Laboratory photochemical processing of aqueous aerosols: formation and degradation of dicarboxylic acids, oxocarboxylic acids and α-dicarbonyls

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Abstract. To better understand the photochemical processing of dicarboxylic acids and related polar compounds, we conducted batch UV irradiation experiments on two types of aerosol samples collected from India, which represent anthropogenic (AA) and biogenic (BA) aerosols, for time periods of 0.5 to 120 h. The irradiated samples were analyzed for molecular compositions of diacids, oxoacids and α-dicarbonyls. The results show that photochemical degradation of oxalic (C₂), malonic (C₃) and other C₈–C₁₂ diacids overwhelmed their production in aqueous aerosols, whereas succinic acid (C₄) and C₅–C₇ diacids showed a significant increase (ca. 10 times) during the course of irradiation experiments. The photochemical formation of oxoacids and α-dicarbonyls overwhelmed their degradation during the early stages of experiment except for ω-oxooctanoic acid (ωC₈), which showed a similar pattern to that of C₄. We also found a gradual decrease in the relative abundance of C₂ to total diacids and an increase in the relative abundance of C₄ during prolonged experiment. Based on the changes in concentrations and mass ratios of selected species with the irradiation time, we hypothesize that iron-catalyzed photolysis of C₂ and C₃ diacids controls their concentrations in Fe-rich atmospheric waters, whereas photochemical formation of C₄ diacid (via ωC₈) is enhanced with photochemical processing of aqueous aerosols in the atmosphere. This study demonstrates that the ambient aerosols contain abundant precursors that produce diacids, oxoacids and α-dicarbonyls, although some species such as oxalic acid decompose extensively during an early stage of photochemical processing.

1 Introduction

Dicarboxylic acids and related polar compounds constitute a significant fraction of water-soluble organic aerosols in the atmosphere (Saxena and Hildemann, 1996; Kawamura and Sakaguchi, 1999; Pavuluri et al., 2010). They have a potential contribution to the formation of cloud condensation nuclei due to their water-soluble and hygroscopic properties (Saxena and Hildemann, 1996; Giebl et al., 2002). Thus diacids and related compounds have an impact on the indirect radiative forcing and hydrological cycle (Twomey, 1977; Albrecht, 1989). They also involve in a series of reactions occurring in gas phase, aerosols and atmospheric waters (Chebbi and Carlier, 1996; Z. H. Wang et al., 2010). Although diacids, oxoacids and α-dicarbonyls can be directly emitted into the atmosphere from incomplete combustion of fossil fuels (Kawamura and Kaplan, 1987) and biomass burning (Narukawa et al., 1999), they are mainly formed by secondary processes of volatile organic compounds of anthropogenic and biogenic origin (Kawamura et al., 1996a; Kawamura and Sakaguchi, 1999; Kanakidou et al., 2005).
They are further subjected to photochemical oxidation during long-range transport; e.g., carbonyls to carboxylic acids (Tilgner and Herrmann, 2010) and breakdown of higher to lower diacids (Kawamura and Sakaguchi, 1999; Matsunaga et al., 1999; G. Wang et al., 2010).

Molecular distributions of diacids in atmospheric aerosols have generally been reported with a predominance of oxalic (C$_2$) acid followed by malonic (C$_3$) or succinic (C$_4$) acid in different environments (Kawamura and Kaplan, 1987; Kawamura and Ikushima, 1993; Kawamura and Sakaguchi, 1999; Narukawa et al., 1999; Pavuluri et al., 2010). The predominance of C$_2$ in different environments is likely because it is an ultimate end product in the chain reactions of diacids and various precursors including aromatic hydrocarbons, isoprene, alkenes and α-dicarbonyls (Kawamura et al., 1996a; Warneck, 2003; Ervens et al., 2004b; Lim et al., 2005; Carlton et al., 2007; Charbouillot et al., 2012). In contrast, C$_4$ was reported to be more abundant than C$_2$ in some aerosol samples collected from Antarctica (Kawamura et al., 1996b), the Arctic (Kawamura et al., 2010) and over the Arctic Ocean (Kawamura et al., 2012) as well as in ice core samples from Greenland (Kawamura et al., 2001). In addition, a significant reduction in C$_2$ diacid concentration and an inverse relationship between C$_2$ and Fe has been reported in stratocumulus clouds over the northeastern Pacific Ocean (Sorooshian et al., 2013). The predominance of C$_4$ over C$_2$ in ice core samples and atmospheric aerosols from polar regions, particularly in the Arctic marine aerosol samples collected under overcast conditions with fog or haze events (Kawamura et al., 2012) and the reduction of C$_2$ in cloud water, suggest that photochemical formation of C$_4$ and/or degradation of C$_2$ (Pavuluri and Kawamura, 2012) should be enhanced in atmospheric waters.

However, the photochemical formation and degradation of diacids and related compounds are not fully understood, particularly in aqueous phase because the composition of aqueous solutions used in laboratory experiments do not reflect the complex mixture of organic and inorganic aerosol constituents in the atmosphere and the experimental conditions are not necessarily atmospherically relevant (Ervens et al., 2011). Hence, it is required to investigate the fate of diacids and related polar compounds with photochemical processing in atmospheric waters. In this study, we conducted a laboratory experiment using two types of ambient aerosol samples collected from Chennai, India, which represent anthropogenic (AA) and biogenic (BA) aerosols. The samples were exposed to UV irradiation in the presence of moisture for different time ranging from 0.5 to 120 h and then analyzed for diacids, oxoacids and α-dicarbonyls. Here, we report their molecular compositions and discuss the photochemical formation and/or degradation of diacids as a function of the irradiation time. Based on the results obtained, we propose possible photochemical formation and degradation pathways of diacids and related compounds with atmospheric implications.

### Figure 1
A map of South Asia with sampling site, Chennai (13.04°N, 80.17°E), India, together with plots of 10-day air mass trajectories arriving at 500 m a.g.l. over Chennai, India.

## 2 Materials and methods

### 2.1 Atmospheric aerosol samples

In this study, we used two types of atmospheric aerosol (PM$_{10}$) samples that were collected in winter on 28 January (IND104) and in summer on 25 May (IND178) 2007 during daytime (ca. 06:00–18:00 h local time) from Chennai (13.03° N, 80.17° E), India, using a high volume air sampler and pre-combusted (450 °C, 4 h) quartz fiber filters. Sampling was conducted on the rooftop of the mechanical sciences building (~18 m a.g.l., above the ground level) at the Indian Institute of Technology Madras (IITM) campus. The details of sampling site and meteorology are described elsewhere (Pavuluri et al., 2010). The sample filter was placed in a preheated glass jar with a Teflon-lined screw cap and stored in darkness at −20 °C prior to the experiment. Figure 1 presents 10-day backward air mass trajectories arriving in Chennai at 500 m a.g.l. for every 6 h during the sampling periods of IND104 and IND178. Table 1 shows concentrations of elemental carbon (EC), organic carbon (OC), levoglucosan and sums of hopanes (specific biomarkers of petroleum and coal) and lipid class compounds (fatty acids and fatty alcohols) in IND104 and IND178 (Fu et al., 2010; Pavuluri et al., 2011).

The air mass trajectories showed that the air masses for the IND104 sample originated from the north Indian subcontinent passing over the Bay of Bengal (Fig. 1). In North India, anthropogenic emissions are mainly derived from fossil fuel combustion and forest fires (Lelieveld et al., 2001; Reddy and Venkataraman, 2002a). This sample is enriched with EC (Table 1). The anthropogenic signature of IND104 is further...
supported by high abundances of hopanes. In contrast, the air masses for the IND178 sample originated from the Arabian Sea passing over the south Indian subcontinent (Fig. 1), where the emissions from marine biota, combustion of biofuels (e.g., cow dung) (Reddy and Venkataraman, 2002b) and livestock (Garg et al., 2001) are important. In addition, emission of volatile organic compounds (VOCs) from tropical plant species in India is enhanced in summer (Padhy and Varshney, 2005). This sample is enriched with OC but EC is less abundant (Table 1). The biogenic signature of IND178 is supported by high abundances of fatty acids and fatty alcohols (Table 1). Hence, we conclude that IND104 represents AA whereas IND178 represents BA.

2.2 Determination of trace elements, metals and water-soluble iron species

Trace elements and metals were determined using an inductively coupled plasma mass spectrometry (Thermo Electron X Series) after the acid microwave digestion of samples (a filter disc of 1.8 cm in diameter) as reported by Theodosi et al. (2010b). Recoveries obtained with the use of certified reference materials ranged from 90.0 to 104.1 %. Water-soluble iron (Fe_{WS}: sum of Fe^{2+} and Fe^{3+} species) was determined spectrometrically using the ferrozine colorimetric method developed by Stookey (1970) as reported by Theodosi et al. (2010a). Fe^{2+} was measured using the same procedure without adding the reducing agent (hydroxylamine hydrochloride), and then Fe^{3+} was estimated indirectly as the difference between Fe_{WS} and Fe^{2+}. The recovery was ∼98.3 % for both Fe_{WS} and Fe^{2+}.

2.3 Irradiation experiment of aerosol samples

Batch UV irradiation experiments using two aerosol samples (AA and BA) were conducted separately for 0.5, 1.5, 3.0, 6.0, 12, 18, 24, 36, 48, 72, 96 and 120 h, because both primary and secondary chemical species associated with aerosols can be subjected to significant photochemical processing throughout their stay (i.e., up to 12 days) in the atmosphere (Warneck, 2003). In each experiment, ∼12 cm² (ca. 3 × 4 cm) of sample filter was cut into 3 × 4 pieces and placed vertically in a cleaned quartz reaction vessel (cylinder, 100 mL) with the sample surface facing to UV light as depicted in Fig. 2. The sample was fully wetted by injecting ∼0.4 mL of ultra pure organic free Milli-Q water and sealed with Teflon-lined screw cap under the ambient pressure. Furthermore, the available excess Milli-Q water (Fig. 2) may promote humid (RH = 100 %) environment in the reaction vessel by equilibrium between water vapor and Milli-Q water. The aqueous ambient aerosol sample was then irradiated with a low-pressure mercury lamp (Ushio, UL0-6DQ) that emits a UV whose spectra are characterized by a main peak at 254 nm and minor peak at 185 nm as well as a broad peak at > 254 nm. The experimental setup (Fig. 2) was covered with a cardboard box containing a hole on each side for the passage of ambient air and placed in a draft chamber. The temperature around the experimental system (i.e., inside cardboard box) was equivalent to room temperature (25 ± 1 °C).

The main objective of UV irradiation with a wavelength primarily at 254 nm, rather than a solar spectrum, was to produce significant amounts of hydroxyl radicals (HO*) from various sources (described below), which should be sufficient enough to act as the main oxidant in our experimental system. Although we do not preclude a minor photolysis of some organic compounds present in the aerosol samples by irradiation at ≥ 254 nm, it is well established that low molecular weight diacids, oxoacids and α-dicarboxylic acids...
including pyruvic acid and methylglyoxal have negligible absorbance at 254 nm and exhibit minimal photolysis, particularly when HO• reactions of organics are significant (Carlton et al., 2006; Yang et al., 2008b; Tan et al., 2012). Because sulfate is abundant in non-irradiated AA and BA (Pavuluri et al., 2011), the production of organosulfates could be significant upon irradiation (Noziere et al., 2010) in both the samples. However, the sulfate contents may not have a significant impact on the production rate of diacids and related compounds (Tan et al., 2009).

Furthermore, the photolysis of organics by the radiation of 185 nm, the intensity of which is 100 times lower than that of 254 nm, should be insignificant during the experiment because the 185 nm light intensity is small and is mostly absorbed by water due to its high absorption coefficient (1.8 cm\(^{-1}\) at 25°C) (Weeks et al., 1963). It is well established that the photolysis rates of organics by the radiation of 185 nm are elevated only in the absence of dissolved oxygen (Shirayama et al., 2001); however, in aerated solutions under irradiation of 185 nm, mainly water undergoes photolysis and produces the hydrogen atoms, which are scavenged by oxygen to form HO• radicals that are finally converted to HO• (Chitose et al., 2003; Yang et al., 2008b). Thus, the minor 185 nm light emitted from the UV source promotes the production of HO• during the experiment rather than the photolysis of the compounds of interest in this study. On the contrary, iron–dicarboxylate complexes (e.g., oxalate and malonate) can photolyze by absorbing both UV-C (254 nm) and UV-A light and their photolysis rate depends on the concentration of Fe in the given sample rather than the UV light wavelength (Zuo and Hoigne, 1994; Z. H. Wang et al., 2010; Pavuluri and Kawamura, 2012). In addition, radiation at 254 nm has been reported to impose only a marginal photolysis of most of the inorganic species except for nitrate, which is one of the HO• sources (Yang et al., 2008b).

The irradiation of wet aerosol sample at 254 nm induces the formation of O\(_3\) from the dissolved O\(_2\), followed by the generation of H\(_2\)O\(_2\); the photolysis of H\(_2\)O, NO\(_3\), NO\(_2\), H\(_2\)O\(_2\), Fe(OH)\(^{2+}\) and certain organic compounds; and Fenton’s reaction of photochemically formed Fe\(^{2+}\) and H\(_2\)O\(_2\) to produce HO• in aqueous phase (Arakaki and Faust, 1998; Carlton et al., 2006; Yang et al., 2008b). In fact, a large amount of Fe, including water-soluble Fe\(^{2+}\) and Fe\(^{3+}\) species, is available in both AA and BA samples (Table 1), which could promote the Fenton’s reaction upon UV irradiation. In addition, O\(_3\), H\(_2\)O\(_2\), HO• and NO\(_2\) formed in aqueous phase reactions may be partitioned into gas phase and generate the gaseous HO• that should be re-partitioned into aqueous phase (Arakaki and Faust, 1998). These sources of HO• are similar to those of atmospheric waters: (i) gas/drop partitioning of HO• and (ii) gas/drop partitioning of O\(_3\) followed by a reaction with peroxy radical (HOО•); (iii) photolysis of H\(_2\)O, NO\(_3\), NO\(_2\), H\(_2\)O\(_2\), Fe(OH)\(^{2+}\) and certain organic compounds; and (iv) Fenton’s reaction of Fe\(^{2+}\) and H\(_2\)O\(_2\) (Arakaki and Faust, 1998).

Unfortunately, we could not approximate the actual concentrations of HO• in our experiments because we did not add any chemical (e.g., a standard compound whose kinetics are known) in order to keep our experimental system as realistic as possible. Furthermore, the formation of O\(_3\) from the initially available O\(_2\) (∼0.94 mM) in the reaction vessel may not cause the deficit of the O\(_2\) that could potentially induce the polymerization of organics during the irradiation on aerosols for several hours, because the additional O\(_2\) could be produced from the gaseous HOО• formed by photolysis of organics and Fenton’s reaction (Arakaki and Faust, 1998) during the experiment.

2.4 Measurements of diacids, oxoacids and α-dicarboxyls

Immediately after the irradiation, samples were analyzed for diacids, oxoacids and α-dicarboxyls using a method reported elsewhere (Kawamura, 1993; Kawamura and Ikushima, 1993). Briefly, the irradiated sample filter was extracted with Milli-Q water (10 mL × 3) under ultra sonication for 10 min and the extracts were concentrated to near dryness using a rotary evaporator under vacuum. The extracts were then derivatized with 14% BF\(_3\)/n-butanol at 100°C to butyl esters and/or butoxy acetals. Both the esters and acetals were extracted with n-hexane and then determined using a capillary gas chromatograph (HP 6890) and gas chromatography–mass spectrometer (Thermo Trace MS). Recoveries of authentic standards spiked to a pre-combusted quartz fiber filter were 73% for oxalic (C\(_2\)) acid and more than 84% for malonic (C\(_3\)), succinic (C\(_4\)) and adipic (C\(_6\)) acids (Pavuluri et al., 2010). The analytical errors in duplicate analysis of the aerosol filter sample are within 9% for major species.
Gas chromatogram of the field and laboratory blanks showed small peaks for $C_2$, phthalic (Ph) and glyoxylic acids. Concentrations of all the species reported here are corrected for the non-irradiated field blanks (Pavuluri et al., 2010).

2.5 Quality control

To examine the possible experimental errors, including the distribution of organic/inorganic constituents over the filter sample, we conducted replicate experiments ($n = 3$) for 18 h irradiation of the AA sample by using the sample cut taken from different parts of the filter sample for each experiment, because a deviation in the results of the irradiation experiment should become large if the impact of potential variance in chemical composition of aerosol at different parts of the single filter, size of the filter sample used (i.e., number of aerosols) and the amount of Milli-Q water added is significant. The experimental errors, including the analytical errors, were found to be within 11 % for major species except for $C_1$ aerosols) and the amount of Milli-Q water added is significant. To examine the possible experimental errors, including the distribution of organic/inorganic constituents over the filter sample, we conducted replicate experiments ($n = 3$) for 18 h irradiation of the AA sample by using the sample cut taken from different parts of the filter sample for each experiment, because a deviation in the results of the irradiation experiment should become large if the impact of potential variance in chemical composition of aerosol at different parts of the single filter, size of the filter sample used (i.e., number of aerosols) and the amount of Milli-Q water added is significant. The experimental errors, including the analytical errors, were found to be within 11 % for major species except for $C_1$ aerosols. These results suggest that organic and inorganic constituents are well distributed over the filter sample and took up water evenly distributed upon wetting. In addition, two irradiation experiments were conducted to check the procedural blank by using a clean quartz filter for 1.5 and 6.0 h. No peaks were detected except for a small peak for $C_2$ and Ph. These results indicate that the occurrence ofbias during the experiment is insignificant.

3 Results and discussion

3.1 Concentrations of trace elements, metals and water-soluble iron species

Concentrations of trace elements, metals and water-soluble Fe species ($Fe^{2+}$ and $Fe^{3+}$) determined in non-irradiated AA and BA samples are presented in Table 1. The trace elements and metals in the AA sample, which mainly originate from soil dust (e.g., P, Al, Ca and Fe), non-ferrous metallurgical industrial activities (Cd, Cu and Zn) and fossil fuel combustion (Cr, Pb and V) (Pacyna and Pacyna, 2001; Mahowald et al., 2008), are significantly more abundant than in BA (by up to several times higher) with the exception of S, Ni and Sb (Table 1). The high abundances of trace metals in AA further suggest that the AA sample should contain high abundances of anthropogenic organic matter. The higher abundances of S, Ni and Sb in BA than in AA may be due to high emissions of S from intensive consumption of biofuels, particularly cow dung that contains higher S content (Reddy and Venkataraman, 2002b), while Ni and Sb are from some specific industrial activities in southern India. Although watersoluble $Fe^{2+}$ and $Fe^{3+}$ species are abundant in both AA and BA, their concentrations in BA are 30–50 % higher than in AA (Table 1). Furthermore, the fraction of water-soluble Fe ($Fe_{ws}$; sum of $Fe^{2+}$ and $Fe^{3+}$) in total particulate Fe ($Fe_{tp}$) is 2.77 % in AA, whereas it is 14.6 % in BA.

3.2 Molecular compositions of diacids, oxoacids and α-dicarboxyls

A homologous series of normal ($C_2$–$C_{12}$) and branched chain (iso-$C_4$–$C_6$) saturated α,ω-diacids were detected in both non-irradiated and irradiated AA and BA samples as well as aliphatic unsaturated diacids such as maleic (M), furmaric (F) and methylnalene (mM) acids and aromatic diacids such as phthalic (Ph), isophthalic (i-Ph) and terephthalic (t-Ph) acids. Diacids with an additional functional group, i.e., malic (hydroxysuccinic, hC$_4$), ketomalonic (kC$_3$) and 4-ketopimelic (kC$_7$) acids, were detected together with ω-oxoacids ($ωC_2$–$ωC_9$), pyruvic acid (Pyr) and α-dicarboxyls, i.e., glyoxal (Gly) and methylglyoxal (MeGly). $ωC_6$ will not be reported here due to the overlapping peak on the gas chromatograph.

Oxalic ($C_2$) acid was found to be the most abundant diacid in non-irradiated samples (accounting for 54 % of total diacids in AA and 53 % in BA), followed by Ph (10 %), $C_4$ (9 %), $C_3$ (8 %) and $C_6$ (4 %) in AA and by maleic ($C_3$) (9 %), $C_4$ (6 %) and t-Ph (6 %) acids in BA. Branched chain diacids were significantly lower than the corresponding normal structures in both samples. Glyoxylic ($ωC_2$) acid is the most abundant oxoacid, comprising 64 and 57 % of total oxoacids in AA and BA, respectively, followed by Pyr (13 %) and 4-oxobutanoic ($ωC_4$) acid (10 %) in AA and $ωC_4$ (18 %) and Pyr (13 %) in BA. MeGly is more abundant than Gly in AA, whereas their abundances are equivalent in BA.

3.3 Changes in concentrations of diacids and related compounds as a function of UV irradiation time

Changes in concentrations of individual and total diacids as a function of UV irradiation time in AA and BA are depicted in Fig. 3, while those of oxoacids and α-dicarboxyls as well as total oxoacids and α-dicarboxyls in Fig. 4. Concentrations of $C_2$ diacid were sharply decreased by a factor of 3–9 (from 553 to 61.7 ng m$^{-3}$ in AA and from 339 to 118 ng m$^{-3}$ in BA) within 6 h and 12 h of UV irradiation, respectively (Fig. 3a). Then the concentrations started to increase to maximize at 24 h (292 ng m$^{-3}$) in AA and 18 h (306 ng m$^{-3}$) in BA on further irradiation. They gradually decreased toward the end (120 h) of the experiment (Fig. 3a). Interestingly, $C_3$ diacid showed a temporal variation similar to $C_2$ in both AA and BA, except for a few points (Fig. 3b). Relative abundances of $C_2$ in total diacids gradually decreased from non-irradiated samples (54 % in AA and 53 % in BA) toward the end (120 h) of the experiment (3.2 % in AA and 9.2 % in BA, Fig. 5).

Concentrations of $ωC_2$, an immediate precursor of $C_2$ (Kawamura et al., 1996a; Warneck, 2003; Lim et al., 2005), increased with irradiation time up to 18 h in both AA and BA, except for two cases (3 and 6 h) of AA, and then gradually decreased until the end (120 h) of the experiment, except for one case (36 h) in AA (Fig. 4a). Pyr, Gly and MeGly,
which are the precursors of $\omega$C$_2$ acid, are all produced by the oxidation of VOCs of anthropogenic and biogenic origin (Warneck, 2003; Ervens et al., 2004b; Lim et al., 2005; Carlton et al., 2006). They also increased with irradiation time up to 18–24 h in both samples and then gradually decreased (except for MeGly in AA) until the end (120 h) of the experiment (Fig. 4g, i, j). However, the other precursor of C$_2$ diacid, kC$_3$ diacid (Kawamura et al., 1996a), showed a decrease with irradiation time throughout the experiment except for in a few cases (Fig. 3v), whereas hC$_4$, a precursor of C$_3$ diacid (Kawamura et al., 1996a), increased up to 18 h in BA and 24 h in AA, remained relatively high until 72 h and then gradually decreased until the end (120 h) of the experiment (Fig. 3u).

In contrast, concentrations of C$_4$ diacid showed a gradual increase with irradiation time up to 72 h in BA and 96 h in AA followed by a slight decrease in the AA and a sharp decrease in BA (Fig. 3c). Relative abundance of C$_4$ diacid in total diacids also increased from 8.9 % (non-irradiated) to 82 % (120 h) in AA and from 6.4 to 88 % in BA (Fig. 5). Similarly, C$_5$ diacid in AA (Fig. 3d) showed a gradual increase with irradiation up to 36 h and stayed almost constant until 96 h followed by a slight decrease. Similar trend was found in BA (Fig. 3d). Both C$_6$ and C$_7$ diacids showed an increase with irradiation up to 6–36 h and then a gradual decrease un-

Figure 3. Changes in concentrations of individual dicarboxylic acids and total diacids as a function of UV irradiation time in anthropogenic (AA) and biogenic aerosols (BA).
Figure 4. Changes in concentrations of individual oxoacids and α-dicarbonyls and total oxoacids and α-dicarbonyls as a function of UV irradiation time in AA and BA.

Figure 5. Changes in relative abundances of straight-chain diacids (C₂–C₁₀) to total diacids as a function of UV irradiation time in AA and BA.
of selected species in AA and BA were similar (Figs. 3–6), α-diacids, oxoacids and ω-dicarbonyls as well as relative abundances of individual diacids in total diacids and mass ratios of selected species in AA and BA were similar (Figs. 3–6), although significant differences are recognized between AA and BA samples during irradiation. Such similarities in the temporal variations of diacids and related polar compounds infer that their photochemical formation and degradation pathways in aqueous aerosols (Fig. 7) are almost same between anthropogenic and biogenic aerosols. However, there were significant differences in the rate of formation and/or degradation of diacids and related compounds between AA and BA, which might have been driven by the differences in the abundances of the diacids and related compounds as well as their precursor compounds in the original (non-irradiated) AA and BA samples. In fact, total diacids, oxoacids and α-dicarbonyls were higher in non-irradiated AA than in BA. On the contrary, OC that contains several precursor compounds (including fatty acids) of diacids and related polar compounds is higher in BA than in AA (Table 1).

3.4 Production and decomposition of short-chain diacids and related compounds

A sharp increase was observed in the concentrations of ωC2, ωC4, Pyr, Gly, and MeGly, but not ωC3, with irradiation up to 18–24 h following a gradual decrease (Fig. 4), demonstrating an enhanced photochemical production of short-chain (≤ C4) oxoacids and α-dicarbonyls during an early stage of photochemical processing. It is likely because ωC2, Pyr, Gly and MeGly are produced by photochemical oxidation of aliphatic olefins and aromatic hydrocarbons and ωC4 was produced from cyclic olefins and unsaturated fatty acids (Bandow et
Figure 7. Possible photochemical formation and/or degradation pathways of diacids, oxoacids and α-dicarbonyls in aqueous aerosols.

al., 1985; Hatakeyama et al., 1987; Kawamura et al., 1996a; Warneck, 2003; Lim et al., 2005), but αC₃ may not be significantly produced from any of these precursor compounds (Fig. 7). The increasing trends of mass ratios of C₂ to its precursor compounds, i.e., αC₂, Pyr, Gly and MeGly as well as C₃ (but not C₄) diacid (Kawamura et al., 1996a; Warneck, 2003; Ervens et al., 2004b; Lim et al., 2005; Carlton et al., 2007), were found for BA near 120 h (Fig. 6a–e and f). It is noteworthy that C₃/αC₇ ratios also showed a slight increase, although they are not clear in the later stages of experiment (Fig. 6g), suggesting a potential formation of C₃ diacid via αC₇ that is derived from unsaturated fatty acids and/or cyclic olefins. In addition, F/M ratios showed an increase with irradiation up to 48 h in AA and 18 h in BA followed by a gradual decrease until the end of experiment (Fig. 6i), indicating a significant photochemical transformation during an early stage of experiment and decomposition in a later stage.

Photochemical degradation of C₂ and C₃ diacids should have overwhelmed their photochemical production even in an early stage of the experiment except for a few cases (Fig. 3a, b). Diacids and other compounds containing a carbonyl group can form stable carboxylate salts with amines upon photochemical oxidation. However, based on laboratory studies, C₂ and C₃ diacids have been reported to decompose in aqueous phase in the presence of Fe³⁺ (and C₂ diacid even in the presence of Fe²⁺) under UV irradiation at 254 nm as well as at a solar spectrum (> 300 nm) (Zuo and Hoigne, 1994; Z. H. Wang et al., 2010; Pavuluri and Kawamura, 2012), but C₂ diacid (and may be C₃ diacid) is relatively stable in the absence of Fe species (Pavuluri and Kawamura, 2012). It is well documented that both C₂ and C₃ diacids have the strongest chelating capacity with Fe³⁺ among all diacids and tend to form mono, di- and trioxalato (equilibrium constant log₁₀(b) = 9.4, 16.2 and 20.4, respectively) and malonato (equilibrium constant log₁₀(b) = 7.5, 13.3 and 16.9, respectively) complexes by acting as ligands in aqueous phase, which exhibit a strong light-absorbing ability (Z. H. Wang et al., 2010). Although the equilibrium
constant of Fe$^{3+}$-malonato complex is slightly lower than that of Fe$^{3+}$-oxalato, both diacids photolyze upon the absorption of UV light to result in Fe$^{2+}$ and CO$_2$ (Zuo and Hoigne, 1994; Z. H. Wang et al., 2010).

We found that non-irradiated AA and BA samples contain significant amounts of water-soluble Fe$^{2+}$ and Fe$^{3+}$ species (Table 1). Because a high abundance of particulate Fe is present in both AA and BA (Table 1), the concentrations of water-soluble Fe$^{2+}$ and Fe$^{3+}$ species in both AA and BA samples may increase upon UV irradiation; the water-insoluble Fe can be transformed into water-soluble forms by photochemical processing of mineral aerosols (Solmon et al., 2009; Srinivas et al., 2012). However, we did not measure the concentrations of Fe$^{2+}$ and Fe$^{3+}$ species in the irradiated samples. In fact, the mass ratio of C$_2$ diacid to Fe$^{3+}$ is 15 : 1 in non-irradiated AA and 7 : 1 in BA, both of which are close to the ratio (10 : 1) used in laboratory experiments conducted by Pavuluri and Kawamura (2012) for Fe-catalyzed photolysis of C$_2$ diacid in aqueous phase, in which the photolysis of C$_2$ is very fast ($k = 206 \text{ mol}^{-1} \text{s}^{-1}$) and 99% of the C$_2$ is degraded in 0.5 h. Therefore, available water-soluble Fe$^{3+}$ and Fe$^{2+}$ in AA and BA should be enough to promote the catalytic photochemical degradation of C$_2$ (and C$_3$) upon UV irradiation (Fig. 7) and thus the degradation rate of C$_2$ (and C$_3$) should have increased with the prolonged experiment due to enhancement in Fe$^{3+}$ and Fe$^{2+}$ levels in the given sample.

The concentration of C$_2$ diacid in AA decreased by 30% in 1.5 h and continued to decline by 90% until 12 h (Fig. 3a). However, the experiment of BA showed that the concentration of C$_2$ decreased by 47 and 51% in 0.5 and 1.5 h, respectively, and then gradually declined. The concentrations of C$_3$ also showed similar trends with C$_2$ (Fig. 3b). Although C$_2$ and C$_3$ diacids decreased sharply during early stages of experiment, they decreased gradually in the later stages despite possibly enhanced levels of water-soluble Fe$^{2+}$ and Fe$^{3+}$ species. These trends imply that photolysis of C$_2$ and C$_3$ diacids is highly significant in the presence of water-soluble Fe$^{3+}$ (and Fe$^{2+}$) (Fig. 7). However, the formation of both C$_2$ and C$_3$ diacids is also intensive with the photochemical processing of their precursor compounds in AA and BA. However, the net rate of production or degradation of C$_2$ and C$_3$ diacids in each experiment (Fig. 3a, b) should depend on the abundances of water-soluble Fe$^{2+}$ and Fe$^{3+}$ species and their precursors in AA and BA.

We found an increase in the mass ratios of MeGly to Gly with irradiation toward the end of the experiment, except for an early stage of the experiment (up to 6 h) in AA, whereas in BA they remained relatively constant up to 36 h and then increased gradually up to 72 h, followed by a rapid decrease (Fig. 6n). As noted earlier, concentrations of Gly and MeGly increased with experiment up to 18–24 h in both AA and BA. Thereafter, Gly decreased toward the end of experiment in both AA and BA, whereas MeGly remained relatively constant in the AA but decreased in BA (Fig. 4i, j). Such differences should be caused by the difference in their production rates depending on the concentrations of potential precursors and their oxidation products in AA and BA, i.e., benzene and glycolaldehyde for Gly and acetone and higher alkanes (> C$_3$) and alkenes (> C$_2$) for MeGly (Fujita et al., 2008), rather than the reaction rates of the Gly $\text{A} \times 10^9 \text{ M}^{-1} \text{s}^{-1}$ and MeGly (6.44 × 10$^8$ M$^{-1}$s$^{-1}$) with HO* in aqueous phase (Tan et al., 2012). Therefore, the higher abundance of MeGly than Gly in AA can be attributed to its enhanced production than the later species during photochemical processing of aqueous aerosols derived from anthropogenic sources. Furthermore, the oligomerization of Gly and MeGly (Tan et al., 2009, 2012; Lim et al., 2010) might have also played an important role on the changes in their concentrations with irradiation time; however, we did not focus on the measurements of oligomers here because of the analytical limitations.

### 3.5 Possible photochemical pathways of long-chain diacids and oxoacids

Enhanced concentrations of normal and branched C$_4$–C$_7$ diacids during an early stage (18–36 h) (Fig. 3c–f), despite degradation of C$_2$ and C$_3$ and longer-chain (> C$_7$) diacids (Fig. 3a, b, g–k), may be caused by photochemical oxidation of the first-generation products derived from the oxidation of anthropogenic and/or biogenic VOCs (e.g., cycloalkanes, monoterpenes and sesquiterpenes) and unsaturated fatty acids (Kalberer et al., 2000; Gao et al., 2004) (Fig. 7). In addition, the photochemical oxidation of the polymers of polyunsaturated fatty acids, if available, can produce significant long-chain (≥ C$_4$) diacids (Harvey et al., 1983), a subject of future research. In fact, polyunsaturated fatty acids (e.g., linolenic acid, C$_{18:3}$) can undergo free radical oxidative cross-linking in the air and produce high-molecular-weight organic compounds (e.g., fulvic acid) (Wheeler, 1972; Harvey et al., 1983). Harvey et al. (1983) found a series of C$_4$–C$_9$ diacids by oxidizing the marine fulvic acid in a laboratory study. The chelating capability of succinate (equilibrium constant log M$^{-1}$ = 7.5; Z. H. Wang et al., 2010) and other long-chain diacids with Fe$^{3+}$ is weak and, hence, their photolysis is insignificant. However, they should be further oxidized to result in lower diacids (Kawamura et al., 1996a; Matsunaga et al., 1999). The degradation of these diacids should be increased with increasing chain length because the oxidation rate of C$_4$ to C$_9$ diacids is increased with increasing carbon number (Yang et al., 2008a).

The relatively constant levels of C$_5$, iC$_4$, and iC$_5$ during 36 h and 72–96 h (Fig. 3d, i, m) may be due to the balance between photochemical production and degradation. The increases in the concentrations of C$_4$ with a prolonged irradiation up to 72 h in BA and 96 h in AA further demonstrate its formation from higher diacids and other precursors in aqueous aerosols (Kawamura and Sakaguchi, 1999; Charbonnet et al., 2012) (Fig. 7). In fact, total diacids stayed relatively constant from 24 h to 72–96 h (Fig. 3x). In addi-
tion, mass ratios of C₄ to C₅–C₇ showed a gradual increase throughout the experiment (until 120 h) in both AA and BA (Fig. 6k–m). These results support a photochemical breakdown of longer-chain (≥ C₅) diacids resulting in C₄ (Matsumaga et al., 1999; Yang et al., 2008a; Charbouillot et al., 2012). Yang et al. (2008a) reported that the production of C₄ diacid is predominant, followed by C₅ diacid during a laboratory photochemical oxidation of C₆–C₉ diacids.

In addition, ωC₈ acid, which can be produced by the oxidation of cyclic olefins and unsaturated fatty acids (Kawamura and Sakaguchi, 1999; Gao et al., 2004), showed a gradual increase (Fig. 4e) similar to that of C₄ diacid (Fig. 3c) in both AA and BA, suggesting a significant photochemical production of C₄ via ωC₈ until the consumption of the precursor compounds derived from anthropogenic and biogenic VOCs and biogenic unsaturated fatty acids (Klaber et al., 2000; Gao et al., 2004). In fact, ratios of C₄ to C₅–C₇ were 10 times higher in BA than in AA whereas those of C₄/ωC₈ were similar in both the BA and AA (Fig. 6j). However, their temporal profiles with irradiation time are similar in both AA and BA. These results suggest that the formation of C₄ and ωC₈ is much higher in biogenic aerosols than in anthropogenic aerosols compared to C₅–C₇ diacids, but their formation/degradation processes may be similar irrespective of the origin of precursors. However, it is not clear from this study whether C₄ is mainly derived (via ωC₈) from cyclic olefins or unsaturated fatty acids (Fig. 7).

It is well established that long-chain (C₈–C₁₂) diacids are formed by photochemical oxidation of unsaturated fatty acids (e.g., oleic acid) (Kawamura and Gagosian, 1987; Matsumaga et al., 1999) (Fig. 7). However, unsaturated fatty acids were not abundant (e.g., oleic acid was 0.89 ng m⁻³ in AA and below detection limit in BA) in non-irradiated samples (Fu et al., 2010). Hence, photochemical formation of long-chain diacids from the oxidation of unsaturated fatty acids should be less important during the experiment, although chemical forms of polymerized and/or partially oxidized unsaturated fatty acids may be abundant in the aerosols. The rate constant of photooxidation of diacids increases with an increase in carbon number of individual diacids (≥ C₄) (Yang et al., 2008a). Hence, photochemical breakdown of C₈–C₁₂ diacids to lower diacids (Matsumaga et al., 1999; Yang et al., 2008a) should be very likely (Fig. 7). The gradual decreases of aliphatic unsaturated diacids, aromatic diacids and o xoacids, except for ωC₈, with irradiation are likely caused by the photochemical degradation (Fig. 7).

3.6 Atmospheric implications

As discussed above, this study reveals that photochemical degradation of C₂ and C₃ (due to Fe-catalyzed photolysis) in aqueous aerosols overwhelmed their production whereas C₄ diacid showed photochemical formation. These results are consistent with the recent atmospheric observations: a significant reduction in C₂ diacid concentration and an inverse relationship between the C₂ and Fe in cloud water (Sorooshian et al., 2013) and the replacement of the predominance of C₂ by C₄ in the Arctic aerosols (Kawamura et al., 2010, 2012).

It was also reported that C₄ and C₅ diacids are most abundant among C₃–C₈ diacids determined during the photochemical oxidation of C₆–C₉ diacids in a laboratory experiment (Yang et al., 2008a).

On the contrary, the enhanced degradation of C₂ and C₃ and the formation of C₄ diacid upon prolonged irradiation are not consistent with previous laboratory, observation and model studies on photochemical production and degradation of diacids and related compounds in aqueous phase (e.g., cloud processing) (Kawamura et al., 1996a; Kawamura and Sakaguchi, 1999; Warneck, 2003; Ervens et al., 2004b; Lim et al., 2005; Carlton et al., 2007; Charbouillot et al., 2012). In fact, previous studies did not consider Fe-catalyzed photolysis of C₂ diacid, which is significant at least in Fe-rich atmospheric waters. However, the formation processes and potential precursor compounds of C₄ diacid have not been fully explored yet. Moreover, previous laboratory experiments on aqueous solutions of specific species did not consider the mixing state of organic and inorganic constituents in atmospheric aerosols (Ervens et al., 2011), although simplified experiments sometimes provide useful information on mechanisms.

Generally, it has been considered that the anthropogenic contributions of α-dicarbonyls to organic aerosols are minor: 8 % for Gly and 5 % for MeGly (Fu et al., 2008). To the best of our knowledge, their production in atmospheric waters has not been well recognized yet. Our laboratory experiments indicate that the photochemical production of Gly and MeGly is significant in aqueous aerosols. The production of MeGly is more pronounced compared to Gly with prolonged photochemical processing of aqueous anthropogenic aerosols. Finally, our findings based on the batch laboratory experiment emphasize the importance of the photolysis of C₂ and C₃ diacids and photochemical production of C₄ diacid and α-dicarboxyls in aqueous aerosols to reconcile the current atmospheric model(s) such as the cloud parcel model (Ervens et al., 2004a) and to better understand the secondary organic aerosol budget and its climatic impacts.

4 Summary and conclusions

In this study, we conducted batch UV irradiation experiments on AA and BA aerosol samples collected from Chennai, India, in the presence of moisture for the reaction time of 0.5 to 120 h. The irradiated samples were analyzed for molecular compositions of diacids, o xoacids and α-dicarboxyls. Concentrations of C₂ and C₃ and C₈–C₁₂ diacids decreased with an increase in reaction time of 12–24 h. In contrast, C₄ diacid (and C₅–C₇) showed a significant increase with reaction time up to 72 h in BA and 96 h in AA. O xoacids and α-dicarboxyls showed a significant increase during an early stage of irradi-

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