Ozone-driven photochemical formation of carboxylic acid groups from alkane groups

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Abstract

Carboxylic acids are ubiquitous in atmospheric particles, and they play an important role in the physical and chemical properties of aerosol particles. During measurements in coastal California in the summer of 2009, carboxylic acid functional groups were highly associated with trajectories from an industrial region with high organic mass (OM), likely from fossil fuel combustion emissions. The concentration of carboxylic acid groups peaked during daytime, suggesting a photochemical secondary formation mechanism. This daytime increase in concentration was tightly correlated with $O_3$ mixing ratio, indicating $O_3$ was the likely driver in acid formation. Based on the diurnal cycles of carboxylic acid and alkane groups, the covariation of carboxylic acid groups with $O_3$, and the composition of the Combustion factor resulted from the factor analyses, gas-phase alkane oxidation by OH radicals to form dihyrofuran followed by further oxidation of dihydrofuran by $O_3$ is the likely acid formation mechanism. Using the multi-day average of the daytime increase of carboxylic acid group concentrations and $m/z$ 44-based Aged Combustion factor, we estimated the lower-bound contributions of secondary organic aerosol (SOA) formed in 12-h daytime of processing in a single day to be 30% of the carboxylic acid groups and 25–45% of the Combustion factor concentration. These unique ambient observations of photochemically-driven acid formation suggest that gas-phase alkanes might be important sources of SOA formation in this coastal region.

1 Introduction

Organic compounds typically account for 10–70% of dry particle mass (Turpin et al., 2000). Understanding the chemistry of particle-phase organic compounds is important for assessing the effects of aerosol particles on air quality, human health, and climate change (Fuzzi et al., 2006). The major organic components identified in ambient particles include alkane, carboxylic acid, hydroxyl, amine, and non-acid carbonyl functional...
groups (Maria et al., 2002; Liu et al., 2009; Russell et al., 2009a). Alkane groups, a large component of ambient organic compounds (Maria et al., 2002), are typically associated with primary gas and particle-phase emissions that originate from fossil fuel combustion emissions, including vehicular exhaust (Rogge et al., 1993) and coal burning (Wang et al., 2009). In addition to being prevalent in primary emissions, alkane groups remain in many oxygenated secondary organic products, e.g. β-pinene oxidation products might include norpinic acid (Yu et al., 1999), which has two oxygenated groups and five hydrogenated carbons, each with 1–3 C-H bonds. Carboxylic acid groups are observed in urban, rural, and remote atmospheric particles and sometimes account for more than 30% of OM (Grosjean et al., 1978; Kawamura and Gagosian, 1987; Kawamura, 1993; Fraser et al., 2002; Wang et al., 2002; Sullivan and Weber, 2006; Claeys et al., 2007; Russell et al., 2009a; Wang et al., 2009; Zhang et al., 2010). Because many carboxylic acids are highly soluble and tend to absorb water under high relative humidity (RH), they could affect physical (e.g. light scattering) and chemical (e.g. aqueous-phase reaction) properties of ambient particles. Carboxylic acid groups are generally SOA components. For example, enhanced abundance of ambient carboxylic acids in summer and in the afternoon suggests that carboxylic acids are formed by photochemical oxidation in the atmosphere (Kawamura and Ikushima, 1993; Kawamura and Yasui, 2005). Laboratory studies have shown condensed-phase carboxylic acid groups formed as secondary organic aerosols (SOA) from anthropogenic and biogenic precursors (Yu et al., 1999; Fisseha et al., 2004; Claeys et al., 2007). Aqueous-phase or in-cloud formation of carboxylic acids has been suggested by several field studies and modeling simulations (Blando and Turpin, 2000; Warneck, 2003; Ervens et al., 2004; Yu et al., 2005; Sullivan and Prather, 2007; Altieri et al., 2006), indicating carboxylic acid groups can be formed through various processes under different ambient conditions. Recently, alkane reactions have been studied by several research groups (Hallquist et al., 2009; Lim and Ziemann, 2009; Miracolo et al., 2010; Presto et al., 2010). Russell et al. (2011) proposed formation of carboxylic acid groups from alkane groups via photochemical oxidation.
Large uncertainty in SOA formation mechanisms makes identification of ambient SOA controversial. Organic carbon (OC) to elemental carbon (EC) ratio has been used to estimate SOA by assuming an average OC/EC from emission source measurements. OC/EC exceeding the average OC/EC is assumed to be SOA (Turpin et al., 1991). The SOA mass estimated from this method is highly uncertain, since OC/EC is highly variable from source to source (Gray et al., 1986) and the average OC/EC is dependent on meteorological conditions. Another approach is to identify SOA products from individual precursors (Yu et al., 1999). This method requires detection of SOA by molecular level speciation and known SOA formation mechanisms, which are often not available.

In this work, organic functional groups are quantified using Fourier Transform Infrared spectroscopy (FTIR) and organic mass fragments are measured using Aerosol Mass Spectrometry (AMS) at a coastal site in southern California. Because of the frequent and consistent aerosol transport from the shipping activities at Los Angeles and Long Beach, the measurements provide an opportunity to use the diurnal trend in aerosol composition to evaluate the contribution of SOA. The formation mechanism of carboxylic acid groups and other oxygenated organic aerosol (OOA) is also considered based on the recurrence of diurnal cycles.

2 Sample collection and instrumentation

Submicron particles were continuously collected at the Scripps Pier (8 m above sea level) in La Jolla (32.87°N, 117.25°W), California, from 15 August to 1 October 2009. Instruments were deployed in a temperature-controlled container at the end of the pier (300 m west of the shoreline) and shared a common 3/8” o.d. stainless steel inlet. Submicron particles were separated by a PM₁ cyclone and were collected on four filters daily for the time periods: 06:00–10:00, 10:00–14:00, 14:00–18:00, and 18:00–06:00 (the next day) Pacific Standard Time (PST, one hour earlier than local daylight time), representing morning, midday, afternoon, and nighttime samples, respectively. A 24-h
sample was simultaneously collected in parallel with shorter samples for each day. 37 mm Teflon filters (Teflo, Pall Inc., Ann Arbor, MI) were used for the FTIR analysis performed using a Bruker Tensor 27 FTIR Spectrometer with a DTGS detector (Bruker, Waltham, MA). Samples were frozen during storage to reduce desorption. Each Teflon filter was scanned before and after sampling using the FTIR and the pre-scanned spectrum was subtracted from the post-scanned spectrum to correct for variability in the polytetrafluoroethylene (Teflon) absorption of the Teflon filters. An automated algorithm was used to conduct spectrum subtraction, baselining, peak-fitting, and error estimation (Russell et al., 2009a). Mass concentrations of alkane, carboxylic acid, hydroxyl, amine, non-acid carbonyl, organonitrate, alkene, and aromatic functional groups were quantified using previously reported algorithms and standards (Russell et al., 2009a; Day et al., 2010).

Concentrations of non-refractory organics, sulfate, ammonium, nitrate, and chloride in submicron particles were measured using a quadrupole AMS (Aerodyne, Billerica, MA). In this instrument, particles passed through a 100 µm orifice are focused by an aerodynamic lens followed by vaporization (600 °C) and ionization at the entrance of a quadropole mass spectrometer. Particle size is measured by time-of-flight between a rotating chopper, which modulates the particle beam (Jayne et al., 2000). The “mass spectrum” (MS) mode and the “time-of-flight” (TOF) mode alternated during the measurements. Complete mass spectra (1-300 amu) and size distributions for selected mass fragments were stored at 5-min resolution. Transmission efficiency was approximately 100% for 60 to 600 nm particles (Jayne et al., 2000). Dry ammonium nitrate particles (350 nm) were used to calibrate the ionization efficiency weekly. Collection efficiency (CE) of the AMS was assigned to each 5-min organic and inorganic measurements to correct for particle loss due to bouncing off the vaporizer. The CE (ranging from 0.45 to 1) was determined as a linear function of ammonium to sulfate molar ratio, with 0.45 and 1 corresponding to ratios 1 and 0, respectively (Quinn et al., 2006). The campaign average ammonium to sulfate molar ratio was 1.9 ± 1.2, and the CE was 0.45 for approximately 85% of the measurements.
Single particles were impacted on Si$_3$N$_4$ windows on 27 August and 4, 14, 20, and 22 September. Four samples were collected (15–30 min) on these days for periods overlapping the four FTIR filter sampling periods. Samples were stored at temperatures below 0°C before analysis at the Advanced Light Source (Lawrence Berkeley National Laboratory, CA) on beamline 5.3.2 (Liu et al., 2009). Single particle image and K-edge X-ray absorption spectrum were acquired using a combination of Scanning Transmission X-ray Microscopy (STXM) and Near-Edge X-ray Absorption Fine Structure (NEXAFS) spectroscopy. Organic functional groups (including alkane, hydroxyl, ketone, alkene, and carboxylic acid groups) and inorganic potassium and carbonate of carbon-containing single particles were measured (Russell et al., 2002; Maria et al., 2004). Particle size, image, and organic functional group abundance were analyzed using an automated algorithm described by Takahama et al. (2010).

Carbon monoxide (CO) dry air mole fractions were measured by a newly installed Horiba APMA-370 NDIR analyzer. Air was continuously pumped through a sampling line (400 m; 1/2″ Dekabon) with an inlet 20 m a.s.l. Measurements were reported as 5-min averages, initially using the factory calibration. The instrument was subsequently calibrated against standards on the NOAA/ESRL 2004 CO calibration scale, and the field campaign results were recalculated based on the measurement of a whole air reference cylinder that was measured during the field campaign and also during the NOAA standard calibrations. In the absence of a more thorough real-time calibration, we estimate that the precision of the 5-min averaged CO measurements is 5–10 ppb.

Ozone (O$_3$) mixing ratio was monitored using a Thermo Environmental Instruments (TEI) 49C analyzer and measurements were recorded as 1-min averages. The O$_3$ measurements were not calibrated during the campaign, and the concentrations relative to the campaign average were reported.
3 Results

This section describes the meteorological conditions under which carboxylic acid groups are formed, the composition of organic mass quantified through complementary measurements by FTIR, AMS, and STXM-NEXAFS. The daily variations in carboxylic acid and alkane groups are discussed. To help to understand these variations, components contributing to organic mass are identified from factor analysis.

3.1 Meteorological conditions during the sampling period

The sampling period was characterized by stable temperature and RH with the averages and standard deviations being $20.2 \pm 2.5 ^\circ C$ and $79.3 \pm 8.6\%$, respectively. Temperature peaked in the afternoon and showed a minimum in the early morning, anti-correlating with RH. Photosynthetically active radiation peaked at noon. Land-sea breeze circulation was observed during the measurement period and was consistent with previous studies (Hughes et al., 2007). In general, surface wind shifted at about 07:00 a.m. (from offshore to onshore wind) and 10:00 p.m. (from onshore to offshore wind).

Back trajectories were calculated hourly using the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Draxler and Rolph, 2003; Rolph, 2003) at 200 m and were used to determine the origin of air masses. The FTIR filter samples were grouped into sectors representing the origin of air masses as indicated by the top bars in Fig. 3a. Each air mass sector (along with consistent daytime onshore flow) allows analysis of particles originating from the same source region (most frequently the ports of Los Angeles and Long Beach) for a consistent number (1–3) of days. Example back trajectories associated with each sector are shown in Fig. 1. To further investigate likely sources of OM, potential source contribution function (PSCF) was applied to the factors resulted from factor analysis. PSCF classifies the back trajectories as “high” and “low” by concentration of the target component and calculates the probability that a source is located at a particular region (Pekney et al., 2006). Examples of PSCF results are shown in Fig. 5.
3.2 Organic and inorganic compositions of submicron particles

Table 1 summarizes the campaign average concentrations of the FTIR and AMS-measured components. Figure 3a shows the time series and the average fraction of organic functional groups measured by FTIR. The OM concentration varied from 0.39 to 11 µg m\(^{-3}\) with an average concentration of 3.3 ± 1.9 µg m\(^{-3}\), which was comparable to the OM concentration in the summer of 2008 (Hawkins and Russell, 2010a) and about twice the OM concentration measured in the winter of 2009 (Day et al., 2010) at the same site. Alkane functional group concentration was 1.5 ± 1.1 µg m\(^{-3}\) and contributed 47% of the OM. Carboxylic acid functional groups accounted for 34% of the OM with an average concentration of 1.1 ± 0.8 µg m\(^{-3}\). The concentration of hydroxyl functional groups was 0.40 ± 0.24 µg m\(^{-3}\), accounting for 12% of the OM. Amine, non-acid carbonyl, and organonitrate functional groups contributed small fractions to the OM (3%, 2%, and 2%, respectively). Alkene and aromatic functional groups were below detection limit for all samples and each was estimated to account for no more than 4% of the OM. These two functional groups were excluded from the analyses in this study. Samples associated with different air mass sectors had similar organic functional group compositions but differed in mass, with the Los Angeles-Long Beach port and the Riverside sectors containing significantly higher OM, indicating transport of pollutants from these heavily polluted regions to the sampling site, which is consistent with the findings of Ault et al. (2009). Only days associated with single air mass sector (32 out of 47 days) were included (Fig. 3a) in the diurnal cycle analysis in order to track the daily changes in compositions caused by chemistry rather than air mixing.

The non-refractory submicron particle mass (nrPM\(_1\)) concentration measured by the AMS showed similar variation to the FTIR OM concentration (Fig. 3b), indicating common sources or formation pathways of organic and inorganic compounds. Non-refractory OM (nrOM) was the largest component of nrPM\(_1\) (43%). Sulfate concentration was 3.1 ± 1.7 µg m\(^{-3}\) and accounted for 39% of nrPM\(_1\). Ammonium and nitrate contributed 12% and 5% of nrPM\(_1\), respectively. Only non-refractory species of chloride...
were measured by the AMS, and these accounted for a negligible amount of nrPM$_1$ with a mass fraction of 0.3% (Table 1).

The FTIR and AMS-measured OM compared reasonably well as shown in Fig. 4. The slope (1.1) and correlation coefficient ($r = 0.7$) of the OM correlation in this study fall into the typical ranges of the FTIR and AMS-measured OM comparison as summarized by Russell et al. (2009b) from eight previous field campaigns. These values are comparable to the average slope (1.1) and $r$ (0.67) derived from the Scripps Summer 2008 and the Scripps Winter 2009 projects (Russell et al., 2009b). Pearson’s correlation coefficient for a reduced major axis regression is used in this study. For samples associated with low sulfate fractions, the trend becomes more scattered from the 1:1 line and the sulfate-based CE has better agreement when sulfate accounts for more than 20% of nrPM$_1$. This is consistent with the fact that the sulfate-based CE used here was developed for ambient samples associated with relatively high sulfate fractions that were about 40% on average (Quinn et al., 2006).

### 3.3 Diurnal cycles of organic and inorganic components

For the diurnal cycle analyses, the concentrations were normalized by the enhancement of carbon monoxide ($\Delta$CO) mixing ratio (with respect to the background or unpolluted CO mixing ratio measured at the site during the study). This background CO mixing ratio was determined as the y-intercept of the linear regression of CO versus OM (DeCarlo et al., 2008), which differed slightly for the FTIR OM and the AMS nrOM at 89 ppb and 80 ppb, respectively. The difference of the two intercepts results from the larger AMS nrOM (10% higher than the FTIR OM). Since the difference of the two intercepts is insignificant (10%), an average value of 85 ppb was used as the background CO mixing ratio. Normalizing the concentration by CO, which does not react significantly on timescales of a few days, is used to separate the variations caused by the change of combustion source strength and effective dilution rates with variable meteorological conditions (which vary with $\Delta$CO) from changes associated with aerosol processing in the atmosphere (which do not vary with $\Delta$CO) (De Gouw et al., 2005, 2008; Gilardoni et al., 2009; DeCarlo et al., 2010).
The diurnal profiles of normalized carboxylic acid group concentrations are classified into four types based on their diurnal pattern (Fig. 6). The features for the four types are (1) the concentration continuously increased during the day and peaked at night (Type 1), (2) the peak concentration occurred in the afternoon (Type 2), (3) the concentration peaked both at noon and at night (Type 3), and (4) the peak concentration occurred at noon (Type 4). The corresponding alkane group diurnal profiles (for the same samples as in Types 1–4 shown in Fig. 6) resemble that of carboxylic acid groups (Fig. 6), indicating alkane and carboxylic acid groups likely were forming in the same molecules, at the same time, and condensing simultaneously.

The four types of diurnal profiles share similar properties that all showed higher concentrations during the solar maximum relative to the early morning period. However, Type 1 and Type 2 both increase from early morning till noon and then peak higher in the late afternoon (“Afternoon High”), but they differ in nighttime concentrations with Type 1 showing high and Type 2 showing low nighttime concentrations. Analogously, Type 3 and Type 4 have similar daytime patterns peaking at noon (“Noon High”) but differing nighttime concentrations. The variation in concentration at night could be explained by land-sea breeze circulations, with Type 3 having high and Type 4 having low nighttime concentrations. The diurnal variations of local wind directions for the four types of days are shown in Fig. 2. The difference in nighttime winds explains the differences between the low and high nighttime concentrations. Nighttime winds coming from the northwest dominated in Type 1 days while easterlies prevailed at night in Type 2 days. This change in wind direction likely resulted in Type 1 being more influenced by marine air masses and Type 2 being more influenced by land air masses at night. Similarly, the profile difference between Type 3 and Type 4 is likely caused by the differing influence of nighttime easterlies. In order to focus on the diurnal changes in chemical composition, our analysis has focused only on the daytime patterns allowing us to group the four types of diurnal cycles into two metaclasses: Type 1 and 2 are merged into the Class A (“Afternoon High”) with carboxylic acid group concentration peaked in the afternoon and similarly, Type 3 and 4 are merged into the Class
B ("Noon High") with noon-high concentrations. The daytime variations of carboxylic groups are consistent with the profiles of O₃ mixing ratios (Fig. 6) for the individual types, with O₃ peaked in the afternoon for the “Afternoon High” cases (Types 1 and 2) and peaked at noon for the “Noon High” cases (Types 3 and 4), indicating O₃-driven acid formation mechanism.

Figure 7 shows diurnal profiles of the AMS-measured fragment m/z 44 and the AMS Aged Combustion factor (identified from the factor analysis in Sect. 3.4.2) divided into the “Afternoon High” (Type 1 and 2) and the “Noon High” (Type 3 and 4) days. Fragment m/z 44, which is CO₂⁺ and representative of highly oxygenated organic components, tracked well with the AMS Aged Combustion factor. These two components peaked in the afternoon (14:00–18:00) in the “Afternoon High” days and the peak concentrations extended from noon to the afternoon (10:00–18:00) in the “Noon High” days, which are reflected in the average diurnal profiles for the two classes and consistent with the diurnal profiles of carboxylic acid group concentrations. The daytime peaks of m/z 44 and the Aged Combustion factor indicate enhanced SOA formation during the day (De Gouw and Jimenez, 2009). Carboxylic acid group concentration correlated well with m/z 44 and the AMS Aged Combustion factor concentrations for both “Afternoon High” and “Noon High” classes (Fig. 8), with the latter having better correlation coefficients of both carboxylic acid groups to m/z 44 and to the Aged Combustion factor (r = 0.8) but similar slopes. The correlation of acid and m/z 44 indicates that for these organic compositions, carboxylic acid groups and m/z 44 were likely associated with the same molecules and same formation mechanisms.

To evaluate evidence for photochemical processing in single particles, the average X-ray absorption spectra of single particles are shown in Fig. 9. Twenty-one morning particles and sixteen afternoon particles (collected on the same days as the morning particles) were analyzed. The afternoon-particle spectrum was characterized by a flat and broad black carbon peak at 285 eV, a broad peak ranging from 287 to 289 eV (peaks at 288.7 eV) indicative of alkane (the shoulder at 287.7 eV) and carboxylic acid (288.7 eV) functional group absorption, a carbonate peak at 290.4 eV, and two potassium peaks.
at 297.4 and 299.9 eV (Russell et al., 2002). These spectra are comparable to the spectrum of CaCO$_3$ (Hawkins and Russell, 2010b) except for the carboxylic acid group absorption, suggesting that carboxylic acid groups may condense on marine particles originating from marine calcareous phytoplankton (Hawkins and Russell, 2010b) when air masses passed over the ocean. Compared to the afternoon spectrum, the morning spectrum had a narrower and sharper black carbon peak with more variations at all energies. The graphite content of particles is represented by calculating the $\%sp^2$ hybridization for each particle (Takahama et al., 2007). Morning particles were associated with greater $\%sp^2$ hybridization than afternoon particles at a 84% confidence level, indicating morning particles were more influenced by primary emissions that likely included black carbon cores.

### 3.4 PMF factors

PMF (Paatero and Tapper, 1994) can be used to separate the contributions of different sources, each with characteristic compositions, to the multicomponent mixtures in ambient organic and inorganic particles (Pekney et al., 2006). We applied PMF to the FTIR spectra and the AMS-measured organic fragment concentrations separately to identify robust, linearly-independent components that compose the OM (Lanz et al., 2007; Russell et al., 2009a).

#### 3.4.1 PMF of the FTIR spectra

The FTIR PMF input matrix consisted of 234 mass-weighted and baselined FTIR spectra. The scaling factors were estimated by baselining errors calculated using the automated algorithm described by Russell et al. (2009a). The robust mode was used and the outliers were downweighted during the fitting procedure. Two to six factors with an FPEAK-range of $[0, \pm 0.2, \pm 0.4, \pm 0.6, \pm 0.8, \pm 1]$ were tested. The Q-value versus FPEAK plot showed a “U” shape with the lowest Q values corresponding to FPEAK of $-0.2, 0, 0.2, 0.4,$ and $0.6$, which resulted in the same factors. The edge-FPEAK...
values (−0.6, −0.4, ±0.8, and ±1) resulted in increased Q values, indicating increased residuals associated with the PMF model (Lanz et al., 2007). Because the sensitivity to rotation was negligible for FPEAK = −0.2, 0, 0.2, 0.4, and 0.6, FPEAK = 0 was selected to represent the solution.

Two- and three-factor solutions were excluded since only 85% of the OM was reproduced, while four or more factors reconstructed more than 95% of the OM. Correlated factors (r > 0.5) with similar compositions were combined to one factor (Liu et al., 2009), resulting in three factors for each of the four-, five-, and six-factor solutions. The three recombined factors resulted from the six-factor solution were selected because these factors captured events that were associated with trajectories from either know wildfires and from Los Angeles-Long Beach ports. In addition, the factor profiles had similar peak structure (r > 0.8) with the known factors derived from the TEXAQS/GoMACCS 2006 and the Scripps Pier 2008 measurements (Hawkins and Russell, 2010a; Russell et al., 2009a).

The first factor spectrum correlated to the fossil fuel Combustion factor profiles of the TEXAQS/GoMACCS (Russell et al., 2009a) and the Scripps Pier 2008 measurements (Hawkins and Russell, 2010a) projects with r being 0.97 and 0.99, respectively, indicating similar organic compositions, likely resulting from similar sources or processes. This factor was characterized by large fractions and co-existence of alkane and carboxylic acid functional groups (51% and 42% of the factor OM, respectively) and was identified as a fossil fuel Combustion factor. The concentration of this factor was 3.0 µg m⁻³, accounting for 62% of the OM on average (Fig. 3c). Hydroxyl and amine functional groups contributed 7% and 1% of the factor OM, respectively. PSCF image (Fig. 5a) shows the origin of this factor was mainly located at the vicinity of Los Angeles region, which are dominated by fossil fuel combustion emissions. The second factor spectrum correlated to the Biomass Burning factor profiles identified from the TEXAQS/GoMACCS (Russell et al., 2009a) and the Scripps Pier 2008 measurements (Hawkins and Russell, 2010a) with r = 0.87 and 0.93, respectively. The factor time series (Fig. 3c) showed three high concentration periods: 26 August–2 September,
8–22 August, and 22–27 September, corresponding to the top three fires in acreage that occurred in California: the Station fire (in Los Angeles County), the La Brea fire (in Santa Barbara County), and the Guiberson fire (in Ventura County), respectively (http://www.fire.ca.gov/fire_protection/fire_protection_fire_info_redbooks_2009.php). PSCF image (Fig. 5b) indicates that this factor is likely from north of Los Angeles – Santa Barbara County, as well as Baja California regions, consistent with fire events that occurred during the sampling period and fire maps from satellite measurements (Fig. 5c). Based on the similarity of this factor spectrum to previously identified biomass burning factors and the increase during fire-influenced time periods, this factor was identified as a Biomass Burning factor. The factor concentration was 0.88 µg m\(^{-3}\) on average and accounted for 18% of the OM. Non-acid carbonyl and alkane functional groups dominated this factor, accounting for 44% and 34% of the factor OM, respectively. The factor spectrum of the third factor was comparable \((r = 0.82)\) to the spectrum of the polluted marine factor described in the Scripps Pier 2008 measurements (Hawkins and Russell, 2010a) and was nearly identical to a sample of seawater atomized and collected for FTIR analysis (Gaston, 2011). This factor was identified as a Marine factor that accounted for 20% of the OM on average and was dominated by hydroxyl functional groups (72%). Alkane, carboxylic acid, and amine functional groups contributed 20%, 4%, and 3% of the OM, respectively. The concentration and composition of the factors are summarized in Table 2.

### 3.4.2 PMF of organic fragment concentrations

The input matrix and the error files for PMF of the AMS measurements were prepared using the Igor Pro 5 (Wavemetrics Inc.) codes based on the work of Zhang et al. (2005). Two to six factors with FPEAK-range of \([0, \pm 0.2, \pm 0.4, \pm 0.6, \pm 0.8, \pm 1]\) were investigated. The Q-value versus FPEAK plot shows a “U” shape with the lowest Q values corresponding to FPEAK of \(-0.2, 0,\) and \(0.2\). The factors generated for each rotation were nearly indistinguishable. FPEAK = 0 was selected to represent the solutions. A distinct factor with significant mass was missing when two factors were
used. For each of the four-, five-, and six-factor solutions, highly correlated factors ($r > 0.7$) were combined, resulting in three recombined factors, which resembled the three factors generated from the three-factor solution, indicating factors were split into smaller but indistinguishable components when more than three factors were used. The three-factor solution was selected and 98% of the OM was reproduced.

The factors were identified by comparing normalized factor spectra with the online AMS reference spectra (Ulbrich et al., 2007, 2009). The first factor spectrum correlated to several LV-OOA (low-volatility OOA) and SOA profiles. For example, the factor spectrum correlated to the Pittsburgh OOA factor profile (Zhang et al., 2005; $r = 0.93$ for m/z > 44 and $r = 0.95$ for all m/z) and SOA profile from the photooxidation of m-xylene with seed aerosols under RH = 55% (Bahreini et al., 2005; $r = 0.94$ for m/z > 44 and $r = 0.90$ for all m/z). The factor could not be further split into LV-OOA and SV-OOA (semi-volatile OOA) factors as shown in many previous studies (Ng et al., 2010), likely because of the relatively low particle concentration at the sampling site. The diurnal cycle of this factor showed a significant increase in concentration during the day and decrease in the morning and night (Fig. 7b), indicating photochemical origins of this factor. This factor likely represented an aged component formed from processed primary emissions. The factor was identified as an Aged Combustion factor, which accounted for 61% of the AMS nrOM and was associated with the largest m/z 44 nrOM fraction and m/z 44 to m/z 43 ratio among all factors (Table 2). The second factor profile strongly correlated with the wood burning spectrum ($r = 0.90$ for both m/z > 44 for all m/z) identified by Lanz et al. (2007) and the brush fire spectrum ($r = 0.94$ for m/z > 44 and $r = 0.92$ for all m/z) described by Bahreini et al. (2005). This factor was identified as a Biomass Burning factor, accounting for 26% of the nrOM. No correlation was found between the third factor spectrum and the spectra from the AMS database. The factor concentration correlated to none of the concentrations of the AMS-measured inorganic compounds. The factor profile correlated moderately ($r = 0.5$) with the Arctic Organic Aerosol (AOA) factor (which likely originates from the ocean) from the ICEALOT study (Frossard et al., 2011) and the time series correlated to that of the FTIR Marine factor.
with \( r = 0.5 \). This factor likely represented a Marine factor, which accounted for 13% of
the nrOM.

### 3.4.3 Comparison of PMF factors identified from the FTIR and AMS measurements

The reconstructed OM from the FTIR and the AMS PMF analyses correlated with
\( r = 0.7 \) and a slope of 1.1 (with the FTIR OM 10% higher). A Combustion factor,
a Biomass Burning factor, and a Marine-derived factor were identified from both the
FTIR and the AMS PMF analyses. The Combustion factors dominated the OM (ap-
proximately 60%) from both analyses and the time series correlated to each other with
\( r = 0.5 \) and a slope of 1.2 (FTIR factor OM was 20% higher). The Biomass Burning
factors resulted from the FTIR and the AMS measurements correlated with \( r = 0.7 \) and
slope = 0.9 (FTIR factor OM was 10% lower). The OM fractions of the two Biomass
Burning factors agreed with less than 10% difference, well within the expected accura-
cies of both FTIR and AMS OM. The two Marine factors correlated to each other with
\( r = 0.5 \), with the FTIR factor OM higher by a factor of 2.1. The weaker correlation and
the lower mass of the AMS Marine factor may be caused by a low collection efficiency
for organic compounds on NaCl particles as well as a lack of marine-aerosol specific
fragments in the AMS measurements. Alternatively it could result from lower absorp-
tivity of carbohydrate hydroxyl groups relative to the hydroxyl standards employed in
the quantification algorithm (Russell et al., 2009a). In summary, the factors identified
by PMF analyses of the FTIR and the AMS measurements agreed well and the mass
differences were within 30% except for the Marine factor. The good agreement shows
the consistency of the measurements and the robustness of the identified factors.
4 Discussion

In this section, we discuss carboxylic acid functional group formation mechanisms based on evidence from the diurnal profiles and size distributions. We use this evidence to estimate the SOA mass fraction in submicron particles associated with the different PMF factors. In addition, we identify factors that drive the variability in hydroxyl functional groups.

4.1 Photochemical formation of carboxylic acids

The daytime concentration of carboxylic acid groups tightly followed the variations of \( \text{O}_3 \) mixing ratios (Fig. 6). The two components correlated well with a correlation coefficient of 0.7 (Fig. 10). The correlation of carboxylic acid groups and odd oxygen (\( \text{O}_3 + \text{NO}_2 \)) could be better (Herndon et al., 2008), however, the \( \text{NO}_2 \) measurements were not available in this study. Atmospheric \( \text{O}_3 \) mixing ratio is affected by a number of parameters, including meteorological variables such as solar intensity, temperature, and RH and gas-phase chemical compositions such as \( \text{NO}_x \), volatile organic compounds (VOC), and other factors (Kleinman et al., 1994; Liu et al., 1994; Talbot et al., 2005). Days with afternoon \( \text{O}_3 \) peaks (the “Afternoon High” days) and noon \( \text{O}_3 \) peaks (the “Noon High” days) were associated with comparable variations and magnitudes of meteorological conditions, including temperature, ambient RH, wind direction and speed, and photosynthetically active radiation, suggesting the peak time of \( \text{O}_3 \) mixing ratio was likely affected by other parameters, such as \( \text{NO}_x \) and VOC mixing ratios (for which the measurements were not available).

The diurnal covariation of carboxylic acids and \( \text{O}_3 \) provides substantial evidence for an \( \text{O}_3 \)-driven oxidation that forms carboxylic acid groups. The potential precursors of carboxylic acids in anthropogenic source dominated regions include alkanes and aromatics, which are emitted from fossil fuel combustion activities. Smog chamber studies show that OH oxidation of aromatics yields multifunctional group products with low alkane group mole fractions (<0.3) and higher non-acid carbonyl group mole fractions.
Large mole fraction of alkane groups (0.85) and the absence of non-acid carbonyl groups of the FTIR Combustion factor rule out the aromatics as acid formation precursors. The carboxylic acid, hydroxyl, non-acid carbonyl, and alkane groups mole fractions of the Combustion factor were 0.11, 0.04, 0.00, and 0.85, comparable to the C_{12} alkane oxidation products with mole fractions of 0.12, 0.13, 0.02, and 0.73 estimated by Russell et al. (2011), suggesting that alkanes are likely carboxylic acid formation precursors. Russell et al. (2011) hypothesize a two-step oxidation of gas-phase alkanes to form carboxylic acids. First, gas-phase alkanes are oxidized by OH radicals to form particle-phase dihydrofuran via H-atom subtraction, isomerization, cyclization, and dehydration processes. Second, dihydrofurans evaporate into the gas phase and react with O_3, producing products that are expected to be similar to cyclic alkene oxidation products, namely multi-functional products with carboxylic acid functional groups (which would be expected to partition into the particle phase due to their low vapor pressures). This mechanism explains the co-variation of particle-phase carboxylic acid groups and alkane groups, and the similarity of their diurnal trends to O_3 mixing ratios in this measurement, suggesting carboxylic acids are second-generation products of alkane oxidation.

Representative size distributions of m/z 44 and the AMS nrOM are shown in Fig. 11 for the “Afternoon High” days (for time period of 14:00–18:00) and the “Noon High” days (for time period of 10:00–14:00). For both cases, the size distributions of the two components showed similar variations and both peaked around 300–500 nm. The OM fraction of m/z 44 was nearly independent of particle size, while nrOM/nrPM decreased with increasing particle diameter, comparable with a theoretical model of a surface-driven process (as shown in Fig. 11 iii and iv). The deviations from the model likely result from density differences of newly condensed mass. This type of size distribution is consistent with the “Condensation and Volume-limited Oxidation” mechanism proposed by Maria et al. (2004), indicating the condensation of newly formed nrOM onto existing particles (the mass fraction of nrOM increased with 1/Dp) and more m/z 44 were formed by further volume-limited oxidation in the particle phase (the mass of
Several chamber studies have suggested acid-catalyzed formation of SOA (Edney et al., 2005; Kleindienst et al., 2006; Surratt et al., 2007), while others have observed the opposite (Bahreini et al., 2005; Tan et al., 2009). Zhang et al. (2007) investigated acidity effects on Pittsburgh SOA and concluded that no enhancement of SOA was observed during high acidity measurement periods. For this study, when the molar equivalence ratio of ammonium to sulfate plus nitrate was used as a surrogate for the acidity of aerosol particles (Quinn et al., 2006), there was no correlation of acidity with carboxylic acid functional group concentration, although particles are in acidic conditions.

### 4.2 SOA mass fraction

The contributions to SOA from carboxylic acid groups and OOA are both estimated by assuming that the minimum concentration which occurs in the early morning is representative of a background value (either from other sources or from processing on previous days) and that the increase that occurs (relative to ΔCO) is from photochemical processing in the atmosphere during a 12-h daytime in a single day. The background and diurnal cycles of carboxylic acid groups and the Combustion factors from the FTIR and AMS measurements are shown in Fig. 12. The SOA mass fraction calculation is summarized in Table 3.

In general, the SOA fractions are comparable for the “Afternoon High” and the “Noon High” days with the differences within 30%. The 12-h daytime increase in carboxylic acid group mass in the “non-morning” periods compared to the low during the morning period approximately accounted for 30% of the carboxylic acid group mass and 10% of OM; similarly the photochemical increase in the FTIR Combustion factor accounted for 28% of the factor mass and 17% of the OM. The AMS Aged Combustion factor showed...
higher SOA mass fractions (41% and 23% for the factor mass and the OM, respectively) than the FTIR Combustion factor, but the two are within the measurement uncertainties of both of the techniques. The lower SOA fractions from acid-group-only-based estimates could be explained by the fact that the molecules contain other components besides the carboxylic acid groups (i.e. likely alkane groups; Lim and Ziemann, 2009). Comparing the estimates from the functional groups and the Combustion factors, we find that approximately 45–70% of the SOA molecules may be attributed to carboxylic acid groups and the remaining would be constituted by other organic functional groups.

4.3 Sources and characteristics of hydroxyl functional groups

Hydroxyl functional groups were mainly from marine sources, since their variation was uncorrelated with sulfate and the AMS and FTIR Combustion factors. Larger contributions from the hydroxyl groups and hence the Marine factor were associated with onshore winds from 07:00 to 22:00 (Fig. 13), which could bring concentrated marine aerosols to the sampling site, indicating the hydroxyl functional group concentration was modulated by the land-sea breeze circulation at the coastal site. The Marine factor concentration weakly correlated with local wind speed ($r = 0.3$ during onshore winds), consistent with the expected increase in particle production from wind-related bubble bursting on the ocean surface (Russell et al., 2010) but sufficiently influenced by other variables to mask a one-to-one link.

5 Conclusions

Measurements at the Scripps Pier showed that OM accounted for nearly 50% of ambient submicron particle mass. The OM was dominated by fossil fuel combustion type sources, which resulted in a mixture of alkane and carboxylic acid functional groups. In this mixture, the abundances of these two functional groups changed diurnally, with the carboxylic acid group concentration increasing significantly during the daytime,
indicating photochemical formation of carboxylic acid groups, consistent with past studies that have argued carboxylic acids are secondary. This daytime increase was tightly correlated with O$_3$ mixing ratios, suggesting O$_3$ played an important role in acid formation. A two-step alkane oxidation mechanism is consistent with the observed diurnal variation of carboxylic acid groups and the composition of the Combustion factor resulted from the PMF analysis. In this mechanism, gas-phase alkane molecules are oxidized by OH radicals, forming dihydrofuran in the particle phase, followed by evaporation of dihydrofuran and further gas-phase oxidation by O$_3$. The O$_3$ peaked both at noon and in the afternoon, likely caused by different NO$_x$ or VOC conditions upstream, resulting in two classes of carboxylic acid group and alkane group daytime profiles (the “Afternoon High” and the “Noon High” days) under comparable meteorological conditions. In both cases, the size distribution of $m/z$ 44/nrOM and nrOM/PM$_1$ revealed that the newly formed nrOM condensed onto existing particles, while $m/z$ 44 may undergo further reactions in the particle phase under high RH at the sampling site.

The fraction of OM that is contributed by secondary carboxylic acid groups is estimated to be 10%. This number represents a lower-bound of the SOA contribution to OM, as other groups are also present in photochemically-generated molecules (i.e. alkane groups). Another estimate is to consider the fraction of OOA that is formed each day as secondary, which is 15–30% of OM. These measurements account for only the SOA formed during an approximately 12-h period, yet it contributes a large fraction of the combustion-associated OM factors (50% for FTIR) and a significant fraction of OM at the Scripps Pier during this study. This large contribution of alkane-derived SOA is not surprising given their large contributions to VOC emissions in the LA/LB source region and the relative absence of significant additional organic precursors during transit in the coastal marine boundary layer. Our study reinforces the importance of gas-phase alkane photochemistry for the air quality of regions downwind of emission sources.

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3. References

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Table 1. Campaign average and standard deviation of FTIR-measured OM, FTIR organic functional group concentrations, and AMS-measured OM, sulfate, ammonium, nitrate, and chloride concentrations in $\mu m^{-3}$. Functional group OM mass fractions (for FTIR) and component PM$_1$ mass fractions (for AMS) are shown in the parentheses.

<table>
<thead>
<tr>
<th></th>
<th>FTIR</th>
<th>AMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>OM</td>
<td>3.3±1.9</td>
<td>3.9±3.0</td>
</tr>
<tr>
<td>Alkane</td>
<td>1.5±1.1 (47%)</td>
<td>3.1±1.7 (39%)</td>
</tr>
<tr>
<td>Carboxylic Acid</td>
<td>1.1±0.77 (34%)</td>
<td>1.0±0.53 (12%)</td>
</tr>
<tr>
<td>Hydroxyl</td>
<td>0.40±0.24 (12%)</td>
<td>0.43±0.49 (5%)</td>
</tr>
<tr>
<td>Amine</td>
<td>0.11±0.09 (3%)</td>
<td></td>
</tr>
<tr>
<td>Non-Acid Carbonyl</td>
<td>0.06±0.21 (2%)</td>
<td></td>
</tr>
<tr>
<td>Organonitrate</td>
<td>0.07±0.12 (2%)</td>
<td></td>
</tr>
</tbody>
</table>
Table 2. Concentration and composition of PMF factors identified from the FTIR and the AMS measurements. Colors in the pie charts indicate alkane (blue), carboxylic acid (green), hydroxyl (hot pink), amine (orange), and non-acid carbonyl (teal) functional groups. Mass fractions of the factors are shown in the parentheses.

<table>
<thead>
<tr>
<th></th>
<th>Aged Combustion</th>
<th>Biomass Burning</th>
<th>Marine Origin</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FTIR</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concentration (µm⁻³)</td>
<td>3.0 (62%)</td>
<td>0.88 (18%)</td>
<td>0.97 (20%)</td>
</tr>
<tr>
<td>O/C</td>
<td>0.46</td>
<td>0.48</td>
<td>1.04</td>
</tr>
<tr>
<td><strong>Composition</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>AMS</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concentration (µm⁻³)</td>
<td>2.5 (61%)</td>
<td>1.1 (26%)</td>
<td>0.51 (13%)</td>
</tr>
<tr>
<td>(m/z 44)/OM</td>
<td>26%</td>
<td>5%</td>
<td>2.6%</td>
</tr>
<tr>
<td>(m/z 60)/OM</td>
<td>0.1%</td>
<td>0.1%</td>
<td>0.6%</td>
</tr>
<tr>
<td>(m/z 44)/(m/z 43)</td>
<td>7.5</td>
<td>0.6</td>
<td>1.5</td>
</tr>
<tr>
<td>(m/z 44)/(m/z 57)</td>
<td>146</td>
<td>2.7</td>
<td>3.6</td>
</tr>
</tbody>
</table>
Table 3. 12-h daytime SOA fraction from a single day calculated from carboxylic acid groups, the FTIR Combustion factor, and the AMS Aged Combustion factor for the “Afternoon High” and the “Noon High” days. The OM fractions of SOA are shown in the parentheses.

<table>
<thead>
<tr>
<th></th>
<th>Carboxylic acid group</th>
<th>FTIR Combustion factor</th>
<th>AMS Aged Combustion factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Afternoon High</td>
<td>27% (9%)</td>
<td>33% (20%)</td>
<td>45% (27%)</td>
</tr>
<tr>
<td>Noon High</td>
<td>30% (10%)</td>
<td>23% (14%)</td>
<td>37% (19%)</td>
</tr>
</tbody>
</table>
Fig. 1. Example back trajectories representing Los Angeles-Long Beach (cyan; air mass coming from Los Angeles and Long Beach regions), Riverside (brown; air mass originating from Riverside vicinity), Inland (orange; easterly/northeasterly air mass), Tijuana-Ensenada (purple; southerly air mass), Mixed coastal (magenta; northerly air mass coming along the coast of California), and Ocean (dark blue; westerly air mass) air mass sectors during the campaign. The black circles (from top to bottom) indicate Riverside, Los Angeles, Los Angeles - Long Beach port, and the sampling site.
Fig. 2. Vector-averaged diurnal cycles of wind direction corresponding to the four types of days shown in Fig. 6. Shaded areas indicate nighttime periods.
Fig. 3. (a) Time series of organic functional group concentrations measured by the FTIR; sectors are indicated by the top color bars (same colors as in Fig. 1), for which the sector associated with each FTIR sample was determined as the air mass origin shown by the majority (>80%) of the back trajectories during the sampling time; top brown bars indicate fire periods corresponding to the La Brea fire (in Santa Barbara County), the Station fire (in Los Angeles County), and the Guiberson fire (in Ventura County), respectively (from left to right); top green bars indicate samples associated with single air mass sector, which were used for diurnal profile analysis. (b) Time series of AMS factors identified by PMF analysis. The inner pie charts in (a) and (b) respectively show campaign average compositions of FTIR components and AMS factors. (c) Time series of normalized O$_3$ (normalized by campaign average) mixing ratio. (d) Mass fractions of the FTIR Combustion factor (red), the Biomass Burning factor (brown), and the Marine factor (blue) during the measurement.
Fig. 4. Comparison of OM measured by the FTIR and the AMS. Colors indicate sulfate mass fraction of nrPM$_1$. Correlation coefficients for high sulfate mass samples (mass fraction >20%) and low sulfate mass samples (sulfate fraction <20%) are 0.8 and 0.6, respectively.
Fig. 5. Potential Source Contribution Function (PSCF) images of (a) the FTIR Combustion factor and (b) the FTIR Biomass Burning factor with warmer colors indicate higher probability. (c) Fire map on 29 August with red points showing fire spots and blue lines indicating back trajectories ending at the Scripps Pier. Fire image was obtained from NOAA’s Aqua satellite.
Fig. 6. Diurnal cycles of normalized carboxylic acid group concentrations (green) and alkane group concentrations (blue) divided into four types (1–4) shown in panels (1), (2), (3), and (4), respectively. Each rectangle represents one FTIR sample with the length of the rectangle indicating the sampling duration. The lines connecting the rectangles show samples collected in the same day. The thinner rectangles and lines represent daily diurnal profiles, while the thicker rectangles and lines show the averages for the days in the corresponding panel. The red dashed lines represent average diurnal profiles of normalized O$_3$ mixing ratio for the four types of days. Shaded areas indicate nighttime periods corresponding to the FTIR nighttime samples, which were excluded from the diurnal cycle analyses.
Fig. 7. Diurnal cycles of (i) $m/z$ 44, and (ii) the AMS Aged Combustion factor for A (“Afternoon High”) and B (“Noon High”) cases. The thinner lines represent daily diurnal cycles and the thicker lines represent the averages. The curves are smoothed using the “Boxcar Smoothing” method with 30-point averaging to reduce high-frequency noise in the measurements. Shaded areas indicate nighttime periods.
Fig. 8. Mass concentration correlation of carboxylic acid groups with (i) the AMS Aged Combustion factor and (ii) m/z 44 for A (“Afternoon High”) and B (“Noon High”) cases. The correlation coefficients are shown in the legends.
Fig. 9. Average X-ray spectra of (a) twenty-one morning and (b) sixteen afternoon particles. Grey vertical bars indicate the standard deviations of absorption at corresponding energies. Red vertical lines indicate alkene group (285 eV), alkane group (287.7 eV), carboxylic acid group (288.7 eV), carbonate (290.4 eV), and K (297.4 and 299.9 eV) absorptions.
Fig. 10. Correlation of normalized carboxylic acid group concentration and normalized O$_3$ mixing ratio (by campaign average) for the “Afternoon High” (orange) and “Noon High” (blue) cases. The correlation coefficient is 0.7.
Fig. 11. Representative size distributions for A ("Afternoon High" days) on 12, 16, and 21 September, in time period 14:00–18:00 and B ("Noon High" days) on 17 and 20 September in time period 10:00–14:00 for (i) m/z 44, (ii) nrOM, (iii) m/z 44/nrOM, and (iv) nrOM/PM$_1$. The curves are smoothed with the “Boxcar Smoothing” method with 20-point averaging and wavelet de-noising method to reduce high-frequency noise in the measurements. Red lines are theoretical models representing (iii) constant with respect to particle diameter and (iv) inversely proportional to the particle diameter (1/Dp), for comparison to results of Maria et al. (2004).
Fig. 12. Daytime profiles of A ("Afternoon High" days) and B ("Noon High" days) for (i) carboxylic acid group, (ii) the FTIR Combustion factor, and (iii) the AMS Aged Combustion factor concentration. Colors indicate POA (grey) and SOA (green), respectively. Red dashed lines in panels A-i and B-i represent average daytime profiles of normalized O$_3$. Black lines in panels A-i and B-i are the average diurnal carboxylic acid group profiles corresponding to the four types shown in Fig. 6 as indicated by the numbers beside the lines. Vertical blue bars in panels A-iii and B-iii show standard deviations of the averaged diurnal cycles.
**Fig. 13.** The FTIR Marine factor concentration versus wind direction. 25th and 75th percentiles are shown by the upper and lower box bounds in each box. The points show individual samples. Median values are represented by the horizontal bars in the boxes.