eq. 1 is still confusing (mixing of parameters and units). Please only use parameters that are defined in the main text. For instance:

MRvoc=Svoc*kbenzene/kvoc*1/Rbenzene.

MR:Mixing Ratio, Svoc: normalized voc signal, kbenzene and kvoc: proton transfer rate constants, Rbenzene: benzene sensitivity

We have revised the equation as suggested. Now it reads.

ppb_{VOC} is the mixing ratio of an analyte.

ncps_{VOC} is the mass discrimination corrected normalized count for an analyte.

k_{benzene} is the proton transfer reaction rate constant for benzene.

k_{VOC} is the proton transfer reaction rate constant for an analyte.

(Line 257 - 260, in the track change version)

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2	Contributions to OH reactivity from unexplored volatile organic compounds measured by
3	PTR-ToF-MS- A case study in a suburban forest of the Seoul Metropolitan Area during
4	KORUS-AQ 2016
5	
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7	Kim, ³ Joonyoung Ahn, ³ Don Blake, ⁴ Scott Herndon, ⁵ Daun Jeong, ¹ John T. Sullivan, ⁶ Thomas
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24 25	To be submitted to Atmospheric Chemistry and Physics

Abstract

comparative reactivity method (CIMS-CRM) instrument in a suburban forest of the Seoul Metropolitan Area (SMA) during Korea US Air Quality Study (KORUS-AQ 2016) from mid-May to mid-June of 2016. A comprehensive observational suite was deployed to quantify reactive trace gases inside of the forest canopy including a high-resolution proton transfer reaction time of flight mass spectrometer (PTR-ToF-MS). An average OH reactivity of $30.7\pm5.1~\text{s}^{-1}$ was observed, while the OH reactivity calculated from CO, NO + NO₂ (NO₃), ozone (O₃), sulfur dioxide (SO₂), and 14 volatile organic compounds (VOCs) was $11.8\pm1.0~\text{s}^{-1}$. An analysis of 346 peaks from the PTR-ToF-MS accounted for an additional $6.0\pm2.2~\text{s}^{-1}$ of the total measured OH reactivity, leaving 42.0 % missing OH reactivity. A series of analyses indicates that the missing OH reactivity most likely comes from VOC oxidation products of both biogenic and anthropogenic origin.

We report OH reactivity observations by a chemical ionization mass spectrometer –

1. Introduction

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Total OH reactivity (s⁻¹), the inverse of OH lifetime, is a measure of the total amount of reactive trace gases in the atmosphere in the scale of reactivity, which allow us to quantitatively evaluate our ability to constrain trace gases by comparing measurements of total OH reactivity with the OH reactivity calculated from a speciated reactive gas measurement dataset. The fraction of observed OH reactivity that cannot be reconciled by calculated OH reactivity is known as "missing OH reactivity" (Di Carlo et al., 2004; Goldstein and Galbally, 2007; Yang et al., 2016). A substantial amount of missing OH reactivity has consistently been reported in forest environments (30 - 80%). Di Carlo et al. (2004) conducted a study in a mixed forest near Pellston, Michigan where they reported missing OH reactivity (~30 %) larger than observational uncertainty. The authors concluded that the missing sources of reactivity were primary biogenic volatile organic compound (biogenic VOC, BVOC) emissions, as the degree of missing OH reactivity followed the temperature dependence of terpenoid emissions. In a boreal forest in Hyytiälä, Finland, Sinha et al. (2010) report a similar result with observed trace gases that account for only 50% of the measured OH reactivity. They argued that oxidation products of BVOCs alone could not account for the missing OH reactivity. Thus, they also concluded that primary emissions were more likely to be the source of missing OH reactivity and they further suggest that this could be the result of the contribution of small amounts of many reactive gases. Follow up studies (Nolscher et al., 2012; Praplan et al., 2019) at the same site have presented a consistent conclusion. Nolscher et al. (2012) observed the highest level of missing OH reactivity during a heat wave episode, possibly inducing a stress emission response from the local forest. A comprehensive analysis by Praplan et al. (2019) using a long-term observation dataset and a

photochemical model framework with the Master Chemical Mechanism illustrates that the model simulated oxidation compound contribution can only contribute 7 % of missing OH reactivity.

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On the other hand, some studies have attributed the sources of the missing OH reactivity to unmeasured oxidation products of well-characterized BVOCs. Edwards et al. (2013) measured OH reactivity in a pristine tropical forest in the Sabah region of Borneo during the Oxidant and Particle Photochemical Processes (OP3) field campaign (Hewitt et al., 2010). This study implemented the Master Chemical Mechanism (MCMv3.2) (Saunders et al., 2003; Jenkin et al., 1997) into a box model framework to quantify potential contributions from unmeasured oxidation products. The model was constrained with VOCs such as isoprene, monoterpenes, and alkanes and alkenes and other observed trace gases such as NO + NO₂ (NO_x) and ozone (O₃). The authors reported that the model simulated oxygenated VOCs (OVOCs) could contribute 47.1% of the calculated OH reactivity – surpassing the contribution from isoprene, the primary emission of this ecosystem. It is notable that 30% of observed OH reactivity could not be accounted for by the box model simulations. After examining the comprehensive observational suite of VOCs, the authors determined that the most significant missing sources of OH reactivity were likely secondary multifunctional carbon compounds rather than primary BVOC emissions. Hansen et al. (2014) suggested that their observed missing OH reactivity were likely from unmeasured oxidation products during the Community Atmosphere-Biosphere INteraction EXperiment (CABINEX 2009) in Michigan. This notion was also consistent with findings reported by Kim et al. (2011) who measured OH reactivity of branch enclosures from four representative tree species in the forest canopy during the CABINEX study. They reconciled most of the measured OH reactivity of four representative tree species with well-known BVOCs, such as isoprene and monoterpenes. Finally, Nakashima et al. (2014) reported that 29.5% OH

reactivity could not be reconciled by the speciated trace gas dataset during the Bio-hydro-atmosphere interactions of Energy, Aerosols, Carbon, H₂O, Organics and Nitrogen-Southern Rocky Mountain 2008 (BEACHON-SMR08) field campaign (Ortega et al., 2014). The campaign took place at the Manitou Experimental Forest (MEF) in Colorado, a ponderosa pine plantation dominated by primary BVOC emissions of 2-methyl-3-butene-2-ol (232-MBO) and monoterpenes (Ortega et al., 2014). The authors also reported that the missing OH reactivity was likely from BVOC oxidation products. In the same context, Kim et al. (2010) conducted PTR-MS mass spectrum analysis for both ambient air and branch enclosures at the MEF site. They reported more conspicuous unidentified signals on PTR-MS mass spectra in the ambient samples than those from branch enclosure samples at this site.

During the Southern Oxidant and Aerosol Study (SOAS) in 2013, Kaiser et al. (2016) used a comprehensive suite of VOC measurements at an isoprene dominant forest site in the southeastern US to examine the role of the OVOC species in missing reactive carbon. The authors used MCMv3.2 embedded in the University of Washington Chemical Box Model (UWCM) to compare OH reactivity from model-generated OVOCs to OH reactivity from measurements of OVOCs. There was no significant discrepancy between the average measured and calculated OH reactivity including observed trace gases and model calculated oxidation products of VOCs. However, it was noted that a small portion (1 s⁻¹) of observed OH reactivity could not be reconciled by the model calculation. As this fraction was not correlated to isoprene oxidation products, it was suggested that the missing OH reactivity may be due to unmeasured primary emissions. One caveat of this analysis pointed out by the authors was that the concentrations of the modeled first-generation isoprene oxidation products (e.g. MVK, MACR, isoprene hydroxy hydroperoxides (ISOPOOH), isoprene nitrates (ISOPN), and hydroperoxy

aldehydes (HPALD)) were significantly overpredicted in the afternoon. Consequently, the uncertainty of the model calculation is likely to be much higher for the multi-generation oxidation products and their contributions to the OH reactivity contributions. This result highlights the uncertainty in relying solely on box-model results to assess OH reactivity. This status quo urges us to take a convergent approach by effectively integrating observational results from novel instrumentation and model outcomes.

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This study examines the OH reactivity observations at Taehwa Research Forest (TRF) supersite from 15 May 2016 to 7 June 2016 during the Korea United States Air Quality Study 2016 (KORUS-AQ 2016) campaign. TRF (37 18' 19.08" N 127 19' 7.12" E, 162 m altitude) is operated by Seoul National University and located in Gwangju in the Gyunggi Province in South Korea (Kim et al., 2013b). The site is about 35 km southeast from the center of Seoul and borders the greater Seoul Metropolitan Area (SMA) with its population of 25.6 million. This geographical proximity to SMA results in a significant level of anthropogenic influence, particularly in elevated NO_x (Kim et al., 2016). Additionally, occasional pollution transport events occur at regional scales. Previous studies at the site have consistently highlighted the importance of BVOC photochemistry at TRF (Kim et al., 2016;Kim et al., 2013a;Kim et al., 2015). Isoprene and monoterpenes are the dominant OH sinks at the site among observed VOCs. The elevated NO_X accelerates the photochemical processing of VOCs (Kim et al., 2015). Thus, this site is an ideal natural laboratory to study contributions towards total OH reactivity from primary trace gas emissions from both natural and anthropogenic processes and their oxidation products. This motivated us to deploy a high-resolution proton transfer reaction time-of-flight mass spectrometer (PTR-ToF-MS) to quantify trace amounts of VOCs with unknown molecular structures by taking advantage of the universal sensitivity of hydronium ion chemistry towards

reactive VOCs (Graus et al., 2010; Jordan et al., 2009a). Therefore, we intend to observationally constrain the contributions of conventionally unidentified or unmeasured VOCs towards OH reactivity.

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2. Methods

2.1.Field Site

The Taehwa Research Forest is a Korean pine (Pinus koraiensis) plantation (300 m × 300 m) surrounded by a deciduous forest dominated by oak trees (Kim et al., 2013b). A flux tower (40 m height) at the center of TRF has air-sampling inlets at multiple heights (4 m, 8 m 12 m, and 16 m) below the canopy top (20 m). Each inlet consists of Teflon tubing (3/8" OD) with ~ 1 second of residence time. The trace gas dataset including VOCs presented is the average of concentrations measured at the inlets inside of the canopy as previous studies illustrate that there is no substantial vertical VOC gradients inside of the canopy (within 3 %, Kim et al. (2013b)). An air-conditioned instrument shack located at the base of the flux tower housed the PTR-ToF-MS for VOC measurements, a mini tunable infrared laser direct absorption spectroscopy (mini-TILDAS) instrument for HCHO, methane, and methanol measurements, and analyzers for carbon monoxide (CO), sulfur dioxide (SO₂), ozone (O₃), and meteorological measurements. The OH reactivity and NO_x analyzers were located in another nearby air-conditioned shack (3 m apart) and sampled air through an extended Teflon inlet line of 4 m (1/4" OD) from the ground with a flow rate of 4 sLpm resulting in a 0.5 second residence time. The height of the ambient air intake was 3.5 m. The analytical characteristics of the instrumentation suite are summarized in Table 1. A ceilometer backscattering characterized boundary layer vertical structure at the site. The ceilometer analysis described by Sullivan et al. (2019) reveals the diurnal boundary layer

height evolution, indicating a maximum in the afternoon around 1-3 km and a minimum in the early morning below 500 m.

2.2.OH Reactivity Measurements

A chemical ionization mass spectrometer – comparative reactivity method (CIMS-CRM) instrument was used to measure OH reactivity. The UCI CIMS-CRM system includes a chemical ionization mass spectrometer with a hydronium reagent ion. The CRM method measures total OH reactivity by quantifying the relative loss of pyrrole, a highly reactive gas ($k_{\rm OH^+}$ pyrrole = 1.07 \times 10⁻¹⁰ cm³ molecule⁻¹ s⁻¹ at 298 K (Dillon et al., 2012)) that is rarely found in the atmosphere (Sinha et al., 2008b). Nitrogen gas flows through a bubbler full of ultrapure liquid chromatography mass spectrometer (LC-MS) grade water to produce water vapor. The water vapor then flows into a glass reactor where it is photolyzed into OH radicals by a mercury lamp (Pen-Ray® Light Source P/N 90-0012-01). The measurement uncertainty is 16.7% (1 σ) with a limit of detection of 4.5 s⁻¹ over 2 minutes (3 σ).

The UCI CIMS-CRM instrument has been deployed on multiple occasions, including the Megacity Air Pollution Study (MAPS)-Seoul 2015 campaign that incorporated previous measurements at the TRF ground site during September 2015 (Sanchez et al., 2018;Kim et al., 2016). During the SOAS 2013 campaign, an ambient OH reactivity intercomparison study was conducted with laser induced fluorescence (LIF) system (Sanchez et al., 2018). The instrument intercomparison showed that the OH reactivity measurements from the CRM and LIF instruments generally agreed within the analytical uncertainty. An average of 16% difference between the techniques was noted in the late afternoons where the CRM measurements were lower than those from LIF. As discussed in Sanchez et al. (2018), this is likely caused by the

difference in sampling strategies, as the CRM measurements relied on a lengthy Teflon inlet (15 m) while the LIF directly sampled air at the top of a walk up tower. As mentioned above, at TRF we used a shorter inlet line to minimize residence time and avoid inlet line loss.

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coming from NO₂.

An extensive intercomparison study was conducted by Fuchs et al. (2017) with various OH reactivity measurement techniques that highlighted potential analytical artifacts in the CRM technique. These artifacts have all been examined and preventive measures have been implemented in the UCI CIMS-CRM system deployed at TRF. This included a laboratory-built catalytic converter (Pt-wool at 350 °C) that minimized the interferences due to changes in air to prevent the interference from the difference in humidity for the zero air characterizations. Hansen et al. (2015) illustrated that NO_X may be generated from the catalytic converter. To prevent potential NO_X interferences, they used a scrubber with Purafil and activated charcoal, which will modulate the humidity in the sample. Our approach to this type of interference has been to determine the maximum NO level, noticeably interfering with the calibration regression line shown in Sanchez et al. (2018). Laboratory tests indicate that the statistical agreement started to veer off when the NO level is 5 ppb in 1 σ of the linear regression between instrument response (unitless) and OH reactivity (s_1^{-1}) as the slope for the calibration curve has changed from 0.238 to 0.246. In addition, Kim et al. (2016) achieved an OH reactivity budget closure in high NO2 condition, which implies no significant interferences from NO2. However, in response to the Fuchs et al. (2017) observation that various CRM configurations suffer from different levels of NO_X interferences, we plan to conduct more systematic NO_X interference tests to more accurately characterize this system. In conclusion, it is possible that our reported OH reactivity may systematically underestimate ambient total OH reactivity as much as ambient OH reactivity

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We consistently kept the pyrrole to OH ratio at 3:1 and so did not achieve a pseudo first order relationship. Even in the field environment with various relative humidity, we have not observed noticeable changes in this ratio as we flow bulk humidified nitrogen (150 standard cc per minute) to the reactor with the total flow of 240 cc, which result in dampening the temporal ambient relative humidity variations. Therefore, we performed multi-point calibrations (5 s⁻¹ to 30 s⁻¹) with a propene mixture using a NIST traceable gas standard (AirLiquide LLC, 0.847 ppm) during the field campaign to avoid any circumstances where the pseudo first-order reaction regime is not established. Detailed calibration procedures for the OH reactivity system including

laboratory multi-component calibration results can be found in Sanchez et al. (2018).

In addition, Fuchs et al. (2017) also described a potential interference from ambient O₃ in some CRM systems. In the 2015 field campaigns conducted in Seoul South Korea (Kim et al., 2016), we conducted a standard addition experiment for the propene standard for additional ~ 30 s⁻¹ in two different ozone environment 65 ppb and 123 ppb. The outcome illustrates an agreement between two additions within the analytical uncertainty although a systematic laboratory study will warrant an accurate uncertainty assessment from ozone. Again, as the CRM method is a relatively new technique, one should keep in mind that the future studies may find potential artifacts that we do not report in this study.

2.3.PTR-ToF-MS Measurements

A high-resolution PTR-TOF-MS (Ionicon Analytik GmbH) (de Gouw and Warneke, 2007b) (Jordan et al., 2009b) was deployed at the TRF site. The instrument was operated with a drift tube temperature of 60 °C, 560 V drift voltage, and 2.27 mbar drift tube to maintain E/N of 126 Td. Background checks were manually conducted about three times a day for a 10-minute

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duration by scrubbing the ambient air through a catalytic convertor (Pt-wool maintained at 350° C). The detectable peaks from the ambient spectra were assessed by subtracting the background spectrum. The instrument was calibrated with a gas mixture manufactured by Apel-Riemer Environmental Inc. The mixture contains ~ 1 ppmv of acetaldehyde, acetone, isoprene, methyl vinyl ketone, methacrolein, benzene, methyl ethyl ketone, toluene, o-xylene, and α -pinene. This standard mixture was only used for the PTR-ToF-MS calibration and not the CRM-CIMS calibration. The concentration of the compounds were assessed in the Blake Lab at University of California, Irvine, who also conducted the airborne VOC analysis using whole air samples during the KORUS-AQ campaign on the NASA DC-8 (Colman et al., 2001).

A mass range of m/z 40 to m/z 267 was analyzed from the recorded PTR-ToF-MS mass spectra. An automatic mass scale calibration was conducted every 5 minutes on the data averaged over 30 seconds. The raw PTR-ToF-MS data were processed using the PTRwid software described by Holzinger (2015). We normalized the mass peaks by 10⁶ reagent ion counts (H₃O⁺). As the majority of the VOC mass peaks could not be directly calibrated, we determined the VOC sensitivities using equation 1 (Eq 1). This method has been employed by a number of previous studies such as Cappellin et al. (2012). The benzene calibration factor was used to calculate mixing ratios by applying its proton transfer reaction rate coefficient (*k*_{benzene}) and sensitivity (ncps ppb⁻¹) for the available compounds. The application of this equation can be justified since PTRwid provides the mass discrimination corrected counts as a final computational product.

$$ppb_{VOC} = ncps_{VOC} \times \frac{k_{benzene}}{k_{VOC}} \times \frac{1}{11.94 \, ncps \, ppb^{-1}} \qquad \text{Eq. 1}$$

where, 11.94 ncps ppb-1 is the assessed sensitivity of benzene.

ppbyoc is the mixing ratio of an analyte.

ncpsyoc is the mass discrimination corrected normalized count for an analyte.

kbenzene is the proton transfer reaction rate constant for benzene.

kyoc is the proton transfer reaction rate constant for an analyte.

For the mass peaks where specific proton transfer reaction rates were unavailable, we estimated the mixing ratios by applying a proton transfer reaction rate coefficient ($k_{\rm H3O^+}$) of 3.00 \times 10⁻⁹ cm³ s⁻¹, the default value for PTRwid calculations. The spectra had a limit of detection of tens of ppt for a 30 second average. The calibrated compounds had a range of detection limits as low as 3.7 ppt for α -pinene and as high as 48 ppt for toluene.

2.4. OH Reactivity Calculation

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OH reactivity was calculated from the concentrations of all the compounds observed by the instrumental suite described in Table 1. The original data can be found in the KORUS-AQ 2016 data archive at https://korus-aq.larc.nasa.gov/. A total of 360 mass peaks measured by the PTR-ToF-MS were analyzed above the background (3 σ or above) to assess their contribution to the calculated OH reactivity. Fourteen of the mass peaks were identified as VOCs commonly reported for PTR-MS measurements (Table 1), leaving 346 unidentified peaks. These remaining mass peaks were grouped into three categories in order to estimate their possible OH reactivity contribution.

Category I (81 peaks) included mass peaks for which the PTRwid software calculated a molecular formula. OH reaction rate coefficients for the individual peaks were obtained from the National Institute for Standards and Technology (NIST) Webbook library. As the only

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information we have is the molecular composition, we identified multiple isomers with different functional groups and thus different reactivity. We have extensively reviewed previous publications (Williams et al., 2001; De Gouw et al., 2003; de Gouw and Warneke, 2007a; Jordan et al., 2009a; Ruuskanen et al., 2011; Muller et al., 2012; Koss et al., 2017a) identifying ambient VOCs using PTR-MS with both quadrupole and time-of-flight systems to identify possible compounds. For example, for the m/z of 75.043, there are four possible compounds including hydroxy acetone, propionic acid, methyl acetate, and ethyl formate. We used the median reaction constant for the set of possible compounds. The detailed description of the OH reaction constant determination process for the Category I peaks is described in Sanchez (2019). If the information was unavailable from the NIST Webbook database, a structure-reactivity relationship described by Kwok and Atkinson (1995) was applied to obtain reaction rate coefficients. This is an empirical calculation system to estimate k_{OH} based upon the number of carbons and the functional groups of given VOCs. The framework is able to calculate k_{OH} within a factor of two according to a thorough assessments presented in Kwok and Atkinson (1995). However, the authors discourage the application of the framework to compounds that were not examined in the study such as halogenated compounds. Although halogenated compounds are not included in this study, one should be aware of a potentially significant uncertainty.

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Category II (28 peaks) included mass peaks for which the PTRwid software could not assess an exact molecular composition due to uncertainty in the data processing system. Nonetheless, this group of compounds illustrated a positive correlation ($R^2 = 0.19$ to 0.88) with either anthropogenic (benzene, toluene) or biogenic (MVK+MACR and monoterpenes) VOCs. Category II compounds are further grouped into subcategories corresponding to these two main VOC sources. OH reaction rate constants ($k_{\rm OH}$) were estimated with equations based on the

relationship between the m/z and the $k_{\rm OH}$ of compounds in Table 1 (Figure S1). More specifically, we assume that $k_{\rm OH}$ is linearly correlated with m/z. To apply this linear relationship, the compounds with known $k_{\rm OH}$ were grouped into $5 \ m/z$ bins and the average $k_{\rm OH}$ of each bin was calculated. The green triangles represent $5 \ m/z$ binned averages from these compounds plotted with their respective average $k_{\rm OH}$. This approach can be justified by the fact that the reaction constants of VOCs towards OH tend to increase as a function of molecular mass within functional groups (Kwok and Atkinson, 1995;Atkinson, 1987). The y-intercepts of the linear regressions were assessed using the $k_{\rm OH}$ values of the biogenic or anthropogenic compounds and their masses.

Category III (237 peaks) included mass peaks with very low mixing ratios (average = 4.8 ppt ± 19.5 ppt) that were above the limit of detection. We applied a k_{OH} corresponding to the dark green best-fit line in Figure S1 to these peaks. The y-intercept of the dark green line was based on that of acetaldehyde, as it was the lowest mass compound used for the OH reactivity calculations in this study.

There are two components that need to be considered for the assessment of uncertainty associated with calculated OH reactivity: the concentration and the reaction constants with OH. The uncertainty of the observed trace gases is in the range of 5 % to 20 % as shown in Table 1 and is associated with the rate constants from laboratory experiments (Atkinson et al., 2006). Combining 15 % uncertainty from reaction constants and 13.5 % from trace gas observations results in 20 % of uncertainty in calculated OH reactivity. This should be considered as a conservative estimate as most VOC concentrations and associated rate constants are empirically estimated.

3. Results and Discussion

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An average OH reactivity of $30.7 \pm 5.1 \text{ s}^{-1}$ was observed from 15 May - 7 June 2016 (Figure 1). This was within the range of OH reactivity observed in urban regions (10 - 33 s⁻¹). (Kovacs et al., 2003;Ren et al., 2003;Sinha et al., 2008a;Dolgorouky et al., 2012;Whalley et al., 2016; Kim et al., 2016; Yang et al., 2017) and in the range of previously reported observations and model calculations at the TRF site (~15 - 35 s⁻¹) (Kim et al., 2016;Kim et al., 2015). The total calculated OH reactivity of $11.8 \pm 1.0 \text{ s}^{-1}$ from the measured compounds in Table 1 resulted in 63.3% missing OH reactivity. However, an additional OH reactivity of $6.0 \pm 2.2 \text{ s}^{-1}$ was further calculated from the reactivity of the VOCs in Categories I – III. The contribution lowered the missing OH reactivity level to 42% of the measured OH reactivity. Kim et al. (2016) had previously measured an average OH reactivity of 16.5 s⁻¹ at TRF during the MAPS-Seoul campaign from 1 September – 15 September 2015, a substantially lower level then what we report during this springtime study. Although small alkanes and alkenes such as ethane, ethene, propane and propene were not observed on the site, we utilized the dataset from the NASA DC-8 that flew at 700 m above the site, which indicates that their contribution was consistently small ($\sim 0.7 \text{ s}^{-1}$ in average). In this suburban forest, we do not think there is any substantial emission sources of these relatively long-lived VOCs. The difference can be attributed to the notably higher reactive trace gas loadings during KORUS-AQ compared to the TRF measurements during MAPS-Seoul. The NO_x, benzene, and toluene concentrations were 3 times higher during KORUS-AQ and CO was 1.4 times higher (Figure S2). Although the average isoprene concentrations were similar between the two campaigns, MVK and MACR concentrations during KORUS-AQ were ~3 times higher, illustrating a higher oxidative environment. There was a persistently high MVK+MACR to

isoprene ratio of 1.8 during the KORUS-AQ campaign at TRF. This ratio was similar to the value reported during the summer in a moderately polluted forest in the Pearl River Delta that was attributed to a strong atmospheric oxidation capacity (Gong et al., 2018). The missing OH reactivity during KORUS-AQ was generally much higher than levels reported during urban observations (up to 50% missing OH reactivity) (Kovacs et al., 2003;Ren et al., 2003;Sinha et al., 2008a; Dolgorouky et al., 2012; Whalley et al., 2016; Kim et al., 2016; Yang et al., 2017) and within the range of previously reported values in forest regions where as much as 80% missing OH reactivity has been reported (Kim et al., 2016;Di Carlo et al., 2004;Nolscher et al., 2012; Edwards et al., 2013; Nolscher et al., 2016; Ramasamy et al., 2018; Nakashima et al., 2014). Figure 2 shows the diurnal average of measured, calculated, and missing OH reactivity from 15 May - 7 June 2016. Isoprene was the largest contributor to VOC OH reactivity in the afternoon and the early evening (36% of the calculated OH reactivity in the evening), consistent with the previous studies conducted in this site (Kim et al., 2016;Kim et al., 2013b;Kim et al., 2015). Among all the trace gases, the largest average contributor to the calculated OH reactivity was NO_x, which contributed 18.2% (5.6 s⁻¹) to the measured OH reactivity. The NO_x contribution to OH reactivity is higher during the morning and evening rush hours and at a minimum in the afternoon, which has been reported consistently in previous reports conducted near megacities (Kovacs et al., 2003; Mao et al., 2010; Dolgorouky et al., 2012; Ren et al., 2003; Shirley et al., 2006). Enhanced OH reactivity during the morning or night and minimum OH reactivity during the afternoon have been reported in urban areas (Kovacs et al., 2003; Ren et al., 2006; Shirley et al., 2006; Dolgorouky et al., 2012; Mao et al., 2010; Whalley et al., 2016). On the other hand, strong light-sensitive biogenic emissions (e.g. isoprene) result in a maximum observed OH reactivity in the afternoon in forested regions (Ren et al., 2006; Sinha et al.,

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2012; Edwards et al., 2013; Hansen et al., 2014; Zannoni et al., 2017; Nolscher et al., 2016). One exception is an OH reactivity observation conducted in Hyytiälä, a forested site that has low isoprene levels, by Sinha et al. (2010). They attributed a flat diurnal OH reactivity variation to the interplay between high daytime emissions and low nighttime boundary layer height. In urban environments, it is mostly anthropogenic trace gases such as aromatics and OVOCs that contribute to OH reactivity. These compounds have a longer lifetime compared to the diurnal boundary layer evolution. This leads to the accumulation of such compounds in the shallow boundary layer during the night. On the other hand, strong emissions of reactive BVOCs in deciduous forest regions enhance OH reactivity during the daytime but then quickly react away. Very subtle diurnal differences observed in this study (Figure 2), therefore, can be understood as the competitive influences of both anthropogenic and biogenic compounds to the OH reactivity. As described in detail in Sullivan et al. (2019) and Jeong et al. (2019), a strong regional stagnation episode occurred during the KORUS-AQ campaign between May 17 – 23. Later, the Korean Peninsula was affected by a period of continental pollution outflow between May 28 and June 1. The diurnal averages of the two periods and their calculated OH reactivity are presented in Figure 3. It is notable that there is very little difference in the observed OH reactivity between the two distinct periods in terms of the amount of OH reactivity and its diurnal pattern (Figure 4). Furthermore, no significant variance of the different classes of reactive gases such as criteria air pollutants (CO, NO_x, O₃, and SO₂), mostly contributed by NO_x, OVOCs (acetone, acetaldehyde, formaldehyde, methylglyoxal, methanol, methyl ethyl ketone), aromatics (benzene, toluene, xylenes, styrene, benzaldehyde, trimethylbenzenes), and BVOCs (isoprene, monoterpenes, sesquiterpenes, MVK+MACR) was observed during the different periods (Figure 5). These different classes of reactive gases generally differed by less than 10% during the two periods

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from the overall campaign. This observation shows that the presence of reactive gases is mostly controlled by relatively short-lived compounds determined by local emissions and their oxidation products.

The diurnal variation behavior of each chemical class reflects the chemical lifetime of the compounds (e.g. aromatics vs BVOCs). The calculated OH reactivity from OVOCs does not show a strong diurnal variation. This reflects the fact that OVOCs are mostly generated or emitted during the daytime and their lifetime is generally longer than their precursors, which allows nocturnal accumulation due to the absence of OH. The differences in the diurnal variation of different classes of reactive gases can also be used to interpret the origin of the compounds in Categories I-III as presented in Figure 6. The diurnal variations of Category I resemble those of relatively long-lived chemical species with a distinct nocturnal accumulation pattern. This diurnal pattern has been previously reported for both anthropogenic VOCs such as toluene and benzene and temperature dependent monoterpenes such as α -pinene. It is notable that the diurnal pattern is enhanced during the stagnation period during early morning hours. This enhancement is also seen in the aromatic trace gases particularly during the stagnation period (Figure 5b).

Indeed, there are both biogenic and anthropogenic contributions towards the Category I compounds, which contribute an average of 3.8 s^{-1} to the OH reactivity assessment, the largest amount among the three categories (Figure 6a). The largest contributors to Category I, which appear to be from a mixture of biogenic and anthropogenic sources, include m/z 89.060, 101.06, and 101.096, and they contributed 0.3 s^{-1} , 0.2 s^{-1} , and 0.2 s^{-1} , respectively. The m/z 89.060 had a molecular formula of $C_4H_8O_2H^+$ and was correlated to the anthropogenic compounds such as benzene and toluene. The m/z 101.06 peak had the molecular formula of $C_5H_8O_2H^+$ and had a diurnal variation similar to that of MVK + MACR. This mass peak has been previously

identified in laboratory (Zhao et al., 2004) and field (Williams et al., 2001) studies as the C₅ hydroxy carbonyl, an isoprene oxidation product. Results from an indoor chamber photooxidation experiment conducted by Lee et al. (2006) showed that m/z 101 is a common fragment of unidentified oxidation products of monoterpenes, sesquiterpenes, and isoprene. Lee et al. (2006) also reported that this mass peak also composed over 5% of the fragments of unidentified α- humulene and linalool oxidation products. The molecular formula of this peak is C₆H₁₂OH⁺, and it has been identified in previous studies as C₆ carbonyls (Koss et al., 2017b) or hexanal (Brilli et al., 2014; Rinne et al., 2005). Furthermore, m/z 99.044 and 113.023 were also among the highest contributors to Category I and were correlated with MVK and MACR. The m/z 99 was previously reported to be a fragment ion of unidentified terpene oxidation products in a chamber experiment (Lee et al., 2006). The m/z 113 was observed by a PTR-MS in a Ponderosa pine forest in central California by Holzinger et al. (2005). In this case, it was formed within the canopy from the rapid oxidation of terpinolene, myrcene, and α -terpinene. Furthermore, m/z 113 was observed to come from the photooxidation and ozonolyis of multiple terpenes in two indoor chamber studies by Lee et al. (2006). The m/z 113 composed over 5% of the oxidation product fragments of myrcene and verbenone. Finally, m/z 83.085 had the molecular formula of C₆H₁₁⁺ and was correlated to benzene. Multiple studies have identified this peak as cyclohexane, methyl-cyclopentane, or methylcyclohexane, typically found in areas rich in oil and gas (Koss et al., 2017b; Gueneron et al., 2015; Yuan et al., 2014). In summary, both the gross diurnal pattern and the individual peak analyses consistently illustrates that both anthropogenic and biogenic compounds comprise Category I, the largest contributor to the previously unexplored compounds in the PTR-ToF-MS spectrum at this research site.

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Category II contributed an average of 0.3 s⁻¹ to the calculated OH reactivity, the lowest amount for the three Categories (Figure 6b). The compounds in category II appear to correlate to either BVOCs or acetone, depending on the time period. In Figure 6b, the maximum during the transport period is enhanced to about 0.2 s⁻¹ higher than the overall campaign and shifted about 3 hours later to ~4:00 PM. The OH reactivity calculated from Category II is strongly correlated to MVK + MACR ($r^2 = 0.82$) during this period as well. On the other hand, during the stagnation period the average OH reactivity from Category II correlates more strongly with acetone (r^2 = 0.62) than with MVK +MACR ($r^2 = 0.28$). In fact, six of the highest contributors to Category II (Figure 6b) are more strongly correlated to acetone ($r^2 > 0.40$) during the stagnation period compared to the transport period. The sources of acetone can be either biogenic or anthropogenic. Biogenic sources include direct emissions from plants or their oxidation products and plant decay (Jacob et al., 2002; Seco et al., 2007). Anthropogenic sources of acetone include vehicular emissions, solvent use, and the oxidation of other anthropogenic VOCs (Jacob et al., 2002). Therefore, this illustrates that the compounds in Category II also have a complex source profile of both biogenic and anthropogenic origin. Category III contributed 1.9 s⁻¹ to the calculated OH reactivity (Figure 6c). The six highest contributors out of 236 mass peaks contributed a total of 0.43 s⁻¹ of the calculated OH reactivity. Overall, Category III compounds had no strong correlations to isoprene, MVK+MACR, benzene, or toluene during either the stagnation or transport periods. However, Category III compounds were highly correlated to methylglyoxal ($r^2 = 0.85, 0.82, \text{ and } 0.78 \text{ for}$ the stagnation, transport, and overall period, respectively), one of the measured OVOCs. A global modeling study illustrated that methylglyoxal is mainly produced from isoprene oxidation processes and the second most important source is acetone oxidation (Fu et al., 2008). In

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addition, aromatics and alkenes are also known to produce methylglyoxal through atmospheric oxidation processes (Henry et al., 2012). As TRF is a high aromatics and high isoprene environment, the source profile of methyl glyoxal in the region is likely complex, which can be applied to interpret the source of the Category III compounds.

Overall, the OH reactivity estimates from Categories I – III contributed an average of 6.0 \pm 2.2 s⁻¹ to the calculated OH reactivity. In summary, there is consistency that both anthropogenic and the biogenic contributions need to be further studied in the PTR-ToF-MS spectrum. Furthermore, by adding this additional signal from Category I, II, and III, VOC contribution to calculated OH reactivity (11.0 s⁻¹) becomes larger than that (6.8 s⁻¹) from criteria air pollutants (CO, NO_x, SO₂ and O₃). This should be considered when evaluating ozone production regimes (Kim et al., 2018).

Even with the inclusion of the additional peaks to the calculated OH reactivity, we still find a missing OH reactivity of 42%. Thus, it is important to investigate the origin of this missing fraction. A correlation can be observed between missing OH reactivity in percentage and OH reactivity from NO_X (R2 = 0.5, Figure 7 A) but not between OH reactivity from NO_X and absolute missing OH reactivity (s^{-1}) (R2 = 0.2, Figure 7 B). This leads us to speculate that there is a consistent presence of unquantified trace gases, likely oxidation products of both anthropogenic and biogenic VOCs as we explored the origin of the unexplored peaks causing missing OH reactivity. In other words, NO_X is relatively well measured with a highly pronounced temporal variation that determines the percentage of missing OH reactivity.

Finally, unaccounted for uncertainty associated with the reaction rate constant estimations described in the method section should be also further explored. For example, to reconcile the averaged missing OH reactivity during the day (10 s^{-1}), it requires $\sim 60 \text{ ppm}$ of

methane but only \sim 4 ppb of isoprene. This clearly demonstrates the importance of rate constant estimation. Indeed, if we apply the reaction rate constant of isoprene with OH ($k_{\rm OH} = 1 \times 10^{-10}$ cm³ molecule⁻¹ s⁻¹at 298 K) to Category II and Category III compounds, then the observed OH reactivity is fully reconciled (Figure S3). Proton ion chemistry may have an intrinsic limitation to quantify highly oxidized OVOCs. Moreover, due to the different inlet configurations for OH reactivity and VOC observations, their contributions towards observed and calculated OH reactivity may not have been consistently evaluated (e.g. Sanchez et al. (2018)). Therefore, a comprehensive analysis along with a dataset from other instrumentation is necessary towards reconciling missing OH reactivity with observational constraints. Finally, it is highly plausible that we may double count for fragmented molecules in the mass spectrum. Although it would not affect concentration evaluation as the intensity of ion signals from the fragmented molecules would be fully accounted for by adding parent ion and fragmented ion signals, the OH reactivity calculated from the fragmented ions is susceptible to underestimation from the assumption that $k_{\rm OH}$ positively correlates with molecular masses.

4. Summary

We present OH reactivity observations at a suburban forest site during the KORUS-AQ field campaign. A comprehensive trace gas dataset including 14 VOCs quantified by PTR-ToF-MS is used to calculate OH reactivity, which only accounts for 36.7 % of the averaged observed OH reactivity.

This study presents a detailed methodology for retrieving OH reactivity contributions from all of the peaks of the PTR-ToF-MS mass spectrum. This decreases the amount of missing OH reactivity as the majority of them have not been accounted towards calculated OH reactivity in

previous studies. First, we converted the raw signals to concentrations using a constant proton transfer reaction rate ($3 \times 10^{-9} \, \mathrm{cm^3 \, s^{-1}}$). Then, we grouped the previously unaccounted peaks into three categories to estimate reaction constants for each compound. The contributions of the unaccounted peaks in the mass spectrum account for a calculated OH reactivity of $\sim 6 \, \mathrm{s^{-1}}$, which decreases missing OH reactivity from 63.3 % to 42.0 %. It is noteworthy that the diurnal variations of observed OH reactivity and calculated OH reactivity from the various groups of trace gases does not have a high variability during the field campaign even though there were several synoptic meteorological configuration changes. This suggests that the reactive trace gas loading is mostly determined by local emission and oxidation processes not influenced by the synoptic meteorological conditions.

In conclusion, this study highlights PTR-ToF-MS as a tool for observationally constraining missing OH reactivity. Further study is required particularly towards characterizing proton reaction rate constants and reaction constants with OH for the many unknown compounds detected on PTR-ToF-MS. In addition, other mass spectrometry techniques, such as nitrate or iodine ion chemistry systems, should be utilized in future studies to complement the PTR technique, which is sensitive to volatile to semi volatile VOCs, to quantify lower volatility compounds and comprehensively constrain OH reactivity contributions from VOCs.

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868 Tables and Figures

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870 Table 1. Description of instrument and measured parameters.

Instrument	Parameters	Measurement Uncertainty (1σ) and lower level of detection limit
Chemical Ionization Spectroscopy - Comparative Reactivity Method (CIMS-CRM)	OH reactivity	16.7% (5 sec-1)
Thermo Scientific 42i	NO	20% (100 ppt)
Cavity Ring Down Spectroscopy	NO ₂	20% (50 ppt)
Thermo Scientific 49i	O ₃	4% (1 ppb)
Lufft 501 C	Temperature	±0.3 °C (NA)
Thermo Scientific 48i TLE	CO	10% (50 ppb)

Thermo Scientific 43i TLE	SO_2	10% (100 ppt)
Mini Tunable Infrared Laser Direct Absorption Spectroscopy (mini-TILDAS) Formaldehyde Monitor(Herndon et al., 2005) (Aerodyne Research, Inc)	HCHO, CH ₄ , CH ₃ OH	5% (few tens ppt)
Proton Transfer Reaction Time of Flight Mass Spectrometer (PTR-TOF-MS 8000, IONICON Analytik, GmbH)	Acetaldehyde, Ethanol, Acetone, Isoprene, MVK + MACR, Methyl ethyl ketone, Benzene, Monoterpenes, Toluene, Furfural, Benzaldehyde, Xylenes, Trimethylbenzenes, Sesquiterpenes	Isoprene 9.8% Benzene 6.9% Toluene 6.5% Monoterpenes 9.2% Xylenes 4.0% Other 16.5% (tens ppt)

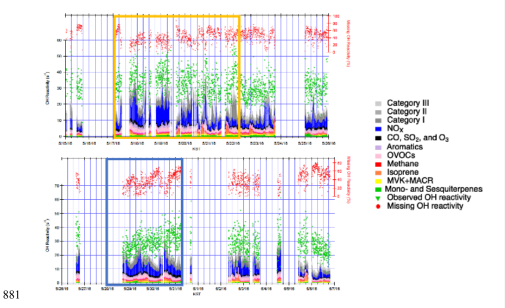


Figure 2. The diurnal average of OH reactivity from 15 May 2016-7 June 2016. The measured and calculated OH reactivity are on the left axis. The blue shading represents uncertainty in the measured OH reactivity. The black bars represent the propagated uncertainty of calculated OH reactivity. The missing OH Reactivity in the percentage scale can be read using the right axis.

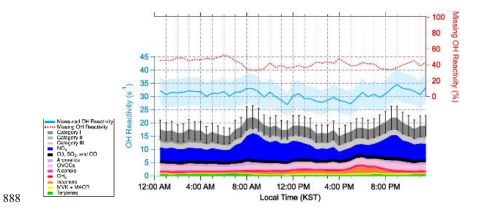


Figure 3. Diurnal averages of OH reactivity during the stagnation period (A) from May 17th – May 22nd in 2016 and the transport period (B) from 28 May – 1 June 2016. The measured and calculated OH reactivity are on the left. The blue shading represents an uncertainty of 16.7% at 1σ . The black bars represent the propagated uncertainty of 20.1% at 1σ from calculated missing OH reactivity. The percent missing OH reactivity is on the right axis.

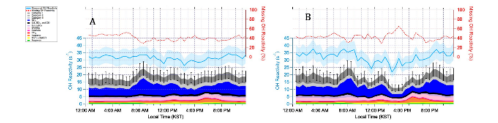


Figure 4. The observed OH reactivity during the overall campaign, stagnation period, and transport period.

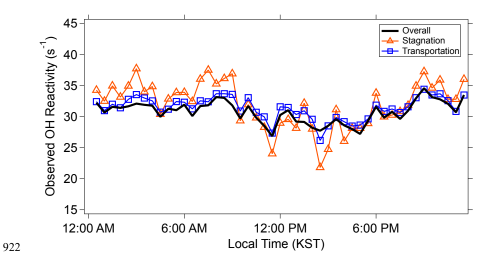


Figure 5. Diurnal profiles for different classes of trace gases during the different periods. A) criteria pollutants NO_x , O_3 , SO_2 , and CO B) Aromatics, C) BVOCs, and D) OVOCs

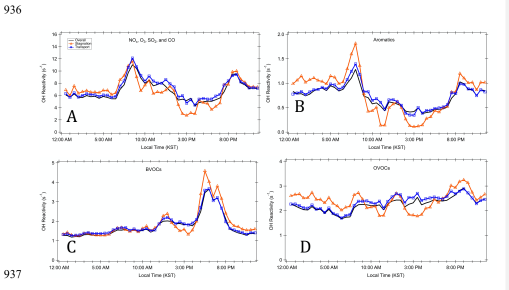


Figure 6. Diurnal averages of the OH reactivity from the compounds in A) Category I, B)

948 Category II and C) Category III

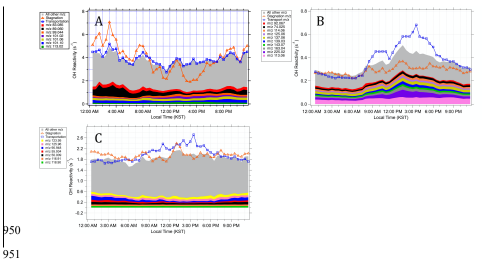


Figure 7. The correlation between A) NOx OH reactivity and absolute missing OH reactivity and B) percent missing OH reactivity

