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Accessibility
Airborne measurements of organosulfates over the continental U.S.

Jin Liao1,2, Karl D. Froyd1,2, Daniel M. Murphy1, Frank N. Keutsch3,4, Ge Yu5, Paul O. Wennberg5,6, Jason M. St. Clair5, John D. Crounse5, Armin Wisthaler7,8, Tomas Mikoviny7,8, Jose L. Jimenez2,9, Pedro Campuzano-Jost2,10, Douglas A. Day5,8, Weiwei Hu5,8, Thomas B. Ryerson1, Ilana B. Pollack1,2, Jeff Peischl1,2, Bruce E. Anderson10, Luke D. Ziembka10, Donald R. Blake11, Simone Meinardi11, and Glenn Diskin10

1Chemical Sciences Division, Earth System Research Laboratory, NOAA, Boulder, Colorado, USA, 2Cooperative Institute for Research in Environmental Sciences, University of Colorado Boulder, Boulder, Colorado, USA, 3Department of Chemistry, University of Wisconsin-Madison, Madison, Wisconsin, USA, 4Now at Department of Chemistry and Chemical Biology, Harvard University, Cambridge, Massachusetts, USA, 5Division of Geology & Planetary Sciences, Pasadena, California, USA, 6Division of Engineering and Applied Science, Pasadena, California, USA, 7Institut für Ionenphysik und Angewandte Physik, Leopold-Franzens Universität Innsbruck, Innsbruck, Austria, 8Now at Department of Chemistry, University of Oslo, Oslo, Norway, 9Department of Chemistry and Biochemistry, University of Colorado Boulder, Boulder, Colorado, USA, 10NASA Langley Research Center, Hampton, Virginia, USA, 11Department of Chemistry, University of California, Irvine, California, USA

Abstract Organosulfates are important secondary organic aerosol (SOA) components and good tracers for aerosol heterogeneous reactions. However, the knowledge of their spatial distribution, formation conditions, and environmental impact is limited. In this study, we report two organosulfates, an isoprene-derived isoprene epoxydiols (IEPOX) (2,3-epoxy-2-methyl-1,4-butanediol) sulfate and a glycolic acid (GA) sulfate, measured using the NOAA Particle Analysis Laser Mass Spectrometer (PALMS) on board the NASA DC8 aircraft over the continental U.S. during the Deep Convective Clouds and Chemistry Experiment (DC3) and the Studies of Emissions and Atmospheric Composition, Clouds, and Climate Coupling by Regional Surveys (SEAC4RS). During these campaigns, IEPOX sulfate was estimated to account for 1.4% of submicron aerosol mass (or 2.2% of organic aerosol mass) on average near the ground in the southeast U.S., with lower concentrations in the western U.S. (0.2–0.4%) and at high altitudes (< 0.2%). Compared to IEPOX sulfate, GA sulfate was more uniformly distributed, accounting for about 0.5% aerosol mass on average, and may be more abundant globally. A number of other organosulfates were detected; none were as abundant as these two. Ambient measurements confirmed that IEPOX sulfate is formed from isoprene oxidation and is a tracer for isoprene SOA formation. The organic precursors of GA sulfate may include glycolic acid and likely have both biogenic and anthropogenic sources. Higher aerosol acidity as measured by PALMS and relative humidity tend to promote IEPOX sulfate formation, and aerosol acidity largely drives in situ GA sulfate formation at high altitudes. This study suggests that the formation of aerosol organosulfates depends not only on the appropriate organic precursors but also on emissions of anthropogenic sulfur dioxide (SO2), which contributes to aerosol acidity.

1. Introduction

Atmospheric aerosols affect climate forcing by directly absorbing and scattering sunlight and acting as cloud condensation nuclei (CCN) to initiate cloud formation [Charlson et al., 1992; Scott et al., 2014]. Aerosols are also atmospheric pollutants because they are harmful to human health, especially the respiratory and cardiovascular systems [Pope et al., 2002], and they decrease atmosphere visibility. Secondary organic aerosol (SOA) accounts for a significant fraction of organic aerosol (OA) mass [Murphy et al., 2006; Zhang et al., 2007a]. Due to our limited knowledge of SOA formation, the modeled SOA mass often has discrepancies with observations [Heald et al., 2005; Volkamer et al., 2006]. The formation mechanism, properties, and fates of SOA must be characterized to evaluate their environmental impact [Hallquist et al., 2009].

Organosulfates are important secondary aerosol components. They are formed by reactions between organic material and sulfate, which are the main chemical components of atmospheric aerosols [Murphy et al., 2006; Zhang et al., 2007a]. Organosulfates have been detected in ambient aerosols [Iinuma et al., 2007; Surratt et al., 2007, 2008; Gomez-Gonzalez et al., 2008; Froyd et al., 2010; Chan et al., 2010; Hatch et al., 2011;...
Olson et al., 2011; Zhang et al., 2012; Worton et al., 2013; Shalamzari et al., 2013] and cloud water [Pratt et al., 2013] and are estimated to comprise up to 5–10% of OA mass over the continental U.S. [Tolocka and Turpin, 2012]. Organosulfates are thought to be good tracers for heterogeneous aerosol phase chemistry and SOA formation since the known formation mechanisms involve reactive uptake of gas phase organic species onto aerosol [Surratt et al., 2010; McNeill et al., 2012; Zhang et al., 2012]. Organosulfates are polar, hydrophilic, and low-volatility SOA compounds, which may help nanoparticle growth [Smith et al., 2008; Yli-Juuti et al., 2013] and increase their potential to become CCN. Therefore, investigation of organosulfate abundance, distributions, sources, formation mechanisms, and fates is an important step to improve our knowledge of SOA.

One of the most well studied and abundant aerosol organosulfates is isoprene epoxydiols (IEPOX) (2,3-epoxy-2-methyl-1,4-butanediol) sulfate (C5H11SO7/C0) (chemical structure shown in Figure 1a). IEPOX sulfate is one of the most abundant individual organic molecules in aerosols (e.g., 1–2% of carbon mass in the southeast U.S.) [Chan et al., 2010; Lin et al., 2013]. IEPOX sulfate was discovered to be a key intermediate in SOA formation from isoprene, the largest nonmethane carbon source [Surratt et al., 2010; Paulot et al., 2009b]. Uptake of IEPOX by acid-catalyzed ring opening of epoxydiol, followed by addition of inorganic sulfate, is known to form IEPOX sulfate [Darer et al., 2011; Eddingsasas et al., 2010; Surratt et al., 2010; Paulot et al., 2009b]. Organosulfates have also been proposed to form by reactive uptake of unsaturated compounds into the particle phase and reaction with the sulfate radical [Rudzinski et al., 2009; Nozière et al., 2010; Schindelka et al., 2013]. The vertical profiles of IEPOX sulfate were measured by the Particle Analysis by Laser Mass Spectrometry (PALMS) instrument during previous airborne campaigns [Froyd et al., 2010]. In those studies IEPOX sulfate accounted for about 2–3% of aerosol mass in the southeast U.S. and higher fractions in the tropical-free troposphere. IEPOX sulfate temporal profiles measured by aerosol time-of-flight mass spectrometry (ATOFMS) in Atlanta [Hatch et al., 2011] have also been reported. The IEPOX sulfate mass loading over the southeast U.S. has been estimated to be 10–60 ng/m3 by a model study [Pye et al., 2013].

A few other organosulfates have been quantified. For example, an organosulfate derived from 2-methyl-3-buten-2-ol, an important biogenic volatile organic emitted from pine trees, was measured to account for 0.25% of OA mass in the Manitou Forest Observatory in Colorado [Zhang et al., 2012]. The aromatic organosulfate benzyl sulfate, thought to form from anthropogenic emissions, was measured in Lahore, Pakistan, and found to account for a very small mass fraction of OA (2 ppm) but might be a useful tracer [Kundu et al., 2013].

![Figure 1.](image-url)
Glycolic acid (GA) sulfate (C$_2$H$_3$SO$_6$$^-$) (chemical structure shown in Figure 1a) is another potentially important organosulfate. Like IEPOX sulfate, the proposed formation mechanism of GA sulfate is the reaction of a gas phase organic precursor with acidic aerosol sulfate, although the formation mechanism remains to be proven. GA sulfate has been detected in ambient aerosols [Olson et al., 2011; Suratt et al., 2008] and in SOA generated by isoprene oxidation in chamber studies [Suratt et al., 2008]. GA sulfate can also form from the particle phase reaction of methyl vinyl ketone, a first generation oxidation product of isoprene, with the sulfate radical through a pathway similar to that proposed for hydroxyacetone sulfate formation [Schindelka et al., 2013], although it is unclear how important this pathway is under ambient conditions. GA sulfate formed from isoprene oxidation and other sources might account for significant aerosol mass. Ambient GA sulfate was measured to be 1.9–11.3 ng/m$^3$ from filters collected on the ground in California, Ohio, and Mexico [Olson et al., 2011]. C$_2$H$_3$SO$_6$$^-$ was also observed as a gas phase ion and may be the only gas-phase observation of an organosulfate [Ehn et al., 2010]. However, the spatial distribution of GA sulfate measurements is limited. The sources, formation mechanisms, and atmospheric importance of these species are unclear.

This study reports measurements of IEPOX sulfate C$_5$H$_{11}$SO$_7$$^-$ and the less studied GA sulfate C$_2$H$_3$SO$_6$$^-$. Measurements were made with the NOAA PALMS instrument on board the NASA DC-8 airplane over the continental U.S. during the Deep Convective Clouds and Chemistry Experiment (DC3) in May and June 2012 and the Studies of Emissions and Atmospheric Composition, Clouds, and Climate Coupling by Regional Surveys (SEAC4RS) in August and September 2013. The potential sources and formation conditions of the two organosulfates in the atmosphere are also discussed.

2. Methods

The NOAA PALMS instrument measures the chemical composition of individual particles using a laser evaporation and ionization technique [Murphy and Thomson, 1995]. Particles with a diameter larger than about 200 nm can be sized by two continuous laser beams (405 nm) and ionized by a pulsed excimer laser (193 nm). The ions are extracted and detected by a time of flight mass spectrometer [Murphy, 2007]. The PALMS instrument detects m/z > 500 with unit mass resolution. However, fragmentation can limit the detection of large molecules. The PALMS instrument is able to detect inorganic (e.g., potassium, sulfate, and metals) and organic compounds in single particles. Most of the organic compounds are fragmented due to excimer laser ionization. However, Froyd et al. [2010] found that IEPOX sulfate is not fully fragmented and can be detected as C$_5$H$_{11}$SO$_7^-$ by PALMS at m/z 215. PALMS measured the aerosol chemical composition on board the NASA DC8 airplane during DC3 and SEAC4RS over the continental U.S. from about 400 m to 12 km in altitude. The organosulfate aerosols were sampled by a forward facing solid diffuser inlet based on the University of Hawaii design [McNaughton et al., 2007] for 90% and 84% of the data in DC3 and SEAC4RS and a High Cross-flow Aerosol Sampler designed by Dr. Suresh Dhaniyala at Clarkson University as an experimental inlet for the remaining data. The solid diffuser inlet sampled up to 2.8 μm at unit efficiency and with a 0.5 s inlet residence time. The experimental inlet sampled particles up to at least 1 μm at unit efficiency and with a 1.4 s inlet residence time. Because almost all organosulfate signals (>95%) were detected in submicron particles, both inlets have the same unit sampling efficiency for organosulfates. The sampling tubing temperature was about 20°C. Additional measurements are reported for flight campaigns over the Alaskan Arctic (ARCPAC, Aerosol, Radiation, and Cloud Processes affecting Arctic Climate) in 2008 and over Central America and nearby oceans: Pre-AVE (Pre-Aura Validation Experiment) in 2004, CR-AVE (Costa Rica Aura Validation Experiment) in 2006, and TC4 (Tropical Composition, Cloud, and Climate Coupling) in 2007.

A peak at m/z 155 was the most intense organosulfate signal on average detected by PALMS during DC3 and SEAC4RS. The ratio of the isotopic signals at m/z 157 to m/z 155 of 0.04 is consistent with the presence of sulfur in the molecule (Figure 1a). An accurate mass analysis [Froyd et al., 2010] of m/z 155 for the DC3 and SEAC4RS data indicates that the empirical formula is C$_2$H$_3$SO$_6$$^-$. The structure of ambient C$_2$H$_3$SO$_6$$^-$ was investigated by Galloway et al. [2009] who suggested that ambient C$_2$H$_3$SO$_6$$^-$ is likely GA sulfate since the mass and elution time of ambient C$_2$H$_3$SO$_6$$^-$ is the same as that of a glycolic acid sulfate standard C$_2$H$_3$O$_2$SO$_4$$^-$. Since PALMS cannot distinguish between isomeric compounds, we adopt the structural identification by Galloway et al. [2009] for this work. No signal at m/z 155 was observed for many spectra that contain IEPOX sulfate at m/z 215 in the field nor when the larger organosulfate BEPOX.
excimer laser power (2.8 mJ or 2.8 × 10^9 W/cm^2). For the simulated aerosol the uncertainty in organosulfate inorganic constituents as long as metal cations are absent. The calibration was also extrapolated to high the calibration plot. The relative signal intensity at ionization ef-

detailed chemical compositions of the solutions are provided in Table S1 in the supporting information. The organosulfates were detected. The mass ratios of organic to inorganic constituents were about 0.1, and the (succinic acid). They were used to mimic the general composition of the atmospheric particles in which compounds besides GA sulfate in the solutions were sulfate, ammonium, and an oxidized organic (succinic acid). They were used to mimic the general composition of the atmospheric particles in which organosulfates were detected. The mass ratios of organic to inorganic constituents were about 0.1, and the detailed chemical compositions of the solutions are provided in Table S1 in the supporting information. The ionization efficiencies of PALMS do not depend strongly on the relative concentrations of the organic to inorganic constituents as long as metal cations are absent. The calibration was also extrapolated to high excimer laser power (2.8 mJ or 2.8 × 10^9 W/cm^2). For the simulated aerosol the uncertainty in organosulfate ionization efficiency due to the aerosol matrix was smaller than the variation of laser power. Figure 2 shows the calibration plot. The relative signal intensity at m/z 155 as a function of aerosol GA sulfate mass fraction is used to determine the sensitivity to GA sulfate and convert the PALMS m/z 155 signal to GA sulfate aerosol mass fraction. Due to the variation of laser power pulses and the inhomogeneous chemical composition of individual aerosols, the calibration has significant uncertainty denoted as the error bars. PALMS is about 3 times more sensitive to GA sulfate than IEPOX sulfate. Increased laser power reduces the organosulfate parent peaks because of increased fragmentation. The effect is less pronounced for GA sulfate than for IEPOX sulfate. This may be due to less fragmentation of the smaller GA sulfate compound.

Figure 2. Average PALMS relative signals at m/z 155 (dots) for different GA sulfate mass fractions at low and medium PALMS laser power. The fits at low power (black line) and medium power (red line) and the extrapolated fit at high power (blue line) are used to convert PALMS relative signals at m/z 155 in ambient air to GA sulfate mass fractions.

at m/z 155, which is the signal intensity at m/z 155 normalized by the total ion intensity, was used to quantify the compounds instead of the individual absolute signal intensity. Aerosols generated from 13 solutions containing different known mass fractions of GA sulfate in the range from 0.3% to 2% were delivered to the PALMS and the corresponding average relative signals of m/z 155 at medium (1.7 mJ or 1.7 × 10^9 W/cm^2) and low (0.84 mJ or 8.4 × 10^8 W/cm^2) excimer laser power were recorded. The chemical compounds besides GA sulfate in the solutions were sulfate, ammonium, and an oxidized organic (sucinic acid). They were used to mimic the general composition of the atmospheric particles in which organosulfates were detected. The mass ratios of organic to inorganic constituents were about 0.1, and the detailed chemical compositions of the solutions are provided in Table S1 in the supporting information. The ionization efficiencies of PALMS do not depend strongly on the relative concentrations of the organic to inorganic constituents as long as metal cations are absent. The calibration was also extrapolated to high excimer laser power (2.8 mJ or 2.8 × 10^9 W/cm^2). For the simulated aerosol the uncertainty in organosulfate ionization efficiency due to the aerosol matrix was smaller than the variation of laser power. Figure 2 shows the calibration plot. The relative signal intensity at m/z 155 as a function of aerosol GA sulfate mass fraction is used to determine the sensitivity to GA sulfate and convert the PALMS m/z 155 signal to GA sulfate aerosol mass fraction. Due to the variation of laser power pulses and the inhomogeneous chemical composition of individual aerosols, the calibration has significant uncertainty denoted as the error bars. PALMS is about 3 times more sensitive to GA sulfate than IEPOX sulfate. Increased laser power reduces the organosulfate parent peaks because of increased fragmentation. The effect is less pronounced for GA sulfate than for IEPOX sulfate. This may be due to less fragmentation of the smaller GA sulfate compound. Because different laser powers are optimal for different aerosol species, PALMS was operated alternately at medium and low laser power during DC3 and at high and low laser power during SEAC4RS.

Other organosulfate signals observed during DC3 and SEAC4RS were much smaller. For example, an organosulfate signal at m/z 169 may be lactic acid sulfate [Olson et al., 2011]. A signal at m/z 169 was first observed by electrospray ionization mass spectrometry (ESI-MS) in isoprene oxidation studies with methylglyoxal as the proposed precursor [Surratt et al., 2007]. The signal at m/z 169 was 1.5% of the signal at m/z 155 on average by PALMS. The vertical distribution of the signals at m/z 169 was similar to that of GA sulfate. Other potential organosulfate species at m/z 139, 141, 153, 183, and 199 had signal intensities of about 9%, 7%, 0.3%, 0.1%, and 0.1% of GA sulfate signals, respectively. Except for m/z 141, the above organosulfate ions were derived mostly from isoprene oxidation chemistry [Surratt et al., 2007]. We estimated they constitute a very small aerosol mass assuming that their sensitivities are between IEPOX sulfate and GA sulfate. Such small peaks were often below the detection limit so we make no further attempt to quantify them in this study. However, these organosulfates may account for a significant mass fraction if their sensitivities are

IEPOX sulfate was quantified using the calibration described by Froyd et al. [2010]. A synthesized GA sulfate standard [Olson et al., 2011] was used to calibrate PALMS signals at m/z 155. The mass spectrum of the GA sulfate standard is shown in Figure 1b. Due to the high variability of laser vaporization and ionization, quantitative PALMS measurements are challenging. The ion signal intensity depends on the excimer laser power and the ablation position, both of which vary from particle to particle. Therefore, the average of the relative signal intensity (2,3-epoxy-1,4-butanediol, C_4H_8O_2) sulfate was sampled in the lab [Froyd et al., 2010], suggesting that C_2H_2SO_4^- is not formed from fragmentation of a larger organosulfate.
orders of magnitude lower. Offline analysis of organosulfates by liquid chromatography and ESI-MS and availability of additional organosulfate standards are required to address this important question.

Aerosol acidity or neutralization can be estimated from the PALMS signal at m/z 195, which corresponds to a sulfuric acid cluster $\text{HSO}_4^- \cdot \text{H}_2\text{SO}_4$. Acidic particles form this peak as well as $\text{HSO}_4^-$, whereas neutralized particles favor formation of only $\text{HSO}_4^-$ ions. Particles in the stratosphere and in volcanic plumes are known to be more acidic and consequently have higher $\text{HSO}_4^- \cdot \text{H}_2\text{SO}_4$ signals compared to tropospheric particles [Murphy et al., 2007; Carn et al., 2011]. The $[\text{HSO}_4^- \cdot \text{H}_2\text{SO}_4]/[\text{HSO}_4^-]$ and $[\text{HSO}_4^- \cdot \text{H}_2\text{SO}_4]/[\text{H}_2\text{SO}_4]$ signals measured in the field generally increased as particle-into-liquid sampler [NH$_4^+$] to [SO$_4^{2-}$] ratios decreased [Froyd et al., 2009, 2010]. Lab experiments also found that the $[\text{HSO}_4^- \cdot \text{H}_2\text{SO}_4]/[\text{HSO}_4^-]$ signals from a similar Aerosol Time-of-Flight Mass spectrometer (ATOFMS) increased with decreased aerosol neutralization [Yao et al., 2011]. This acidity indicator signal also varies with laser power. Due to the different laser powers used during these two campaigns, the acidity response ion $\text{HSO}_4^- \cdot \text{H}_2\text{SO}_4$ was biased high (more acidic) during DC3 and biased low (less acidic) during SEAC4RS. The aerosol pH values were also estimated from the thermodynamic model extended aerosol inorganic model (E-AIM) [Clegg et al., 1998; Wexler and Clegg, 2002; Zhang et al., 2007b] for DC3 and SEAC4RS. Hennigan et al. [2014] and Guo et al. [2014] evaluated the aerosol acidity estimated from thermodynamic models ISORROPIA and E-AIM with and without gas phase inputs. Submicron aerosol sulfate (SO$_4^{2-}$), ammonium (NH$_4^+$), and nitrate (NO$_3^-$) from a High-Resolution Aerosol Mass Spectrometer (HR-AMS), gas phase nitric acid (HNO$_3$) from the California Institute of Technology (Caltech) chemical ionization mass spectrometer (CIMS), ambient relative humidity, temperature, and pressure were used as inputs. The impact of organic compounds on pH values was not considered. The predicted gas phase HNO$_3$ levels agreed well with the measurements. The predicted pH values generally anticorrelated with the PALMS acidity signal, as expected (Figures 3c, 3d, 3g, and 3h). Since the pH predictions from the thermodynamic model probably become less accurate as sulfate approaches complete neutralization, and since gas phase NH$_3$ measurements were not available to constrain the model, the PALMS acidity signal was used as the main aerosol acidity indicator.

To investigate the potential formation mechanisms and precursors of GA sulfate, laboratory experiments were carried out to synthesize the organosulfate at m/z 155 from both glyoxal and glycolic acid. Aerosols were nebulized from a (0.4M: 0.2M: 0.2M) mixture of glyoxal, ammonium bisulfate (NH$_4$HSO$_4$), and sulfuric acid (H$_2$SO$_4$) or a (0.2M: 0.2M: 0–0.2M) mixture of glycolic acid, NH$_4$HSO$_4$, and H$_2$SO$_4$. Some solutions were exposed to UV light (254 nm) from a standard mercury pen lamp to initiate photochemistry for a range of time periods (Table 1). A differential mobility analyzer was used to select the sizes representative of ambient particles containing GA sulfate. The size-selected aerosols were sampled by PALMS. Table 1 shows the lab experiment cases, number fraction of acidic aerosols containing m/z 155, relative m/z 155 signals in acidic aerosols, and size distribution of the aerosols detected by PALMS. Figure 1c shows the mass spectrum of a particle generated from a solution of glycolic acid, NH$_4$HSO$_4$, and H$_2$SO$_4$.

Other trace gas and aerosol concentrations were measured by a suite of instruments on board the NASA DC8 airplane during DC3 and SEAC4RS. Isoprene was measured by both proton transfer reaction mass spectrometry (PTR-MS, University of Innsbruck) [Hansel et al., 1999] and by whole air sampling (WAS; University of California Irvine) followed by laboratory analysis using gas chromatography [Colman et al., 2001]. PTR-MS measurements have a higher spatial and temporal resolution compared to WAS data. Isoprene measurements by PTR-MS do, however, suffer from furan interference in biomass burning (BB) plumes [Christian et al., 2004]. In BB-impacted air masses (identified by elevated acetonitrile levels), WAS data were used for the analysis. Two gas phase isoprene oxidation products, hydroxyhydroperoxides (ISOPOOH) and dihydroxyepoxides (IEPOX), were measured by the Caltech CIMS [Paulot et al., 2009a; St. Clair et al., 2010]. NO and O$_3$ were measured by a NOAA chemiluminescence instrument [Ryerson et al., 2001]. The aerosol mass of organic material, sulfate, and ammonium was measured by a University of Colorado Aerodyne HR-AMS [DeCarlo et al., 2006] and is reported under standard temperature and pressure conditions (1 atm and 273 K). Submicron total aerosol mass was derived from the total submicron aerosol volume measured by the Ultra-High Sensitivity Aerosol Spectrometer and the laser aerosol spectrometer aerosol sizing instruments operated by the NASA LARGE group [Ziemba et al., 2013] with the assumption of an average aerosol density of 1.4 g/cm$^3$. The submicron total aerosol mass was used to multiply the PALMS organosulfate mass fraction products to get to the absolute organosulfate mass loading.
3. Results and Discussion

3.1. Vertical Distribution of IEPOX Sulfate and GA Sulfate Over the Continental U.S.

The average vertical aerosol mass fractions of IEPOX sulfate and GA sulfate in the eastern U.S. and western U.S. during DC3 and SEAC4RS are shown in Figures 3a and 3e, respectively. Flights during the DC3 campaign targeted continental convective inflow and high altitude outflow. Most of the SEAC4RS flights targeted a wide range of tropospheric environments. SEAC4RS campaign data were excluded for marine flights and for the flight on 2 September that sampled convective outflow, where the vertical profile was similar to DC3 with enhanced IEPOX sulfate at high altitudes. IEPOX sulfate is estimated to account for 1.3% and 0.2% of submicron aerosol mass on average near the ground in the eastern and western U.S., respectively, during DC3, and 1.4% and 0.2% during SEAC4RS. Near the ground is defined as from the lowest GPS altitudes sampled by DC-8 (about 400 m) to 1500 m since the typical daytime boundary layer height was about 1–2 km. The IEPOX sulfate mass fraction decreased in the upper troposphere and lower stratosphere (to < 0.2% of submicron aerosol mass). GA sulfate accounted for 0.6% and 0.3% of submicron aerosol mass near the ground and increased to 0.7% and 0.6% of submicron aerosol mass in the upper troposphere in the eastern and western U.S., respectively, during DC3, compared to 0.4% and 0.6% near the ground and 0.6% and 0.4% near the ground and 0.6% and 0.4%.
in the upper troposphere in the eastern and western U.S. during SEAC4RS. Total IEPOX sulfate mass accounted for 2.2% and 0.7% of total organic mass measured by AMS near the ground in the eastern and western U.S. and for 5.0% and 1.9% of total sulfate mass measured by AMS on average in these two campaigns. Total GA sulfate mass accounted for 0.9% and 0.7% of total organic mass measured by AMS near the ground in the eastern and western U.S. and for 2.0% and 2.5% of total sulfate mass measured by AMS on average in these two campaigns. Most of the eastern U.S. flights were over the southeast U.S. There were several low-altitude flight segments over the “isoprene volcano” region in the Ozark Mountains of Arkansas and Missouri during SEAC4RS. Although isoprene observed in the “isoprene volcano” region was significantly enhanced, there was no enhancement of IEPOX sulfate observed compared to average levels in the eastern U.S.

The IEPOX sulfate mass fraction was highest near the ground in the eastern U.S. where the isoprene emissions were most intense. Compared to IEPOX sulfate, the vertical profiles of GA sulfate mass fraction were nearly constant with altitude and peaked in the upper troposphere, suggesting that GA precursors are more widely distributed and probably longer lived than the IEPOX-derived sulfate. Accordingly, GA sulfate may account for significant aerosol mass throughout the global troposphere. The average IEPOX sulfate mass fraction of 1.4% of total submicron aerosol mass or of 2.2% of total OA mass in the southeast U.S. from 400 m to 1500 m was close to the aircraft measurements in 2004 by Froyd et al. [2010] (3% of submicron aerosol mass at low altitude) and the ground measurements by Lin et al. [2013] (1–2% of OA mass). The estimated mass loading of GA sulfate of 20 ng/m³ at low altitudes was near the upper limit of GA sulfate detected from filter measurements collected from several ground sites [Olson et al., 2011]. To our knowledge, these are the first-online and the first-airborne measurements of GA sulfate reported.

### 3.2. IEPOX Sulfate and Its Precursors

The IEPOX sulfate vertical profiles had a similar pattern to those of isoprene and its gas phase oxidation product IEPOX (Figures 3b and 3f). Average isoprene mixing ratios were similar during DC3 and SEAC4RS in the eastern U.S. The gas phase IEPOX (and ISOPOOH) was significantly lower during SEAC4RS, which was probably due to higher NO concentration observed in the
southeast U.S. during SEAC4RS than DC3. Considering lower levels of IEPOX (523 pptv in DC3 versus 285 pptv in SEAC4RS on average) during SEAC4RS, a similar IEPOX sulfate mass fraction in both campaigns, and twice higher submicron aerosol mass loading during SEAC4RS near the ground in the southeast U.S., production of IEPOX sulfate from gas phase IEPOX seemed to be much more efficient during SEAC4RS. The lower pH near the ground estimated during SEAC4RS may contribute to the more efficient IEPOX sulfate formation. Another possibility is that reactive uptake of IEPOX formed other species such as 2-methyltetrols and C₅ alkene triols [Surratt et al., 2010] in particle phase under the DC3 conditions. Further study is needed to explore the reasons for the higher IEPOX sulfate production efficiency during SEAC4RS.

The vertical profiles of IEPOX sulfate and its precursors may provide clues about the lifetime of IEPOX sulfate, which has not been investigated. Generally, isoprene, gas phase IEPOX, and aerosol phase IEPOX sulfate dropped to insignificant levels above 3.5 km, 4 km, and 8 km, respectively. The chemical lifetime of isoprene is about 1–3 h depending on hydroxyl radicals (OH) concentrations [Paulot et al., 2009a]. The lifetime of gas phase IEPOX due to OH oxidation is about 3–28 h [Jacobs et al., 2013; Bates et al., 2014], and loss to aerosol uptake varies from about 1 h to days depending on the aerosol acidity [Surratt et al., 2010; Gaston et al., 2014]. The IEPOX sulfate vertical profile suggests that the lifetime of IEPOX sulfate is longer than those of isoprene and gas phase IEPOX. Higher IEPOX sulfate mass fractions in the upper troposphere observed during DC3 compared to SEAC4RS may be due to shortened vertical transport time of either IEPOX sulfate or isoprene from convection because DC3 flights were designed to target convection, and SEAC4RS mostly sampled nonconvectively active air masses. The observed vertical profiles of isoprene, IEPOX, and IEPOX sulfate and the relevant HOₓ and NO levels may help modelers to estimate the IEPOX reactive uptake rates under ambient conditions that can be compared with rates measured in the laboratory at different conditions (e.g., aerosol acidity).

Two case studies of IEPOX sulfate and isoprene measurements are shown in Figure 4. The case study on the 11 June 2012 flight to the southeast U.S. during DC3 (Figures 4a and 4b) further confirms that IEPOX sulfate is formed from isoprene. IEPOX sulfate mass fraction generally tracks the concentrations of isoprene and gas phase IEPOX. The consistent low levels of isoprene (<200 pptv), IEPOX (<200 pptv), and IEPOX sulfate (<0.3% mass fraction) at low altitudes near 10:25 P.M. UTC are due to flying over the low vegetation and low isoprene emission area between Arkansas and Mississippi (Figure 4a). This indicates that isoprene is required in IEPOX sulfate formation and is consistent with the IEPOX sulfate formation from oxidation of isoprene as proposed by Surratt et al. [2010] and Paulot et al. [2009a]. This also indicates that IEPOX sulfate formation can be isoprene-limited in low-leaf areas in the southeast U.S. Meanwhile, the levels of IEPOX sulfate generally followed the OA mass concentrations. This may indicate that SOA mass formed from biogenic VOCs oxidation contributed substantially to OA mass in the southeast U.S. and that IEPOX sulfate can be used as a tracer for SOA formed from isoprene oxidation. Figures 4c and 4d demonstrate a case study near Northern California and southern Oregon on 6 August 2013 where there were significant isoprene and gas phase IEPOX concentrations, but no IEPOX sulfate was detected. In the enhanced isoprene periods, the aerosols were mostly neutralized, and the airplane was sampling biomass burning plumes. A similar case (not shown) with high isoprene and no detectable IEPOX sulfate was found on the 26 August 2013 flight. In general, little IEPOX sulfate was observed in biomass burning plumes because the aerosols were neutralized. This indicates that either IEPOX does not effectively react with neutralized aerosols or the reactive uptake forms other SOA products (e.g., 2-methyltetrols and C₅ alkene triols) not IEPOX sulfate [Surratt et al., 2010].

In addition to the appropriate organic and sulfate precursors, the formation of IEPOX sulfate from isoprene is modulated by sulfate aerosol acidity and gas phase NO levels. Figures 5a and 5b show IEPOX sulfate mass fraction versus isoprene mixing ratios when the aerosols were acidic (PALMS acidity signal > 0.002) (red) and near neutralized (PALMS acidity signal < 0.002) (black). Isoprene concentrations in Figure 5 are from PTR-MS with a detection limit of 6 pptv (2σ for 5 min data) and were filtered for acetonitrile less than 200 pptv to exclude biomass burning plumes. More IEPOX sulfate was formed on average at the same isoprene levels when aerosols were acidic compared to near neutralized. The same data are plotted for low (<100 pptv) and high NO (>200 pptv) levels in Figures 5c and 5d. Generally, higher IEPOX sulfate mass fractions were observed at low NO conditions for both field campaigns, which is consistent with efficient formation of gas phase IEPOX from isoprene oxidation under low NO conditions. The significant IEPOX sulfate present at high NO conditions (even > 500 pptv) during SEAC4RS may be due to transport or formation of IEPOX from isoprene hydroxynitrate oxidation [Jacobs et al., 2014], which probably plays a small role. Aerosol SO₄²⁻ concentrations are known to be important for IEPOX sulfate formation, and NH₄⁺
concentrations may also be important [Nguyen et al., 2014]. The average IEPOX sulfate mass fraction increased with aerosol $SO_4^{2-}$ or $NH_4^+$ at low levels ($SO_4^{2-}/C_0 < 2$ or $NH_4^+ < 1 \mu g/m^3$) but not at higher mass loadings. This indicates that the aerosol $SO_4^{2-}$ and $NH_4^+$ levels are important but may not be limiting factors of IEPOX sulfate formation in most ambient conditions. It is worth noting that almost all IEPOX sulfate and GA sulfate signals were observed in particles classified as sulfate-organic mixtures.

### 3.3. Aerosol Acidity

The PALMS acidity signal has undergone previous validation as a qualitative indicator of sulfate acidity [Murphy et al., 2007; Froyd et al., 2009, 2010; Carr et al., 2011; Yao et al., 2011]. Our data show that under some conditions aerosol acidity as measured by PALMS is important in both ambient IEPOX sulfate and GA sulfate formation. At low altitudes (<1000 m) in the eastern U.S. where isoprene is more abundant, the IEPOX sulfate and GA sulfate signals were well correlated with PALMS aerosol acidity signal ($HSO_4^{-}/C_0 \cdot H_2SO_4$) during DC3 and SEAC4RS when IEPOX sulfate and GA sulfate signals were above their respective detection limits (see Figure 6). This correlation indicates that higher acidity tends to promote formation of IEPOX sulfate and GA sulfate near the ground when the organic precursors are abundant. This provides field evidence for an important role of aerosol acidity in ambient IEPOX sulfate formation.

The importance of acidity agrees with both the acid-catalyzed epoxydiol ring opening formation mechanism [Surratt et al., 2010] and the sulfate radical initiated organosulfate formation because efficient formation of sulfate radicals also requires acidity [Schindelka et al., 2013]. At comparable IEPOX concentrations, higher levels of IEPOX sulfate were generally observed in particles with higher acidity (see Figures 5a and 5b), concentrations may also be important [Nguyen et al., 2014].
also demonstrating the important role of acidity in IEPOX sulfate formation. Conversely, a correlation was not observed with aerosol sulfate near the ground in the southeastern U.S. In a recent study at a rural ground site in southeastern U.S., Xu et al. [2014] reported a correlation of biogenic OA mass with aerosol sulfate but not with aerosol acidity. This apparent discrepancy is at least partially explained by the inherent differences in the measured aerosol properties. PALMS gives a direct measure of two organosulfate species that comprise 1–2% of aerosol mass, whereas Xu et al. [2014] used Aerodyne AMS bulk measurements to extract a biogenic OA factor representing 10–30% of aerosol mass. However, other factors besides aerosol acidity such as gas phase IEPOX partitioning, aerosol water content, and subsequent neutralization during particle aging may affect IEPOX sulfate concentrations and the apparent connection to aerosol acidity. Accordingly, the correlation with acidity was not generally observed above the boundary layer. The important role of relative humidity in IEPOX sulfate formation is discussed in section 3.4.

3.4. The Impact of Relative Humidity

Ambient relative humidity determines aerosol liquid water content, which affects the partitioning of gas phase compounds to aerosols, aerosol pH, and ionic strength [Zhang et al., 2007b]. IEPOX sulfate and GA sulfate had opposite trends with relative humidity (RH). Most (95%) of the data collected during these two...
aircraft campaign were during the daytime so that the RH trends were likely not due to the daytime and nighttime RH differences. The IEPOX sulfate mass fraction generally increased as RH increased up to 60%–80% (Figure 7). The same trend was observed if IEPOX sulfate mass fraction was normalized by isoprene mixing ratios or if only low-altitude data were plotted (not shown). This may indicate that aerosol liquid water promotes IEPOX partitioning and suggest that uptake can be an important step in IEPOX sulfate formation. Conversely, the GA sulfate mass fraction generally decreased with increasing RH (Figure 7). This may indicate that more concentrated or more acidic aerosols promote formation of GA sulfate at low RH levels [McNeill et al., 2012] and that partitioning of organic precursors may not limit GA sulfate formation in ambient air.

3.5. GA Sulfate in the Lower Stratosphere and Evidence for In Situ Formation

In stratospheric air (\([\text{O}_3]/[\text{CO}] > 2.3\)), the GA sulfate mass fraction decreased as \(\text{O}_3\) increased (Figure 8a). This indicates that stratospheric GA sulfate or its gas phase precursors likely come from the troposphere and were diluted or removed via physical and chemical processes as they traveled further into the stratosphere. The GA sulfate mass fraction (Figures 3 and 8a) reached a maximum (≈1%) near the tropopause because this region contains both highly acidic aerosols and more abundant organic precursors than the lower stratosphere. Particles that originate in the stratosphere or tropopause region are clearly identified by their high sulfate content and high acidity [Murphy et al., 2007]. The number fraction of such stratospheric origin aerosols decreased dramatically (Figure 8b, red) at lower altitudes. However, when examining just these stratospheric origin particles, the number fraction containing GA sulfate increased from 16% to 45% when they mixed into the troposphere (Figure 8b, black). This indicates that the maximum GA sulfate concentration near the tropopause is likely formed in situ when gas phase organic precursors from lower altitudes encounter highly acidic particles from the stratosphere.
3.6. Investigation of Potential GA Sulfate Organic Precursors

Ambient measurements provide some clues about the sources of GA sulfate. Although GA sulfate had a different spatial distribution from IEPOX sulfate, GA sulfate had a correlation ($r^2 = 0.34$ for SEAC4RS; $r^2 = 0.33$ for DC3 and SEAC4RS together) with IEPOX sulfate at low altitudes (below 1000 m) (Figure 9). Moreover, aerosols containing high IEPOX sulfate always had significant GA sulfate, but aerosols with significant GA sulfate may or may not contain IEPOX sulfate. This pattern probably indicates that the organic precursors of GA sulfate are more diverse than IEPOX sulfate, which is formed only from isoprene.

Lab experiments were performed to investigate some potential organic precursors and formation mechanisms of GA sulfate (see Table 1). Glyoxal was proposed to be a potential organic precursor of GA sulfate under UV irradiation [Galloway et al., 2009]. Aerosols generated from a mixture of glyoxal, ammonium bisulfate, and sulfuric acid were delivered to PALMS to investigate the formation of GA sulfate. No signal at $m/z$ 155 was detected when the solution containing glyoxal was newly made and not exposed to UV radiation. The relative signal at $m/z$ 155 increased to ~3 × 10$^{-4}$ when the solution was a few days old or exposed to UV radiation. This indicates that GA sulfate can form in a glyoxal/ammonium bisulfate/sulfuric acid solution with low efficiency and that UV radiation can accelerate the reaction. Aerosols generated from a solution containing glycolic acid instead of glyoxal had higher relative signals (1.5–6 × 10$^{-3}$) at $m/z$ 155 even when the solution was newly made and not exposed to UV radiation. Lowering the solution acidity (pH > 3) significantly decreased the relative signals of GA sulfate ($8 × 10^{-5}$). The laboratory experiments suggest that the organosulfate at $m/z$ 155 is formed from a glycolic acid solution more efficiently than from a glyoxal solution, and acidity largely promotes formation of GA sulfate.

Although glyoxal in aerosol can be converted to glycolate by a disproportionation reaction [Fratzke and Reilly, 1986] and thereby provides a possible route to GA sulfate formation, the disproportionation is negligibly slow in acidic solutions.
Glycolic acid, a potential important precursor of GA sulfate identified in lab experiments, has both biogenic and anthropogenic sources [Warneck, 2005]. Methyl vinyl ketone, another potential gas phase precursor of GA sulfate [Schindelka et al., 2013], has isoprene as a biogenic source. Glycolic acid can form from oxidation of glycolaldehyde (CHOCH2OH) [Perri et al., 2009; Warneck, 2005] and acetic acid (CH3COOH) [Tan et al., 2012], both of which come from biogenic and anthropogenic sources [e.g., de Gouw et al., 2009; Goldstein et al., 1996; Chebbi and Carlier, 1996; Paulot et al., 2011]. Glycolaldehyde is water soluble and can undergo reactive uptake on aerosols (Heff. = 107 M/atm) [Nguyen et al., 2013]. In the aqueous phase, the oxidation of glycolaldehyde by OH may be more efficient than that of acetic acid [Lim et al., 2005]. The reaction of glycolaldehyde in the aerosol phase forms glycolic acid on a time scale of minutes [Ortiz-Montalvo et al., 2012]. The lifetime of glycolaldehyde due to photolysis and reaction with OH is relatively long (1 day and > 2.5 days) [Racher et al., 2001]. Glycolaldehyde is the second generation product from isoprene oxidation by OH, and about 10–30% of isoprene oxidation yields glycolaldehyde at high NO conditions [Bates et al., 2014; Galloway et al., 2011; Paulot et al., 2009a] but only 3% yields glycolaldehyde at low NO conditions [Bates et al., 2014]. Glycolaldehyde can also form from oxidation of anthropogenic emissions of ethene [Spaulding et al., 2003; Warneck, 2005]. Although the organic precursors of GA sulfate in the ambient are not well known, glycolic acid and its precursors (such as glycoaldehyde, acetic acid, ethene, and isoprene) are likely precursors according to known photooxidation pathways, lab experiments, and ambient measurements. However, no clear correlation was found in the aircraft data between potential GA sulfate gas phase organic precursors (glycoaldehyde or ethene) and aerosol phase GA sulfate mass fraction. This is not completely unexpected because GA sulfate vertical profiles seem to correlate with aerosol acidity, and gas phase GA sulfate organic precursors may not be the limiting factor of GA sulfate formation in most of the troposphere. Compared to the precursor of IEPOX sulfate, the precursors for GA sulfate have more diverse emission sources and longer photochemical lifetimes. These probably contribute to a much wider spatial distribution of GA sulfate in the troposphere.

### 3.7. GA Sulfate Measurements in the Tropics and Arctic

The IEPOX sulfate measurements in the tropics and eastern U.S. were summarized by Froyd et al. [2010]. GA sulfate and PALMS acidity signal measurements in the tropics during the CR-AVE, PRE-AVE, and TC4 field studies and in the Arctic during the ARCPAC field study are shown in Figure 10. The GA sulfate vertical profiles in these campaigns also generally correlated with the PALMS acidity signal. The highest mass fraction of GA sulfate in the upper troposphere in CR-AVE was likely driven by the very acidic aerosols at those altitudes. Lower aerosol acidity during ARCPAC (3–7 km) also contributed to the lower GA sulfate mass fraction. GA sulfate was significantly lower near the ground in the Arctic compared to midlatitudes (e.g., DC3) even with similar aerosol acidity to DC3. This is consistent with the notion that GA sulfate originates from biogenic and anthropogenic sources, both of which are much weaker in the Arctic region (Alaska and the Arctic Oceans). The measurements of GA sulfate in the Arctic, midlatitudes, and tropics demonstrate that GA sulfate is widespread in locations that have acidic aerosols and significant biogenic and anthropogenic emissions or outflow.

![Figure 10. Vertical profiles of GA sulfate aerosol mass fraction and the PALMS acidity signal during previous field campaigns in the Central American tropics (CR-AVE, Pre-AVE, and TC4) (red), in the Arctic (ARCPAC) (blue), and in the midlatitudes (DC3) (black).](image-url)
4. Conclusions

IEPOX sulfate and GA sulfate were quantified in aerosols sampled over a wide variety of tropospheric and lower stratospheric environments. IEPOX sulfate was most abundant near the ground in the southeast U.S. where it was estimated to account for about 1.4% of submicron aerosol mass. The measurements confirm that IEPOX sulfate can be used as a tracer of SOA formed primarily from low-NO isoprene oxidation chemistry in regions of acidic aerosols such as southeast U.S. GA sulfate was more spatially uniform and was estimated to account for about 0.5% of submicron aerosol mass on average from near the ground to the upper troposphere and lower stratosphere. Considering its spatial distribution, GA sulfate may be even more abundant than IEPOX sulfate globally. The wide spatial distribution of GA sulfate is consistent with multiple organic precursors of both anthropogenic and biogenic origin. This study provides field evidence of the importance of acidity in formation of both organosulfates under some conditions. Aerosol acidity is measured by the PALMS acidity signal. Higher relative humidity also promotes formation of IEPOX sulfate. The different RH dependence of IEPOX sulfate and GA sulfate may indicate that the rate-limiting steps in their ambient formation are different. The in situ formation of GA sulfate in the upper troposphere and lower stratosphere provides evidence for aerosol chemical reactions at high altitudes and low RH and for the presence of gas phase oxygenated organic compounds in the upper troposphere. Other potential organosulfate species have concentrations a factor of >10 lower than IEPOX sulfate and GA sulfate, assuming similar detection sensitivities. Although the total organosulfate burden is limited to a few percent of aerosol mass over the continental U.S., organosulfates are good tracers for SOA formation. The emission of SOx, which contributes to aerosol acidity and water uptake, likely promotes the formation of SOA compounds such as IEPOX sulfate. Regions with both elevated VOCs and SO2, such as Asian pollution plumes or continental convective outflow in the upper troposphere, may produce significantly more aerosol mass via these mechanisms.

References


