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Strong anthropogenic control of secondary organic aerosol formation from isoprene in Beijing

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Abstract. Isoprene-derived secondary organic aerosol (iSOA) is a significant contributor to organic carbon (OC) in some forested regions, such as tropical rainforests and the Southeastern US. However, its contribution to organic aerosol in urban areas that have high levels of anthropogenic pollutants is poorly understood. In this study, we examined the formation of anthropogenically influenced iSOA during summer in Beijing, China. Local isoprene emissions and high levels of anthropogenic pollutants, in particular NO_x

and particulate SO_4^{2-} , led to the formation of iSOA under both high- and low-NO oxidation conditions, with significant heterogeneous transformations of isoprene-derived oxidation products to particulate organosulfates (OSs) and nitrooxyorganosulfates (NOSs). Ultra-high-performance liquid chromatography coupled to high-resolution mass spectrometry was combined with a rapid automated data processing technique to quantify 31 proposed iSOA tracers in offline $PM_{2.5}$ filter extracts. The co-elution of the inorganic ions in the

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extracts caused matrix effects that impacted two authentic standards differently. The average concentration of iSOA OSs and NOSs was 82.5 ng m⁻³, which was around 3 times higher than the observed concentrations of their oxygenated precursors (2-methyltetrols and 2-methylglyceric acid). OS formation was dependant on both photochemistry and the sulfate available for reactive uptake, as shown by a strong correlation with the product of ozone (O₃) and particulate sulfate (SO_4^{2-}). A greater proportion of high-NO OS products were observed in Beijing compared with previous studies in less polluted environments. The iSOA-derived OSs and NOSs represented 0.62 % of the oxidized organic aerosol measured by aerosol mass spectrometry on average, but this increased to $\sim 3\%$ on certain days. These results indicate for the first time that iSOA formation in urban Beijing is strongly controlled by anthropogenic emissions and results in extensive conversion to OS products from heterogenous reactions.

1 Introduction

Rapidly developing countries such as China often experience very poor air quality. Beijing regularly experiences periods of very high particle pollution, with annual and 24 hr levels well above the World Health Organization guidelines (Chan et al., 2008; Hu et al., 2014). Premature mortality, as a result of respiratory illness, cardiovascular disease and cancer, has been associated with exposure to poor air quality (Dockery et al., 1993; Pope et al., 2000; Pope and Dockery, 2006; Jerrett et al., 2009; Beelen et al., 2014; Laurent et al., 2014; Ostro et al., 2015). Lelieveld et al. (2015) estimated that 1.36 million premature deaths in China in 2010 were a result of exposure to outdoor air pollution. By far the most dangerous pollutant to health in China is particulate matter less than 2.5 µm in diameter, known as PM_{2.5}, with a recent study suggesting that a 50 % reduction in excess mortality requires a 62 % reduction in PM_{2.5} in the Beijing-Tianjin-Hebei (BTH) region (Hu et al., 2017a).

Previous measurements using aerosol mass spectrometry (AMS) indicate that PM_1 in Beijing is mainly composed of sulfate, nitrate, ammonium and organics (Hu et al., 2016; Zhang et al., 2013). Positive matrix factorization of AMS measurements indicates that oxidized or secondary organic aerosol (SOA) can make up a substantial fraction of the PM_1 mass (>25%), even in urban areas, but the sources of this material are still poorly understood (Q. Zhang et al., 2011; Sun et al., 2018). Hu et al. (2017a) estimated that exposure to SOA was responsible for 0.14 million deaths in China in 2013 based on mass contribution alone, ranging from <1% to 23% source contributions to $PM_{2.5}$ depending on location. Zhang et al. (2017a) used ^{14}C measurements to determine that non-fossil fuel emissions are generally a dominant contributor of secondary organic carbon (SOC) in Beijing, with

a larger contribution in summer as a result of increased biogenic volatile organic compound (VOC) emissions.

Hu et al. (2017b) updated the Community Multiscale Air Quality (CMAQ) model with revised SOA yields and a more detailed description of SOA formation from isoprene oxidation. Removing all anthropogenic pollutants from the model resulted in a huge drop in isoprene SOA concentrations, indicating that controlling anthropogenic emissions would result in a reduction of both anthropogenic and biogenic SOA. The predicted SOA was dominated by isoprene in summer across China and in four cities (Beijing, Guangzhou, Shanghai and Chengdu) with concentrations of up to $30 \,\mu \mathrm{g} \,\mathrm{m}^{-3}$ in Beijing. However, there is currently very little observational evidence to support such high SOA mass concentrations from isoprene oxidation in these Chinese cities. The widely used SOA tracer method (Kleindienst et al., 2007) has been extensively utilized to estimate the fraction of isoprene-derived SOA (iSOA) across China. Ding et al. (2014) studied SOA at 14 Chinese sites and found that iSOA dominated the apportioned SOA mass ($46 \pm 14\%$), contributing between 0.4 and $2.17 \,\mu \text{g m}^{-3}$ with an average of $1.59 \,\mu \text{g m}^{-3}$ in Beijing. However, only a very limited subset of VOC precursors were included, and this method fails to account for heterogeneous formation processes. To overcome some of these limitations, Q. Wang et al. (2017) used tracer-based source apportionment of PM_{2.5} with positive matrix factorization in the Pearl River Delta region during summer. They identified an iSOA factor that contributed up to 4 µg m⁻³ in Guangzhou and up to 11 % of the total SOC.

A multitude of studies have examined iSOA formation (Pandis et al., 1991; Edney et al., 2005; Kroll et al., 2006; Dommen et al., 2006; Kleindienst et al., 2006; Ng et al., 2006, 2008; Surratt et al., 2007a, b, 2008, 2010; Paulot et al., 2009; Chan et al., 2010; Chhabra et al., 2010; T. B. Nguyen et al., 2011, 2014, 2015; H. Zhang et al., 2011, 2012; Lin et al., 2013; Xu et al., 2014; Krechmer et al., 2015; Clark et al., 2016; Riva et al., 2016a, b); however, the magnitude of iSOA formed can be vastly different from study to study (Carlton et al., 2009). Furthermore, there have been limited field measurements undertaken to establish if these estimates are representative of urban environments (Wang et al., 2018; Li et al., 2018; Glasius et al., 2018; Le Breton et al., 2018; Hettiyadura et al., 2018, 2019; Rattanavaraha et al., 2016; Budisulistiorini et al., 2013).

iSOA formation during the daytime is dominated by reaction with hydroxyl radicals (OH), with the concentrations of NO having a strong influence on the reaction products (Wennberg et al., 2018, and references therein). Under low-NO conditions, isoprene peroxy radicals (RO₂) can react with hydroperoxy radicals (HO₂) to form isoprene hydroxyhydroperoxides (ISOPOOH). The ISOPOOH isomers can react further with OH to form isoprene epoxydiol isomers (β - or δ -IEPOX; Paulot et al., 2009), which can undergo uptake into acidified sulfate particles to form 2-methyltetrol organosulfates (2-MT-OS; Surratt et al., 2010; Lin et al.,

2012; T. B. Nguyen, 2014). Under high-NO conditions, isoprene RO2 can react with NO to form alkoxy radicals (RO), producing methacrolein (MACR) and methyl vinyl ketone (MVK) as the main reaction products. The reaction of MACR with OH and the subsequent addition of NO₂ leads to methylacrloylperoxynitrate (MPAN), which reacts with OH to produce hydroxymethyl-methyl- α -lactone (HMML; Nguyen et al., 2015) or methacrylic acid epoxide (MAE; Lin et al., 2013). HMML is thought to be the more abundant product compared with MAE (Nguyen et al., 2015). The subsequent uptake of HMML into wet sulfate aerosols is proposed to lead to either 2-methylglyceric acid (2-MG) or its organosulfate derivative (2-MG-OS), as well as their dimers and higher order oligomers (Surratt et al., 2006, 2010; Lin et al., 2013; Nguyen et al., 2015). Recently, Schwantes et al. (2019) showed that under ambient conditions the formation of SOA from low-volatility nitrates and dinitrates, formed via reactions of isoprene-derived RO2 with NO, is also important. Chamber-derived SOA yields for OH chemistry are variable depending on the experimental conditions, but they are generally low (< 10 %). The addition of acidified sulfate aerosol, accounting for wall losses and using more atmospherically relevant radical lifetimes can lead to significantly higher SOA yields in chamber studies (Surratt et al., 2010; Lin et al., 2012; Gaston et al., 2014). However, recent work has revealed that isoprene SOA formation can be suppressed when viscous organic coatings are present on acidified sulfate aerosol, as they impede the multiphase chemistry of IEPOX that yields additional SOA (Riva et al., 2016b, 2019; Zhang et al., 2018).

Observations using aerosol mass spectrometry (AMS) indicate that IEPOX-derived SOA can make up a significant fraction of organic aerosol in isoprene-rich environments, such as Borneo (23 %; Robinson et al., 2011), the Amazon (34%; Chen et al., 2015) and the Southeastern US (33 %-40 %; Budisulistiorini et al., 2013, 2016; Rattanvaraha et al., 2017). Hu et al. (2015) compared previous AMS studies and found a magnitude lower average IEPOX-SOA signal in urban studies ($f_{C_5H_6O}$ of 0.17 %) compared with those in isoprene-rich regions ($f_{C_5H_6O}$ of 2.2%). The average IEPOX-SOA concentration measured in Nanjing, a polluted city in eastern China, in August 2013 was $0.33 \,\mathrm{\mu g}\,\mathrm{m}^{-3}$ (Zhang et al., 2017b). This represented only 3.8 % of the total OA, indicating that there is limited formation of IEPOX under high-NO_x conditions (average NO_x of 21 ppb). He et al. (2018) found higher concentrations of low-NO isoprene SOA tracers (average of 121 ng m⁻³) than high-NO iSOA tracers (average of 9 ng m⁻³) at a regional background site (Wanqingsha) situated within the heavily polluted Pearl River Delta region. Only two high-NO iSOA tracers were measured (2-MG and 2-MG-OS), which could lead to a significant underestimate of the strength of the high-NO_x pathway. Wang et al. (2018) measured a range of OSs at a regional site 38 km northeast of Beijing during May and June 2016. Isoprene-derived OSs ranged from 0.9 to 20 ng m⁻³ with a mean isoprene-derived OS concentration of 14.8 ng m⁻³. In both of these studies, the ratio of the average concentration of the commonly used OS tracers from the low-NO versus the high-NO pathways was close to 1.5 (2-MT-OS:2-MG-OS; Beijing was 1.47 and Wanqingsha was 1.57), indicating that low-NO oxidation chemistry can even play a significant role in iSOA formation in polluted environments.

SOAs formed from anthropogenic and other biogenic (monoterpenes and sesquiterpenes) sources have also been studied. Thousands of organic species including hundreds of OS and nitroxy OS (NOS) species have been identified in studies from a range of precursors using ultra-highperformance liquid chromatography coupled to ultra-highresolution mass spectrometry equipped with electrospray ionization (UHPLC/ESI-HR-MS) from ambient aerosol samples (X. Wang et al., 2016, 2017). Brüggemann et al. (2019) quantified both monoterpene OS (MT-OS) and sesquiterpene OS (SQT-OS) species in Melpitz, Germany, and Wangdu, China, with authentic standards. They found median daytime concentrations of 52 MT-OS species of 12.15 and 38.19 ng m⁻³ for Melpitz and Wangdu respectively. For the five SQT-OS species, the median concentrations were 0.3 and 3.90 ng m⁻³ for daytime concentrations in Melpitz and Wangdu respectively, which are much lower values than the iSOA OS species quantified in this study. Riva et al. (2016c) identified OS species from the photooxidation of C₁₀-C₁₂ alkanes, which were then characterized in ambient aerosol samples collected in Lahore, Pakistan, and Pasadena, CA, USA. High concentrations of OS species were identified in Lahore, with the largest observed concentration arising from a cyclodecane OS species (C₁₀H₁₆O₇S) with a concentration of $35.93 \,\mathrm{ng} \,\mathrm{m}^{-3}$.

The lack of molecular-level measurements of iSOA in highly polluted urban areas makes it difficult to determine the role of isoprene in summer haze episodes in Beijing. To investigate the formation of iSOA in Beijing, offline PM_{2.5} filter samples were collected during summer 2017 as part of the Atmospheric Pollution and Human Health programme (Shi et al., 2019). The filters were extracted and then screened using a sensitive and selective high throughput method based on UHPLC/ESI-HR-MS. High time resolution filter sampling allowed for the formation and evolution of iSOA to be studied, with observed concentrations strongly controlled by levels of anthropogenic pollutants.

2 Experimental

2.1 PM_{2.5} filter sampling and extraction

Aerosol samples were collected between 18 May and 24 June 2017 at the Institute of Atmospheric Physics (IAP) in Beijing, China. This sampling was part of the Sources and Emissions of Air Pollutants in Beijing (AIRPOLL-Beijing)

project, as part of the wider Atmospheric Pollution and Human Health in a Chinese Megacity (APHH-Beijing) programme (Shi et al., 2019). PM_{2.5} filter samples were collected using an Ecotech HiVol 3000 (Ecotech, Australia) high-volume air sampler with a selective PM_{2.5} inlet and a flow rate of 1.33 m³ min⁻¹. Filters were baked at 500 °C for 5h before use. After collection, samples were wrapped in foil, stored at -20 °C and then shipped to the laboratory for offline analysis. Samples were collected at a height of 8 m on top of a building in the IAP complex. Samples were collected every 3 h during the day (between approximately 08:30 and 17:30 LT) and then one sample was collected overnight (between approximately 17:30 and 08:30 LT). Hourly samples were also taken on certain high-pollution days towards the end of the sampling period. A number of 24 h samples were also collected using a Digitel high-volume PM2.5 sampler at the same location.

The extraction of the organic aerosol from the filter samples was based on the method in Hamilton et al. (2008). Initially, an eighth of the filter was cut up into roughly 1 cm² pieces and stored in a vial. Next, 4 mL of LC-MS-grade H₂O was added to the sample and left for 2 h. The samples were then sonicated for 30 min. A small subset (three) of the filter samples were also extracted via orbital shaker, and no appreciable difference was found in the concentrations of the iSOA tracers compared with sonication. Using a 2 mL syringe, the water extract was then pushed through a 0.22 µm filter (Millipore) into another sample vial. An additional 1 mL of water was added to the filter sample and then extracted through the filter to give a combined aqueous extract. This extract was reduced to dryness using a vacuum solvent evaporator (Biotage, Sweden). The dry sample was then reconstituted in 1 mL of 50: 50 MeOH: H₂O solution for offline chemical analysis.

2.2 Ultra-high-performance liquid chromatography tandem mass spectrometry (UHPLC-MS²)

The water-soluble fractions of the filter samples were analysed using UHPLC full-scan ddMS² via an Ultimate 3000 UHPLC (Thermo Scientific, USA) coupled to a Q Exactive Orbitrap MS (Thermo Fisher Scientific, USA) with heated electrospray ionization (HESI). The UHPLC method uses a reversed-phase 5 µm, 4.6 mm ×100 mm, Accucore column (Thermo Scientific, UK) held at 40 °C. The mobile phase consists of LC-MS-grade H₂O and 100 % MeOH (Fisher Chemical, USA). The H₂O was acidified using 0.1 % formic acid to improve peak resolution. The injection volume was 2 μL. The solvent gradient was held for 1 min at 90:10 H₂O: MeOH; changed linearly to 10: 90 H₂O: MeOH over 9 min; held for 2 min at this gradient before returning it to 90:10 H₂O: MeOH over 2 min; and then held at 90:10 for the remaining 2 min, with a flow rate of $300 \,\mu\text{L min}^{-1}$. Due to the wide range of compounds studied, poor retention was observed for some species (retention time, RT, of less than 0.8 min). These species closely eluted to the dead time of the column where inorganic sulfate ions eluted (0.67 min). To check for ionization artefacts, an aqueous solution containing 20 ppm ammonium sulfate, 1 ppm 2-methyltetrol and 1ppm 2-methylglyceric acid was run under the same conditions as the filter samples to check for organosulfate formation (2-MT-OS and 2-MG-OS respectively). No MG-OS formation was observed, and less than 0.5 % conversion was seen for 2-MT. Therefore, this ruled out adduct formation for the two most important iSOA species, 2-MT-OS and 2-MG-OS; however, due to the lack of authentic standards and the complexity of the samples, adduct formation throughout the entire chromatogram could still be occurring. At this stage, there is not enough evidence to definitively say if adducts are forming or not. The mass spectrometer was operated in negative mode using full-scan ddMS². The scan range was set between 50 and $750 \, m/z$, with a resolution of 70 000. The ESI voltage was 4 kV, with capillary and auxiliary gas temperatures of 320 °C. The number of the most abundant precursors for MS² fragmentation per scan was set to 10. The samples were run in batches of 70, in a repeating sequence of 5 samples followed by one blank, and each filter sample was run only once. The calibrations were run separately after the samples were finished; the following sequence was used for the calibration runs: $(3 \times \text{ same concentration})$ multiplied by the number of standards in the calibration curve from the lowest concentration to the highest followed by two blanks. The quantification method will be discussed in Sect. 3.3.

2.3 Construction of accurate mass library

A mass spectral library was built using the compound database function in TraceFinderTM 4.1 General Quan software (Thermo Fisher Scientific, USA). Each compound was input into the compound library in the generic form: $C_cH_hO_oN_nS_s$ (where c, h, o, n and s represent the number of carbon, hydrogen, oxygen, nitrogen and sulfur atoms respectively). From the literature, species were identified, searched for in the ambient samples according to their accurate mass, and then the retention time (RT) of each isomer was obtained. Using previously observed iSOA products from the literature, extracted ion chromatograms were plotted for each m/zvalue from a small subset of ambient samples, and the RT of the observed species/isomer was obtained. For most of the OS species in this study, the separation was not good enough to see individual isomers and only one peak was observed, which was added to the library. For the NOS species, individual isomers could be resolved, and each isomer was added to the library based on its RT. The accurate masses, RT and literature references for iSOA tracers are shown in Table 2.

2.4 Automated method for SOA tracer analysis

The UHPLC/ESI-HR-MS data for each ambient sample and standard were analysed using TraceFinderTM software.

TraceFinderTM extracted the OS/NOS tracer peak areas from each ambient sample chromatogram using the library based on the RT and accurate mass. The mass tolerance of the method was set to 2 ppm, and the RT window was set to 30 s, although for species with multiple isomers present, the integration was checked to make sure the same peaks were not being integrated twice, and the window changed accordingly. The peak tailing factor was set to 2.0 to reduce the integration of the peak tails. The minimum signal-to-noise ratio (S/N) for a positive identification was set to 3.0. Using the output from TraceFinderTM, an in-house R code script was developed to combine the identified species and peak areas with the correct filter sampling date/time midpoint and volume of air sampled. Calibration curves from the standards were then obtained (as discussed in Sect. 3.3), and the intercept and gradient were inputted to quantify the iSOA tracer concentrations in the extract. These quantified values were then converted to the mass on the whole filter, divided by the volume of air sampled for that filter sampling period and converted to units of nanograms per cubic metre (ng m^{-3}). Higher time resolution data were averaged to the filter sampling times. It should be noted that MS² was used to check that the iSOA species fragmented to give typical OS fragment ions.

2.5 Hydrophilic liquid interaction chromatography (HILIC)

A subset of filters (n = 15) were also analysed at the University of North Carolina (UNC) using a newly developed HILIC method interfaced to high-resolution quadrupole time-of-flight mass spectrometry equipped with ESI (i.e. HILIC/ESI-HR-QTOFMS; Cui et al., 2018). Briefly, filters were extracted with 22 mL of LC/MS-grade methanol by 45 min of sonication; the samples were first extracted for 23 min, the water bath was replaced with cool water, and then the samples were extracted again for 22 min. This was done to make sure that the water bath contained within the sonicator did not exceed 30 °C. Extracts were filtered through polypropylene membrane syringe filters in order to remove insoluble filter fibres and soot particles. The extracts were dried under a gentle stream of nitrogen gas. Dried methanol extracts were reconstituted with 150 µL of 95 : 5 (v/v) LC/MS-grade acetonitrile/Milli-Q water. Operating details of the HILIC/ESI-HR-QTOFMS used for these samples are also summarized by Cui et al. (2018).

2.6 Gas chromatography–mass spectrometry

Details of the measurement procedure used can be found elsewhere (Fu et al., 2010). Briefly, filter samples were extracted with dichloromethane/methanol $(2:1\ v/v)$, filtered through quartz wool packed into a Pasteur pipette, concentrated using a rotary evaporator under vacuum and blown down to dryness with pure nitrogen gas. The extracts were derivatized and diluted with n-hexane contain-

ing the internal standard prior to GC-MS analysis. Separation was performed on a fused silica capillary column (DB-5MS: $30\,\mathrm{m}\times0.25\,\mathrm{mm}\times0.25\,\mathrm{\mu m}$). The MS detection was conducted in electron ionization (EI) mode at $70\,\mathrm{eV}$, scanning from 50 to 650 Da. Individual compounds were identified via comparison of the mass spectra with those of authentic standards or literature data. 2-Methylglyceric acid, C₅-alkene triols (the sum of *cis*-2-methyl-1,3,4-trihydroxy-1-butene, *trans*-2-methyl-1,3,4-trihydroxy-1-butene and 3-methyl-2,3,4-trihydroxy-1-butene) and 2-methyltetrols (the sum of 2-methylthreitol and 2-methylerythritol) were quantified using the response factor of *meso*-erythritol. Field blank filters were treated as the real samples for quality assurance. Target compounds were not detected in the blanks.

2.7 High-resolution aerosol mass spectrometry measurements

The size-resolved non-refractory submicron aerosol species at the same site were measured by an Aerodyne high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) at a time resolution of 5 min. The elemental ratios of hydrogen to carbon (H:C) and oxygen to carbon (O:C) of OAs were determined, and the sources of OAs were analysed with positive matrix factorization. Seven OA factors were identified in summer, including two primary factors; hydrocarbon-like OA (HOA); cooking OA (COA); and three oxidized OA factors with increasing degrees of oxidation, OOA1 (O:C = 0.53), OOA2 (O:C = 0.74) and OOA3 (O:C = 1.18).

2.8 Iodide CIMS

A time-of-flight chemical ionization mass spectrometer (ToF-CIMS; Lee et al., 2014; Priestley et al., 2018) using an iodide ionization system coupled to a filter inlet for gases and aerosols (FIGAERO) was deployed here to make near-simultaneous, real-time measurements of both the gas- and particle-phase chemical composition. The instrument was originally developed by Lopez-Hilfiker et al. (2014) and is described and characterized in more detail by Bannan et al. (2019). The experimental set-up employed by the University of Manchester ToF-CIMS is described in Zhou et al. (2018). Only gas-phase data are presented herein.

Field calibrations were regularly carried out using a known concentration of formic acid in gas mixtures produced in a custom-made gas-phase manifold. A range of other species were calibrated for after the campaign, and relative calibration factors were derived using the measured formic acid sensitivity during the in situ calibrations (Bannan et al., 2015). Offline calibrations after the field work campaign were performed specific to the isoprene oxidation species observed here. IEPOX ($C_5H_{10}O_3$) synthesized by the University of North Carolina, Department of Environmental Sciences and Engineering, was specifically calibrated for. Known concen-

trations were deposited on the FIGAERO filter in various amounts and thermally desorbed using a known continuous flow of nitrogen over the filter. For the isoprene nitrate, $C_5H_9NO_4$, there was no direct calibration source available, and concentrations using the calibration factor of $C_5H_{10}O_3$ are presented here.

2.9 Gas-phase measurements

Additional gas-phase measurements were collected at the site from an elevated inlet at 8 m. Data included nitrogen oxide (NO) measured by chemiluminescence with a Thermo Scientific Model 42i NO_x analyser and nitrogen dioxide (NO₂) measured using a Teledyne Model T500U cavityattenuated phase-shift (CAPS) spectrometer. The sum of the NO_v species was measured using a Thermo Scientific Model 42C NO_x analyser and a heated molybdenum converter at the sample inlet. The molybdenum converter reduces NO_v compounds to NO and allows for measurement by chemiluminescence. Ozone (O₃) was measured using a Thermo Scientific Model 49i UV photometric analyser. All instruments were calibrated throughout the measurement period using a "zero" or "background" calibration with a Sofnofil/charcoal trap. Span (high-concentration) calibrations were carried out using gas standards. Both the Thermo Scientific 42i and 42C instrument calibrations are traceable to the National Physical Laboratories (NPL) NO scale. The meteorological variables of wind speed, wind direction, relative humidity (RH) and temperature were measured at 102 m on the IAP 325 m meteorological tower.

Observations of VOCs were made using a dual-channel GC with flame ionization detection (DC-GC-FID). Air was sampled at $30\,\mathrm{L\,min^{-1}}$ at a height of 5 m, through a stainless-steel manifold (1/2'' internal diameter). Next, $500\,\mathrm{mL}$ subsamples were taken, dried using a glass condensation finger held at $-40\,^{\circ}\mathrm{C}$ and then pre-concentrated using a Markes Unity 2 pre-concentrator on a multi-bed O_3 precursor adsorbent trap (Markes International Ltd). These samples were then transferred to the GC for analysis following the methods described by Hopkins et al. (2011).

Further details on the following additional gas-phase instrumentation can be found in the Supplement and in Shi et al. (2019). Isoprene was also measured at a height of $\sim 102~\text{m}$ using a Voice200 selected ion flow tube mass spectrometer (SIFT-MS, Syft Technologies, Christchurch, New Zealand). OH, HO2 and RO2 concentrations were measured using fluorescence assay by gas expansion (FAGE; Whalley et al., 2010, 2018), and NO3 concentrations were measured using broadband cavity enhanced absorption spectrometry (Le Breton et al., 2014).

3 Results and discussion

The field campaign was conducted at the Institute of Physics, Beijing, situated between the third and fourth ring roads (Shi et al., 2019). The site is typical of central Beijing: it is surrounded by residential and commercial properties, and is near several busy roads. It is also close to several green spaces, including a tree-lined canal to the south and the Olympic forest park to the northeast, that provide sources for local isoprene emissions.

3.1 Isoprene gas-phase concentrations and loss processes

Isoprene was measured hourly using the DC-GC-FID between 18 May and 20 June 2017, and the observed concentrations are shown in Fig. 1 alongside NO, NO₂ and O₃. The mean mixing ratio of isoprene was 0.53 ppb, with a maximum of 2.9 ppb on 16 June 2017. The ambient temperature ranged from 16 to 38 °C. Daytime isoprene mixing ratios increased with temperature, with all isoprene mixing ratios above 1 ppb occurring when the temperature was above 25 °C. The average diurnal profile of isoprene in Fig. 3a shows low values overnight (< 50 ppt) with a rapid increase at 06:00 LT that reaches a maximum of around 1 ppb by the afternoon. The mixing ratio rapidly decreased after 18:00 LT and returned to very low values by around 22:00 LT. There was a strong correlation between the isoprene mixing ratio measured at 8 m using the DC-GC and at 102 m using the SIFT-MS ($R^2 = 0.77$). Therefore, the SIFT-MS measurements were used to investigate the correlation with iSOA tracers when no DC-GC data were available. The slope of the linear fit between the two datasets was 0.67, indicating a loss of around 30 % of the isoprene during transport from the ground to the tower (100 m).

Using the average observed diurnal profiles of the main atmospheric oxidants, OH, O₃ and NO₃ (shown in Supplement Fig. S1), and isoprene (Fig. 3a), the isoprene loss rate was calculated (rate of loss = k_{ox} [oxidant][isoprene]) and is shown in Fig. 4a. The IUPAC rate constants that were used in the calculation for NO₃, O₃ and OH were 7×10^{-13} , 1.27×10^{17} and 1×10^{10} cm³ molecule⁻¹ s⁻¹ respectively. The percentage contribution of each oxidant to the average diurnal isoprene loss rate is shown in Fig. 4b. During the day, OH is responsible for over 90 % of isoprene loss, with NO₃ becoming relatively more important from 18:00 until around 03:00 LT, although the amount of isoprene available to react rapidly decreases during this time period. OH chemistry is still an important loss route at night (> 30 %) owing to nighttime OH sources, such as the ozonolysis of alkenes (Lu et al., 2014). The loss of isoprene via ozonolysis, however, is a minor route, contributing less than 15 %. During the daytime (10:00-15:00 LT), the lifetime of isoprene was found to be around 20 min on average, increasing to a maximum of around 6 h at 03:00 LT. While the high levels of oxidants

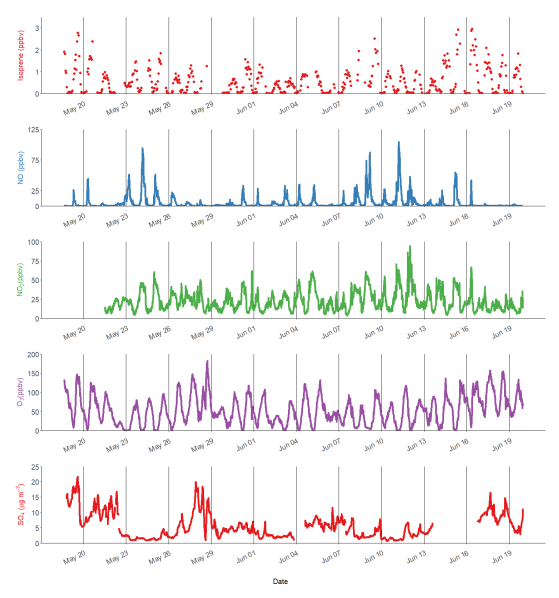


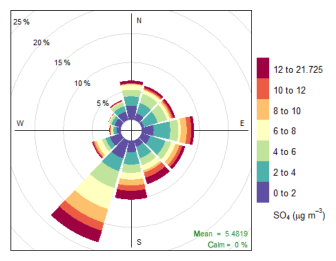
Figure 1. Time series of isoprene, nitric oxide (NO), nitrogen dioxide (NO₂), ozone (O₃) and particulate sulfate (SO₄). The black lines represent midnight every 72 h.

lead to a short isoprene lifetime during the day, the ambient concentrations of isoprene are still maintained at the parts per billion level. This indicates that there are significant local emissions of isoprene impacting the measurement site; therefore, there is a high potential for the formation of iSOA in this urban environment.

3.2 Anthropogenic tracers

A range of gas-phase anthropogenic tracers were measured during the campaign, as discussed in Shi et al. (2019). Figure 1 shows the time series of NO, NO₂, O₃ and particulate sulfate during the part of the campaign analysed in this study. Table 1 shows the average, maximum and minimum concentrations of these anthropogenic pollutants. NO mixing ratios

ranged from less than 10.1 to 104 ppbv with a mean concentration of 5.1 ppb during the filter sampling period. The highest concentrations generally occurred in the morning between 04:00 and 07:00 LT and steadily decreased during the day. On some days, the mixing ratio of NO was very low in the afternoon as a result of reaction with O₃ and other unknown sinks (Newland et al., 2020). The mean mixing ratio of NO₂ was 22.3 ppbv, which was much higher than NO, with a range from 3.7 to 95 ppbv. NO₂ peaked between 06:00 and 07:00 LT and decreased to a minima at 14:00 LT before steady increasing until about 20:00 LT. High afternoon concentrations of O₃ (>80 ppb) were found on most days, with a maximum observed mixing ratio of 182 ppbv. Night-time O₃ levels were much lower due to reduced photochemistry



Frequency of counts by wind direction (%)

Figure 2. Pollution rose plot (openair) of particulate sulfate measured by AMS for the sampling period, highlighting the wind conditions under which the highest concentrations of sulfate occur.

Table 1. Supplementary anthropogenic pollutants measured during the sampling period analysed in this study.

Pollutant	Mean ± SD	Max	Min
O ₃ (ppbv)	54.0 ± 37.5	181.8	2.0
NO (ppbv)	5.1 ± 11.3	104.1	0.1
NO ₂ (ppbv)	22.3 ± 13.0	94.5	3.7
$SO_4 (\mu g m^{-3})$	5.5 ± 4.1	21.7	0.7

and reaction with NO, although O_3 levels were maintained above 40 ppbv on some nights, as shown in Fig. 1. Particulate sulfate concentrations measured by AMS are also shown in Fig. 1. Sulfate ranged from 0.7 to $21.7 \,\mu g \, m^{-3}$ with an average of $5.5 \,\mu g \, m^{-3}$. The time series shows a number of periods of high sulfate concentrations, and these generally matched periods of increased $PM_{2.5}$ (see Fig. 9). Figure 2 shows the wind direction-dependent concentrations of particulate sulfate for the sampling period in a pollution rose plot (R, openair package). There is a strong source of sulfate from the south of the sampling site, which is enhanced under the highest wind speeds. Previous studies have shown a strong source of pollution from the southwest of Beijing, which is where many industrial factories are located (Wang et al., 2005).

3.3 Isoprene SOA in Beijing

Using the high throughput screening method described, the peak areas of 31 potential isoprene-derived OSs and NOSs, which are known iSOA tracers, were measured in 132 PM_{2.5} filter extracts. The full list of iSOA tracers, along with their measured m/z values and molecular formulas, is shown

in Table 2, ordered by descending average concentration (weighted by filter sampling time and reported in nanograms per cubic metre) during the campaign. The isoprene SOA tracers identified in this study are correlated towards themselves as well as common anthropogenic tracers in a correlation plot (R, openair, corPlot), which is shown in Fig. 5. The correlation plot highlights the correlations of the iSOA tracers to each other as well as the moderate to strong correlations towards some of the anthropogenic pollutants, as discussed in further sections.

3.4 Quantification of isoprene OS tracers

Initially, two synthesized isoprene-derived OS standards (2-MT-OS and 2-MG-OS; Cui et al., 2018; Rattanavaraha et al., 2016) were used to produce calibration curves. Both standards gave strong linear calibration curves (R^2 of 0.980 and 0.996 respectively) across an appropriate range of concentrations for the peak areas in the samples. The gradient obtained for the 2-MT-OS standard was \sim 4 times higher than that of the 2-MG-OS, as shown in Fig. S2. To investigate the potential for matrix effects from the large amounts of inorganic sulfate, nitrate and other particulate components that co-elute due to the poor retention of OS in reversed-phase UHPLC, standard addition calibrations were used. Five-point standard addition calibrations were run on six different filter extracts, covering both daytime and night-time samples during periods of both high and low concentrations of iSOA species. Thus, this provides a representative sample of filters for the entire sampling period. A total of 50 µL of filter sample extract and 50 µL of the calibrant solution were combined, giving a dilution factor of 2. The five-point calibration range of standard added to each sample was between 0 and 3 ppm for 2-MG-OS and between 0 and 1 ppm for 2-MT-OS. Two examples of the standard addition calibrations are shown in Figs. S3 (2-MG-OS) and S4 (2-MT-OS), with good linear fits observed (R^2 of 0.997 and 0.997 respectively). A strong matrix effect was observed for 2-MT-OS: the concentration measured by standard addition calibration was 8.6-10 times higher than that measured using the external calibration, which was carried out on the same day. In contrast, 2-MG-OS showed a much lower matrix effect: the concentrations were only 1.1–1.5 times higher when using the standard addition calibration. A further comparison using camphorsulfonate, which has a longer RT (3.74 min) and, therefore, does not experience high inorganic ion concentrations in the source, showed no matrix effects when using standard addition. Tables S1 and S2 in the Supplement provide a comparison of the concentrations calculated from the standard additions and the two external calibrations. Table S1 shows the concentration of 2-MT-OS in the three filter sample extracts (144, 204 and 208) calculated via the standard addition of 2-MT-OS to the filter sample extract and via the external calibrations using both 2-MT-OS and 2-MG-OS. The ratio of the standard addition to the external calibrations then gives an

Table 2. Molecular formulas, negative ion masses, retention times (RT) and time-weighted means (ng m $^{-3}$) for the entire sampling period as well as the original reference where each proposed iSOA tracer was found. BD denotes "below detection". The estimated uncertainties are 60%, as discussed in Sect. 3.4, accounting for the use of the matrix correction factors.

Isoprene tracer	[M-H] ⁻¹	RT (min)	Time- weighted mean (ng m ⁻³)	Maximum (ng m ⁻³)	Minimum (ng m ⁻³)	Reference
C ₂ H ₄ O ₆ S	154.9656	0.73	38.4	366.1	BD	Surratt et al. (2008)
$C_5H_{10}O_6S$	197.0125	0.79	28.7	336.2	0.25	Surratt et al. (2007a)
$C_5H_{10}O_5S$	181.0176	0.93	26.5	448.5	2.91	Nguyen et al. (2010)
$C_4H_8O_6S$	182.9969	0.73	21.7	229.1	0.50	Riva et al. (2016a)
$C_4H_8O_7S$	198.9918	0.73	21.5	180.5	0.32	Surratt et al. (2007a)
$C_3H_6O_5S$	152.9863	0.73	20.5	327.9	0.98	Surratt et al. (2008)
$C_3H_6O_6S$	168.9812	0.73	14.5	137.7	0.25	Surratt et al. (2008)
$C_5H_8O_7S$	210.9918	0.73	14.0	136.4	0.27	Surratt et al. (2008)
$C_5H_{11}O_9NS$	260.0082	0.86	12.6	154.1	0.10	Surratt et al. (2008)
$C_5H_{12}O_7S$	215.0231	0.71	11.8	110.9	0.77	Surratt et al. (2008)
$C_5H_{10}O_7S$	213.0075	0.73	10.6	104.7	0.38	Surratt et al. (2008)
$C_5H_9O_{10}NS$	273.9874	0.94	9.17	53.8	BD	Nestorowicz et al. (2018)
$C_4H_8O_5S$	167.0019	0.73	9.10	114.5	0.68	Surratt et al. (2007a)
$C_5H_8O_5S$	179.0020	0.85	6.59	144.2	0.43	Riva et al. (2016a)
$C_5H_{10}O_5S$	181.0176	1.24	4.90	36.3	1.21	Riva et al. (2016a)
$C_5H_{10}O_8S$	229.0024	0.75	4.59	40.9	BD	Nestorowicz et al. (2018)
$C_5H_8O_9S$	242.9816	0.64	1.55	13.9	BD	Nestorowicz et al. (2018)
$C_5H_{10}O_{11}N_2S$	304.9783	2.18	1.04	8.62	BD	Surratt et al. (2008)
$C_{10}H_{20}O_8S$	299.0806	1.65	1.01	8.38	BD	Riva et al. (2016b)
$C_5H_{10}O_{11}N_2S$	304.9783	1.89	0.83	7.69	BD	Surratt et al. (2008)
$C_8H_{14}O_{10}S$	301.0235	0.73	0.57	4.16	BD	Surratt et al. (2007a)
$C_5H_{10}O_{11}N_2S$	304.9783	1.56	0.42	2.90	BD	Surratt et al. (2008)
$C_{10}H_{18}O_{7}S$	281.0701	1.03	0.33	6.76	BD	Riva et al. (2016b)
$C_5H_{10}O_{11}N_2S$	304.9783	3.60	0.31	3.32	BD	Surratt et al. (2008)
$C_5H_9O_{13}N_3S$	349.9783	5.90	0.19	2.04	BD	Ng et al. (2008)
$C_{10}H_{18}O_8S$	297.0650	0.75	0.14	5.25	BD	Riva et al. (2016)
$C_5H_{11}O_8NS$	244.0133	1.93	0.11	1.46	BD	Nestorowicz et al. (2018)
$C_5H_9O_{13}N_3S$	349.9783	5.49	0.02	0.17	BD	Ng et al. (2008)
$C_5H_9O_{13}N_3S$	349.9783	5.34	0.008	0.10	BD	Ng et al. (2008)
$C_5H_{12}O_8S$	231.0180	0.75	0.005	0.50	BD	Riva et al. (2016b)
$C_{10}H_{20}O_{9}S$	315.0755	1.46	0.002	0.21	BD	Riva et al. (2016b)

estimate of the under- or overestimation that the external calibrations cause when calculating the concentration of 2-MT-OS in the samples. Both of the external calibrations would lead to an underestimation of the concentration of 2-MT-OS in the filter samples. 2-MG-OS provided a closer quantification of 2-MT-OS in the samples, with an average factor of 2.3 underestimation, whereas the 2-MT-OS external calibration gives a sample concentration that is a factor of 10 lower than the concentration determined by standard addition.

It is not realistic to carry out standard addition calibrations for all samples and all SOA tracers. When the 2-MG-OS external calibration was used to predict the 2-MT-OS concentrations during the standard addition experiments, the concentrations were within a factor of 1.5–2.5. Therefore, the 2-MG-OS external calibration was used as a proxy for

all isoprene SOA tracers, with scaling factors applied to account for matrix effects (1.33 for 2-MG-OS, 2.33 for 2-MT-OS and an average of 1.83 used for all other OSs). Therefore, we estimate an uncertainty on our measured concentrations of 60%, this uncertainty was calculated to account for the difference in the measured correction factors used when correcting for the matrix effects. The uncertainty was calculated as 2σ of the six values used to calculate the average correction factor of 1.83. The matrix effects identified in this study are likely due to the extracted samples being a complex mixture of different compounds, including a high proportion of inorganic ions that are extracted into water. This is likely to change the surface tension of the droplet produced in the ionization source and the ion distribution. Further work is needed to fully understand the reasons for

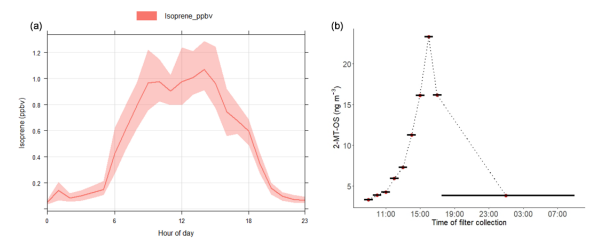


Figure 3. (a) Average diurnal profile of the isoprene mixing ratio measured using DC-GC-FID. **(b)** Diurnal profile of 2-methyltetrol sulfate (2-MT-OS) in particulate matter (PM_{2.5}) collected on filters hourly over the period from 11 to 12 June 2017. Black lines indicate the length of the filter sampling period.

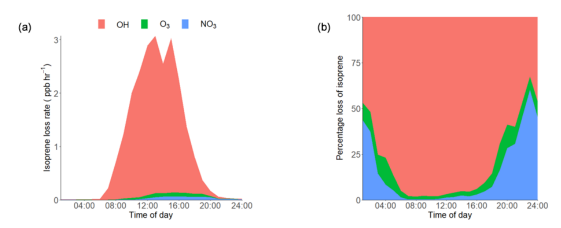


Figure 4. (a) Diurnal loss rate of isoprene calculated using the measured average diurnal profiles of isoprene, OH, NO₃ and O₃. (b) Average diurnal percentage loss of isoprene from reactions with OH, O₃ and NO₃ radicals. The IUPAC rate constants used for the calculations are as follows: NO₃, 7×10^{-13} molecule⁻¹ s⁻¹; O₃, 1.27×10^{17} molecule⁻¹ s⁻¹; OH, 1×10^{10} cm³ molecule⁻¹ s⁻¹ (Atkinson et al., 2006).

this. Without these additional standard addition calibrations, the iSOA concentrations would have been largely underestimated. The dinitrate and trinitrate NOS species eluted after the sulfate peak (RT>1.6 min). In the absence of authentic standards for these species, camphorsulfonic acid was used as a proxy for calibration. This work highlights an additional difficulty of calibration when using ESI-MS to study OSs and indicates that future studies using reversed-phase LC (RPLC) should consider the impacts of matrix effects.

3.5 Organosulfates

3.5.1 2-Methyltetrol OS (2-MT-OS)

The 2-MT-OS (C₅H₁₂SO₇) formed from the uptake of IEPOX into the particle phase is often used as a marker of low-NO isoprene photochemistry (Wennberg et al., 2018).

The time series of 2-MT-OS is shown in Fig. 6a. The particle concentration ranged from $0.7 \,\mathrm{ng}\,\mathrm{m}^{-3}$ to a maximum of $111 \,\mathrm{ng}\,\mathrm{m}^{-3}$ with a mean concentration of $11.8 \,\mathrm{ng}\,\mathrm{m}^{-3}$. The mean concentrations of 2-MT-OS and 2-MG-OS are compared to observations from previous studies in Table 3. The mean concentration observed in Beijing was much lower than those observed in the Amazon (Riva et al., 2019) and the Southeastern US (Budisulistiorini et al., 2015; Hettiyadura et al., 2019), but it was higher than summertime observations at polluted regional sites in China (Wang et al., 2018; He et al., 2018). The lower amounts of IEPOX-derived SOA result in an average AMS $f_{\mathrm{C_5H_6O}}$ in Beijing during the APHH project of only $0.2 \,\%$, which is similar to observations in other urban studies (Hu et al., 2015).

Hourly samples were collected on selected high-pollution days and used to obtain information on the diurnal evolution of the iSOA tracers. The findings on these days are consis-

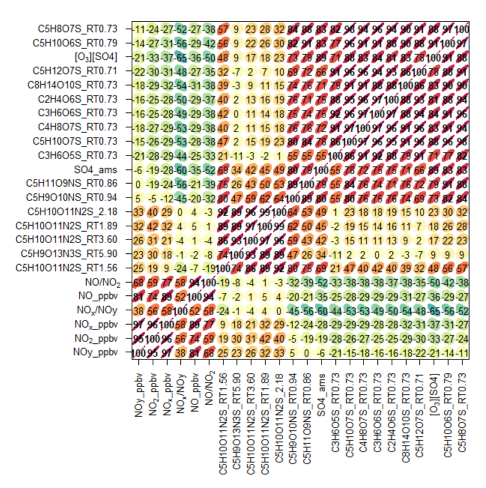


Figure 5. Correlation plot (R, openair, corPlot) highlighting the correlations between known iSOA tracers and anthropogenic pollutants. The numbers represent the R correlations between the two species; the redder, more elongated circles highlight a higher correlation.

Table 3. Isoprene CHO tracer concentrations measured via GC-MS using 24 h samples between 22 May and 22 June 2017. 2-MTs is equal to the sum of 2-methylthreitol and 2-methylerythritol, and the C5-alkene triols are equal to the sum of *cis-*2-methyl-1,3,4-trihydroxy-1-butene, 3-methyl-2,3,4-trihydroxy-1-butene and *trans-*2-methyl-1,3,4-trihydroxy-1-butene.

Isoprene tracer	Min (ng m ⁻³)	$\max_{(\text{ng m}^{-3})}$	Average (ng m ⁻³)
2-MTs	4.55	52.67	17.29
MG	1.38	15.53	7.24
C ₅ -alkene triols	0.23	1.08	0.51

tent with the 3-hourly data. The particulate 2-MT-OS measured by UHPLC-MS, on 11–12 June 2017, had a strong diurnal profile (Fig. 3b), peaking in the late afternoon (between 15:30 and 18:30 LT) and displaying a minimum overnight. This is consistent with the average diurnal profile of the gasphase precursors IEPOX+ISOPOOH (C₅H₁₂O₃) measured using the I-CIMS (Fig. S5). High levels of O₃ were observed

in the afternoon (up to 180 ppb), leading to relatively low levels of NO observed for a highly polluted environment (in some cases below 500 ppt). Thus, although the mixing ratio of NO_x was high, less than 2 % was in the form of NO on most afternoons. High levels of peroxy radicals were observed, with mean afternoon concentrations of HO2 and RO2 of around 3×10^8 and 1.5×10^9 molecule cm⁻³ respectively. Zero-dimensional box modelling indicates that up to 35 % of the isoprene-derived RO2 radicals can react with HO2 in the afternoon on some days (Newland et al., 2020). Thus, the diurnal profile in Fig. 3b, measured in samples during the measurement period, suggests that IEPOX was formed at this urban location by the reaction of OH with local isoprene emissions, with a fraction of the RO₂ radicals formed reacting with HO₂ rather than NO, and subsequent uptake to aerosol forming 2-MT-OS. OH + isoprene hydroxynitrate also has a small yield of IEPOX (Jacobs et al., 2014). The average diurnal profile of isoprene hydroxynitrates (C₅H₉NO₄) in the gas phase measured using the I-CIMS peaks at around 11:00–12:00 LT followed by a reduction during the afternoon into the evening/night (Fig. S6). This is likely to be a result of

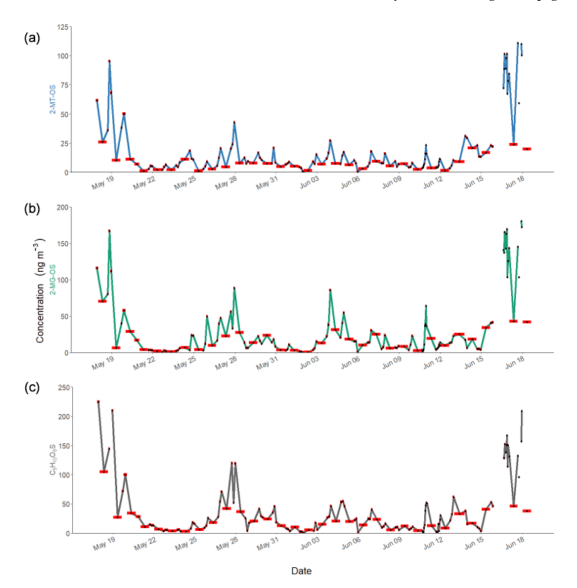


Figure 6. Time series of observed concentrations of iSOA tracers in Beijing during APHH for (a) 2-MT-OS ($C_5H_{12}O_7S$), (b) 2-MG-OS ($C_4H_8O_7S$) and (c) $C_5H_{10}SO_6$. The red bars indicate the length of the sampling period. The individual sample times will be given in the accompanying dataset (https://doi.org/10.15124/7c1a54dd-ce72-4b1a-bed1-812cf25bc407; Hamilton and Bryant, 2019).

Table 4. Comparison of iSOA tracer concentrations and ratios from previous studies in the Amazon, the Southeastern US and China.

	Mean concentration (ng m ⁻³)			Ratio low to high NO		Ratio CHO: CHOS			
Location	2-MT	2-MT-OS	2-MG	2-MG-OS	2-MT: 2-MG	2-MT-OS: 2-MG-OS	2-MT: 2-MT-OS	2-MG: 2-MG-OS	Reference
Amazon, Manuas (2016)	137*	390*			1		0.35		Cui et al. (2018)
Amazon, T3 (2014)		83 (wet)/ 399 (dry)		0.7 (wet)/ 30 (dry)		118 (wet)/ 13 (dry)			Glasius et al. (2018)
Southeastern US, Centreville (2013)		217		10.7		20.3			Riva et al. (2019)
Southeastern US, Look Rock (2013)	163.1	169.5	7.5	10	21.7	17.0	0.96	0.75	Budlisulistiorini et al. (2015)
Southeastern US, Look Rock (2013)	861*	2334*					0.37		Cui et al. (2018)
Southeastern US, Atlanta (2015)		1792		53		33.8			Hettiyadura et al. (2019)
China regional, PRD (2008)	91.5	2.2	7.7	1.4	11.9	1.57	41.6	5.51	He et al. (2018)
China regional, Beijing (2016)		5.3		3.6		1.47			Wang et al. (2018)
China rural, North China Plain (2013)	44		19.3		2.30				Li et al. (2018)
China urban, Beijing (2017)	17.3	11.8	7.2	21.5	2.40	0.55	1.47	0.33	This work

^{*} Selected sample not an average concentration.

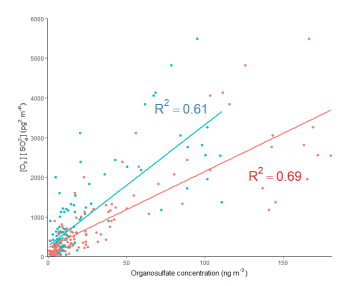


Figure 7. Plot of 2-MT-OS ($C_5H_{12}O_7S$, blue) and 2-MG-OS ($C_4H_8O_7S$, red) concentrations versus $[O_3][SO_4]$. The high time resolution data (O_3 and AMS SO_4^{2-}) have been averaged to the filter sampling time. The line was calculated with the "stat_smooth" function in the "ggplot2" R package, using the "lm" method.

the relatively low levels of NO during the afternoon, which will reduce isoprene nitrate formation from $RO_2 + NO$ reactions; thus, isoprene hydroxynitrates are unlikely to be a significant source of 2-MT-OS in Beijing.

The 2-MT-OS showed a moderate correlation with particulate sulfate ($R^2 = 0.44$) and a weak anti-correlation with photochemical age, which was estimated using the NO_x/NO_y ratio ($R^2 = 0.23$), as shown in Fig. 5. All correlations between species are shown in Fig. 5. By taking the product of the concentration of O₃, as a proxy of photochemistry, with the amount of particulate sulfate measured using AMS, [O₃][pSO₄], a much stronger correlation with 2-MT-OS was observed ($R^2 = 0.61$), as shown in Fig. 7. This observation highlights the role of both local photochemistry and particulate sulfate mass in the formation of 2-MT-OS (Fig. 7). The correlation of $[O_3][pSO_4]$ with 2-MT-OS is likely to be weaker at longer photochemical ages when the O₃ concentration is not directly related to the photochemical formation of the OS. Again, this highlights the strong role of local photochemistry in the production of low-NO iSOA (2-MT-OS) in Beijing. Elevated levels of 2-MT-OS were observed at the start and end of the measurement period which were influenced by strong southwesterly winds. There were also elevated isoprene concentrations (up to 2.9 ppb) and high particulate SO_4^{2-} levels. Therefore, these spikes in 2-MT-OS could be a result of either higher 2-MT-OS in regional aerosol transported to the site or a high isoprene emission source to the southwest of the site (i.e. producing IEPOX locally) that then reacts with increased regional sulfate pollution. The I-CIMS data show that the IEPOX/ISOPOOH (Fig. S5 and Newland et al., 2020) signal increases during the afternoon as the NO levels drop to below 1 ppb. The low NO levels indicate that up to 30 % of the isoprene peroxy radical from OH oxidation can react with HO₂ rather than NO at this site, meaning IEPOX can be formed locally (Newland et al., 2020). There is also likely to be a regional source of IEPOX and 2-MT-OS, suggesting both local and regional anthropogenic influences.

Analysis of the 2-MT-OS isomer distribution using HILIC/ESI-HR-OTOFMS, on a subset of 15 samples, indicates that β -IEPOX is the dominant ambient IEPOX isomer, which is in line with other recent observations (Cui et al, 2018; Krechmer et al., 2016) see Fig. S7). The MT-OS derived exclusively from δ -IEPOX-OS isomers could not be observed in any of the samples. The four IEPOX-OS isomers in Fig. S7 showed similar temporal trends, although small changes in the relative proportions were observed. The sum of peak areas from the 2-MT-OS isomers measured by HILIC and the quantified 2-MT-OS (sum of isomers) measured via UHPLC/ESI-HR-MS were compared and showed a high degree of correlation (R^2 of 0.84), even though the two methods used different solvents. The agreement indicates that the UHPLC/ESI-HR-MS method captures the sum of the isomers and that there is no evidence of ion source-induced artefacts.

3.5.2 2-Methyl glyceric acid OS (2-MG-OS)

The most commonly targeted SOA tracer for high-NO isoprene chemistry is 2-methylglyceric acid (2-MG) and its derivatives. As such, this tracer is the result of a direct biogenic-anthropogenic interaction. Two observed SOA tracers related to this chemistry are the OS derivatives of 2-methylglyceric acid (2-MG-OS) and the unresolved C₈ dimers of 2-MG-OS (C₈H₁₄SO₁₀) that have been identified previously in chamber-derived iSOA (Surratt et al., 2006, 2010). 2-MG-OS had an average concentration during the campaign of 21.5 ng m^{-3} , ranging from $0.3 \text{ to } 180.5 \text{ ng m}^{-3}$, with the time series shown in Fig. 6b. These values are within the range of 2-MG-OS measured in other urban locations (Q. T. Nguyen et al., 2014; Rattanavaraha et al., 2016; Hettiyadura et al., 2019). However, these concentrations are considerably higher than those previously observed at two Chinese regional background sites (Wang et al., 2018; He et al., 2018). At these locations, the ratio of the low-NO to high-NO isoprene OS tracer average concentrations was close to 1.5 (2-MT-OS:2-MG-OS; Beijing was 1.47 and Wangingsha was 1.57). However, in central Beijing, this ratio was considerably lower (2-MT-OS:2-MG-OS was 0.55), reflecting the higher proportion of RO₂ radicals reacting with NO at this location compared with the regional measurements. The ratio of 2-MT-OS:2-MG-OS observed in Beijing is compared to previous studies in Table 3 and is considerably lower than measurements taken in a range of isoprene-dominated environments (Southeastern US showed a 2-MT-OS:2-MG-OS ratio of 17, Budisolistiorini et al., 2015; the Amazon

showed a 2-MT-OS:2-MG-OS ratio of 13–118, Glasius et al., 2018; Atlanta showed a 2-MT-OS:2-MG-OS ratio of 33, Hettiyadura et al. 2019) reflecting the strong impact of urban NO emissions on iSOA formation. Future work will investigate how to use these ratios to quantify the effect of NO emissions on iSOA formation in different regions.

The mean concentration of the 2-MG-OS dimer $(C_8H_{14}SO_{10})$ was $0.57\,\mathrm{ng\,m^{-3}}$. A strong linear relationship was observed between the 2-MG-OS monomer and dimer concentrations ($R^2=0.83$) with a dimer to monomer ratio of 0.02. The formation of oligomers from reactions of 2-MG and HMML has been shown to be reduced in chamber experiments under humid conditions (Schwantes et al., 2019; Nestorowicz et al., 2018). The average RH during the afternoon of the campaign was $\sim 40\,\%$, which may account for the relatively low formation of the dimer OS compared with the monomer (see Fig. S8).

The diurnal profile of the 2-MG-OS, as shown in Fig. 8, was similar to the 2-MT-OS, peaking during the early afternoon samples but with an enhanced signal at night. There was also a strong correlation between these two species ($R^2 = 0.92$) during the campaign. The 2-MG-OS showed a stronger correlation with particulate sulfate ($R^2 = 0.52$) than 2-MT-OS ($R^2 = 0.44$), and there was also a weak anti-correlation with photochemical age ($R^2 = 0.28$). A strong correlation was also observed for 2-MG-OS with $[O_3][pSO_4]$ ($R^2 = 0.69$), as shown in Fig. 7, highlighting that formation is dependent on both photochemistry and the sulfate aerosol availability.

3.5.3 Other isoprene-related OSs and NOSs

A total of 24 additional OSs species, with molecular formulae consistent with iSOA tracers seen in chamber experiments, were also observed in Beijing (as shown in Table 2). For C₅ compounds, the most abundant species were C₅H₁₀SO₆ and C₅H₁₀SO₅, with mean concentrations of 28.7 and $26.5 \,\mathrm{ng}\,\mathrm{m}^{-3}$ respectively. The identity of the OS at m/z 182 (C₅H₁₀SO₅) is currently unknown, and the product ion MS provides little additional information other than sulfate-related fragment ions at m/z 97 and m/z 80. The OS at m/z 198 (C₅H₁₀SO₆) was identified as an IEPOX-related OS in chamber experiments by Nestorowicz et al. (2018), although at relatively low concentrations compared with the 2-MT-OS (1 %-4 %). This is very different to the observed ratio in Beijing, where the C₅H₁₀SO₆ average concentration was more than double that of 2-MT-OS, as shown in Fig. 6c. This compound showed a strong correlation with 2-MT-OS $(R^2 = 0.77)$, but it is currently unclear why this compound is the most abundant C₅ species. The molecular weight of this species is 18 Da (-H₂O) lower than 2-MT-OS, which may indicate that it is a dehydration product enhanced under acidic aerosol conditions. In addition, this species may also be enhanced if it is formed from additional VOC precursors.

Potential low-NO iSOA tracers, seen in chamber experiments, correlated strongly with the 2-MT-OS, including unresolved isomers of cyclic hemiacetals [$C_5H_{10}SO_7$] ($R^2=0.92$) and lactones [$C_5H_8SO_7$] ($R^2=0.83$) (Spolnik et al., 2018). These compounds were similar in concentration to the 2-MT-OS, with lactones at MW 212 having a mean concentration of 14 ng m⁻³ and cyclic hemiacetals at MW 214 having a mean concentration of 10.6 ng m⁻³. These compounds were also observed to be the dominant type of isoprenederived OSs in Atlanta, Georgia, although concentrations were \sim 15 times lower than the observed 2-MT-OS (Hettiyadura et al., 2019).

Additional small OS compounds, previously identified during high-NO chamber experiments, were also observed in Beijing, including (in order of decreasing concentration) glycolic acid sulfate (C₂H₄SO₆, mean of 38.4 ng m⁻³), hydroxyacetone sulfate ($C_3H_6SO_5$, mean of 20.5 ng m⁻³) and lactic acid sulfate ($C_3H_6SO_6$, mean of 14.5 ng m⁻³) (Surratt et al., 2007, 2008). These concentrations are in line with measurements made in other urban locations (Rattanavaraha et al., 2016; Huang et al., 2018; Hettiyadura et al., 2018). While all three C₂-C₃-OS compounds had strong correlations with the other iSOA OS tracers ($R^2 = 0.6-0.94$), the relative strength of isoprene versus other VOC precursors, such as aromatics, cannot be determined. As such, they cannot be definitively assigned as iSOA tracers and are, therefore, included in the potential iSOA portion of Fig. 9. The sum of the C₂ and C₃ OSs had an average concentration of 73 ng m⁻³ with a range from 2.0 to 831 ng m $^{-3}$.

In addition, nine NOS species related to isoprene were identified, as shown in Table 2 (Ng et al., 2008; Rollins et al., 2009). Some of the NOSs observed peaked in the day-time and some were enhanced at night. In total, they had a mean concentration of 24 ng m^{-3} during the campaign. NOS species are formed via the heterogenous uptake of isoprene nitrates (IN) into the particle phase. Nitrate radicals play a key role in the formation of IN, with nitrate radicals forming from reaction of NO₂ with O₃, which are both key anthropogenic pollutants. Therefore, emissions of NO_x and the formation of particulate sulfate will enhance the production of isoprene NOS species.

3.6 Contribution of isoprene SOA in Beijing

In order to estimate the total amount of isoprene-derived OSs and NOSs, labelled here as iSOA, 13 species (including isomers) were chosen that could be confidently identified as being predominately from isoprene (2-MT-OS, 2-MG-OS, $C_5H_{10}SO_7$, $C_5H_8SO_7$, $C_5H_{11}NSO_9$, $C_5H_9NSO_{10}$, $C_5H_9N_2SO_{11}$ and $C_5H_8N_3SO_{13}$). Although there were a number of other compounds with formulas similar to iSOA tracers, their trends compared to previous studies and the potential for alternative sources made a confident assignment of a VOC precursor difficult. Therefore, the estimated contribution of iSOA to the observed total particulate mass deter-

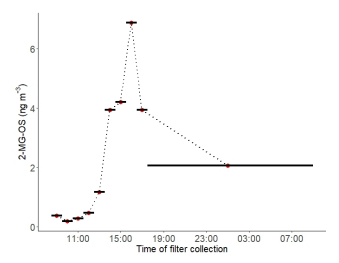
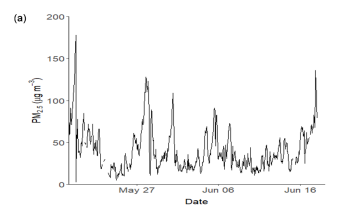


Figure 8. Diurnal profile of 2-methylglyceric acid sulfate (2-MG-OS) in particulate matter collected on filters hourly from 11 to 12 June 2017. The black lines indicate length of the sampling periods.

mined here should be taken as a lower limit. Figure 9 shows the time series of the iSOA observed in Beijing. The average concentration was 82.5 ng m^{-3} during the campaign, ranging from 718 ng m^{-3} on 19 May 2017 (11:38–14:30 LT) to 1.9 ng m^{-3} on 2 June 2017 (14:36–17:28 LT). The contribution of iSOA to the OOA factors measured by the AMS was obtained by assuming that all OS and NOS species fragment in the ion source to lose the sulfate and nitrate groups. Across the whole measurement period, the iSOA tracers represented only a small fraction of the total OOA measured by AMS (0.62% of \sum [OOA1-3]). However, towards the end of the measurement period, this increased up to a maximum of 3% on 17 June 2017 (13:32–14:23 LT).

Additional iSOA tracers containing only CHO (Table 3), including 2-methyltetrols, 2-methylglyceric acid and C₅alkene triols, were measured in separate 24h filter samples via the commonly used derivatization GC-MS method (Claeys et al., 2004; Wang et al., 2005). The average ratio of the 2-methyltetrols to their corresponding OS (2-MT:2-MT-OS) was 1.4, indicating extensive heterogeneous conversion of isoprene oxidation products within the particles. The observed ratio is slightly higher than those measured in the Southeastern US (~ 0.37 –0.96 as shown in Table 3) but much lower than that measured in the Pearl River Delta region (\sim 40) where the 2-methyltetrols dominated. In contrast, the average ratio of the high-NO iSOA tracer (2-MG) and its corresponding organosulfate (2-MG:2-MG-OS) observed in Beijing was 0.33, indicating more extensive transformation to products from heterogeneous reactions. This ratio may also reflect the more volatile nature of 2-MG compared with 2-MT. Overall, the combined concentrations of these isoprene CHO compounds were generally low (mean of 25 ng m^{-3} and max of 69 ng m^{-3}) in comparison with the



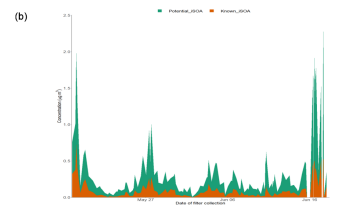


Figure 9. (a) Time series of PM $_{2.5}$ over the sampling period. (b) Time series of the total known isoprene SOA signal, 2-MT-OS, 2-MG-OS, C $_5$ H $_1$ 0SO $_7$ (MW 214), C $_5$ H $_8$ SO $_7$ (MW 212), C $_5$ H $_1$ 1NSO $_9$ (MW 261), C $_5$ H $_9$ NSO $_{10}$ (MW 275), C $_5$ H $_{10}$ O $_{11}$ N $_2$ S (MW 306) and C $_5$ H $_9$ O $_{13}$ N $_3$ S (MW 351); and the total signal from the other iSOA tracers quantified in this study.

heterogeneous iSOA compounds (i.e. isoprene-derived OSs and NOSs) targeted in this work. In addition, the concentrations of these CHO species may be overestimated based on recent studies demonstrating that thermal decomposition leads to these products being detected by GC-MS and FIGAERO-CIMS methods (D'Ambro et al., 2019); thus, the conversion to products from heterogeneous reactions (i.e. OSs and NOSs) may in fact be larger (2MT:2MT-OS of 0.5–0.91 using the overestimates of 160 %–288 % observed in Cui et al., 2018).

The study presented here shows for the first time that OS species derived from isoprene oxidation can make a significant contribution to oxidized organic aerosol in Beijing in summer. There is significant anthropogenic control, from both NO_x and sulfate aerosols, on the products and concentrations of iSOA in Beijing. The majority of the OS species showed a strong correlation towards the product of $[O_3][pSO_4]$, highlighting the role of both photochemistry and the availability of particulate sulfate in heterogeneous reactions. When the observed concentrations of all of

the OS and NOS species measured in this study (including the additional 19 compounds not confidently assigned to iSOA) are combined, they contribute 2.2% to the total OOA (\sum [OOA1-3]) on average, increasing to a maximum of 10.5%, indicating the extensive heterogeneous conversion of VOC oxidation products in Beijing in summer.

Data availability. Concentration data are available at https://doi.org/10.15124/7c1a54dd-ce72-4b1a-bed1-812cf25bc407 (Hamilton and Bryant, 2019). Filter collection times are shown in Table S3.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/acp-20-7531-2020-supplement.

Author contributions. DJB analysed the aerosol samples and quantified iSOA tracers. WJD and KLP developed the UHPLC-MS method. JRH, RED and MS provided the VOC measurements. FAS and JDL collected the NO, NO₂ and O₃ data. TJB, AM, SDW, AB, CJP and HC collected and analysed the CIMS data. LKW, DEH and EJS provided the OH and HO₂ data. BO provided the NO₃ measurements. TC, JDS and WJD carried out the offline HILIC analysis. DL, ZS and RH provided the GC-MS iSOA data. YS and WX carried out the AMS measurements and the PMF analysis. ACL and RH lead the APHH projects. DJB, ARR and JFH wrote the paper with input from all co-authors.

Competing interests. The authors declare that they have no conflict of interest.

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