Long-term (2001–2013) observations of water-soluble dicarboxylic acids and related compounds over the western North Pacific: trends, seasonality and source apportionment

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To better understand the impact of East Asian pollutants on the molecular composition of marine organic aerosols, we conducted long-term (2001–2013) observations of water-soluble dicarboxylic acids and related compounds in total suspended particulate samples collected at Chichijima Island in the western North Pacific (WNP). Seasonal variations of all the diacids and related compounds showed maxima in winter and spring and minima in summer, except for azelaic acid (C9), which maximized in summer to autumn. The overall annual concentrations of the total diacids, ω-oxoacids and α-dicarbonyls showed an increase during 2001–2013. We found a significant (p < 0.05) decadal increase in the inter-annual trends of pyruvic and glyoxylic (p > 0.05) acids, and methylglyoxal (MeGly). In contrast, phthalic acid (p < 0.05) and glyoxal (Gly) showed a decrease in their trends. We also found a significant decrease in the trend of the Gly/MeGly mass ratios. These results demonstrate that the enhanced concentrations of diacids over the WNP are majorly attributed to the aqueous-phase photooxidation of biogenic volatile organic compounds from East Asia followed by long-range atmospheric transport. Further, positive matrix factorization analysis showed a biogenic photochemical contribution (42%) was the dominant source of oxalic acid in the WNP.

Low molecular weight dicarboxylic acids (diacids) are ubiquitous and important constituents of atmospheric organic aerosols. Due to the low vapor pressures (approximately less than 10^{-7} Pa), these compounds are almost exclusively partitioned into a particulate phase and contribute significantly to the water-soluble fraction. Thus, particles enriched with diacids can act as cloud condensation nuclei and have an impact on the earth’s radiative forcing. Diacids also play an important role in many biological processes in the ocean. Despite the great importance, they continuously represent a challenge to atmospheric scientists due to their complexity, although the first unambiguous observations of diacids in the atmosphere were made three decades ago.

Diacids are the late photooxidation products of hydrocarbons and other organics. Therefore, photochemical chain reactions are the most important sources of diacids. Aqueous-phase oxidation and cloud processing of aerosols are potentially important sources of organic aerosols and could explain the high abundances of oxalic acid in the atmosphere. High levels of diacids were observed in motor exhausts and ambient aerosols influenced by forest fires, indicating important primary sources of diacids. However, there is still a knowledge gap on the relative contribution of primary and secondary sources of diacids in the atmosphere. On the other hand, iron-carboxylate complexes and their photochemical degradation in the water mediated atmosphere may be an important sink of diacids in addition to wet and dry deposition.

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A large fraction of secondary organic aerosol (SOA) is derived from the oxidation of isoprene, monoterpenes and sesquiterpenes of biogenic origin. Global aerosol modelling studies suggest an increase in isoprene emissions of 22 to 55% by 2100 in response to the temperature increase following a global warming of 33–35. Recent modelling studies simulate an increased trend of isoprene emissions with 0.16% per year over Asia and this trend is more pronounced over China with 0.52% per year during 1979 to 2012. All these models, however, have significant uncertainties in the projection of SOA formation due to a series of fundamental assumptions and lack of long-term observational studies. The models assume that particles are liquids and are exposed to changes in the surrounding atmosphere by rapid condensation or evaporation and by in-particle mixing during long-range atmospheric transport.

East Asia is one of the most populated regions on the globe with a population density of 133 inhabitants per square kilometer, which is about three times higher than the world population density (45 per km²). Increasing levels of air pollutants in East Asia and their outflows to the Pacific have received a significant attention due to the potential impacts on regional and global climate and atmospheric circulations. However, the effects of East Asian pollutants and their outflows over the western North Pacific (WNP) are still unclear, especially with regard to the sources and formation of SOA during long-range transport.

In this study, we carried out long-term (2001–2013) measurements of diacids and related compounds in total suspended particulate (TSP) aerosols collected at Chichijima Island in the WNP. Chichijima is a remote island 2000 km away from East Asia and exists in the outflow region of Asian dusts and pollutants from East Asia, especially China. Here, we discuss the inter-annual trends and seasonality of diacids over the WNP. We also discuss the sources and formation pathways of the diacids using a positive matrix factorization (PMF) model.

Results and Discussion

Air mass origin and meteorological parameters. Figure 1(a–d) presents daily 7-day isentropic air mass back trajectories at an altitude of 500 m above the ground level using the HYSPLIT model for different seasons during the year 2013 over the WNP as an example. Air mass transport from East Asia to the sampling site in the Pacific is stronger during winter (December to February) and spring (March to May) than during summer and autumn to deliver continental air masses via long-range atmospheric transport. The continental air mass transport is almost absent in summer (June to August). Air masses mostly come from the central Pacific carrying pristine air masses to the observation site in summer, whereas in autumn (September to November) the air mass pathway shifts from southeasterly to northwesterly towards winter. More detailed information of air mass transport over the WNP is described elsewhere.

The meteorological parameters such as air temperature (°C), wind speed (m s⁻¹), solar radiation (MJ m⁻²), precipitation (mm) and cloud cover are downloaded from the Japan Meteorological Agency (JMA) for Chichijima...
Island during 2001–2013 (Figure S1). We found a clear seasonal variation in air temperature, solar radiation and precipitation with summer maxima and winter minima. Wind speed and cloud cover were higher in spring and lower in summer. However, no significant decadal trends were observed in any meteorological parameters throughout the sampling period, therefore, we used meteorological parameters in this study to better understand the seasonal variations of organic compounds but not to discuss the trends.

**Molecular distributions.** In order to better understand the sources, formation pathways, and physicochemical properties of organic aerosols, we studied the molecular distributions of the diacids and related compounds at Chichijima Island between 2001 and 2013. Molecular distributions are shown in Figure S2. Throughout the observation period, we found the predominance of oxalic acid (C2) followed by malonic (C3) and/or succinic (C4) acids. This molecular distribution is consistent with our previous study for 1990–1993 at the same observation site44 and other East Asian sites such as Okinawa Island49, 50, the Gosoan site, Jeju Island in South Korea51, Mt. Tai in North China52, urban sites in China53, 54, as well as different sites in the world including Tanzania, East Africa55, remote western European continental sites56, west-east transect in the European atmosphere57, and Los Angeles58.

Glyoxylic acid (ω-C2) was the dominant species among all ω-dicarboxylic acids and the fourth most abundant species detected, whereas methylglyoxal (MeGly) was more abundant than glyoxal (Gly) in the WNP aerosols. Throughout the observation period, we found the molecular distributions of the diacids and related compounds at Chichijima as C2 > C3 > C4 > ω-C2 > MeGly > Ph > ω-C7 > C5 > ω-C9 > Gly > C6 > ω-C8 > C10. However, seasonal molecular distributions provided a different picture depending on the source strengths and formation mechanisms.

**Trends.** Table 1 summarizes the regression statistics such as range, mean ± SD, and the trend (slope) for water-soluble diacids, ω-dicarboxylic acids, pyruvic acid, α,ω-dicarbonyls, and their diagnostic ratios during the period of 2001 to 2013. Figure 2 presents the temporal trends of the major diacids and related compounds and diagnostic mass ratios during the whole study period.

C2, C3, and C4 acids are the end or near-end products of photochemical reaction chains of hydrocarbons and biogenic unsaturated fatty acids, accounting for approximately 74%, 11%, and 6% of the total diacids, respectively, during the 13-year study period at Chichijima. From Table 1, it is obvious that C4 shows an increasing trend (+0.003% yr−1), whereas those of C2–C3 diacids (particularly, C6, C7, C8, C9, and C12) show significant (p < 0.05) decreasing trends during the study period. These results suggest a photochemical conversion of higher to lower molecular weight diacids during the long-range atmospheric transport over the WNP. This point is further supported by significant increasing trends of diagnostic mass ratios of diacids, as discussed below.

It has been suggested that maleic acid (cis configuration) (M), a photo-oxidation product of aromatic hydrocarbons such as benzene and toluene69, may be isomerized to fumaric acid (trans configuration) (F) under a high solar radiation in the atmosphere61. Therefore, the F/M ratio may be a good indicator of photochemical processing. In our study, we found a significant (p < 0.05) increasing trend (+0.016% yr−1) of F/M ratios during 13 years of the study period (Fig. 2g), indicating an intensive photochemical aging or increased oxidant levels over the WNP. Further, it has been suggested that C2 and C3 are likely produced in the marine atmosphere by the photooxidation of C4 through intermediates such as hydroxysuccinic or malic acid (hC4) and ketomalonic acid (kC3)56, 59. In this study, we found that C2/C3 ratios showed an increasing trend, although it is not significant (p > 0.05). However, C3/C2 and C4/C2 ratios show increases (+0.004% yr−1 and +0.005% yr−1) during the 13 year study period. This observation suggests that C2 could be largely produced by the photochemical degradation of C3 and C4 diacids. A significant (p < 0.05) increase (+0.002% yr−1) of C4/C2 (C2–C12) further supports the photochemical aging of diacids over the remote marine aerosols (Table 1). All these results suggest that the production of diacids over the WNP may be closely linked with an increased photochemical oxidation of biogenic and anthropogenic precursors that are delivered from the Asian continent by long-range atmospheric transport.

Previous studies suggested that oxidation of aromatic hydrocarbons such as naphthalene and o-xylene, which originate from an incomplete combustion of fossil fuel18, is one of the major sources of phthalic acid (Ph) in the atmosphere61, 62. High abundances of aromatic hydrocarbons are reported over China during winter63, 64. Similarly, adipic acid (C6) is probably produced through the oxidation of cyclohexene by ozone in the atmosphere and has been proposed as an anthropogenic tracer65. It is also well documented that glyoxal (Gly) is largely produced in the atmosphere by the oxidation of many aromatic anthropogenic hydrocarbons65–67, although it has small contribution from biogenic and marine origin68–70. In this context, we found a decreasing trend in the concentrations of Ph (p < 0.05; −0.007% yr−1), C6 (p > 0.05; −0.003% yr−1) and Gly (p < 0.05; −0.006% yr−1) (Fig. 2b, e, and Table 1). These results suggest that combustion (fossil fuel) derived aerosols have declined (or constant) over the WNP during 2001–2013. This point is further supported by the study of Boreddy, et al.71, who reported that declined concentrations of elemental carbon (EC) over WNP occurred during 2001–2012 over the WNP. However, it should be noted that although Gly concentrations are decreased, its processing to C2 has not decreased over the WNP, as evidenced by the increasing trend (+0.002% yr−1; p > 0.05) of C2/Gly ratios (Table 1).

On the other hand, a modeling study by Stavrakou, et al.72 observed a continuous increase of isoprene emissions over Asia during 1979–2012. They found a strong correlation (r > 0.90) between isoprene emissions and above-canopy solar radiation, suggesting that enhanced solar radiation intensifies isoprene emissions from terrestrial higher plants over Asia (particularly in China). Similarly, Zhang, et al.72 have recently reported an increase of biogenic isoprene emissions in northern China during 1982–2010 using the biogenic emission model. It has also been documented that 79% of MeGly may come from biogenic isoprene emissions globally, as inferred from modeling studies73. Further, pyruvic (Pyr) and ω-C2 have been suggested as in-cloud oxidation products of isoprene, which are subsequently oxidized to C2. Therefore, it may be possible that biogenic isoprene derived volatile organic precursors (e.g., MeGly) over East Asia/China are taken up by aqueous-phase aerosol particles in the atmosphere and transported to the WNP.
### Organic compounds

|              | Range (mean ± SD) | Slope (m, diacid year⁻¹) | Uncertainty (σm) | Trend (% year⁻¹) |
|--------------|-------------------|--------------------------|------------------|------------------|

#### Dicarboxylic acids (ng m⁻²)

**Normal chain saturated dicarboxylic acids**

- Oxalic, C₂: 2.21–514 (73.9 ± 66.8) ± 0.0032 ± 0.0020 + 0.004
- Malonic, C₃: 0.28–55.6 (11.5 ± 9.4) ± 0.00005 ± 0.0002 + 0.0004
- Succinic, C₄: 0.05–52.4 (6.12 ± 6.41) ± 0.00002 ± 0.0002 + 0.0003
- Glutaric, C₅: 0–7.48 (1.11 ± 1.19) ± 0.00003 ± 0.00003 + 0.0027
- Adipic, C₆: 0.01–5.08 (0.59 ± 0.60) ± 0.00002 ± 0.000017 + 0.0033
- Pimelic, C₇: 0–1.34 (0.15 ± 0.16) ± 0.000014* ± 0.000047 + 0.009
- Sebacic, C₈: 0–1.24 (0.13 ± 0.14) ± 0.00003* ± 0.000038 - 0.023
- Azelic, C₉: 0.01–2.50 (0.53 ± 0.34) ± 0.00003* ± 0.0001 ± 0.005
- Decanedioic, C₁₀: 0–0.95 (0.06 ± 0.08) ± 0.000003 ± 0.000023 - 0.005
- Undecanedioic, C₁₁: 0–8.03 (0.08 ± 0.34) ± 0.000002 ± 0.000098 - 0.002
- Dodecanedioic, C₁₂: 0–0.34 (0.01 ± 0.04) ± 0.000004* ± 0.00001 ± 0.04

**Branched chain saturated dicarboxylic acids**

- Methylmalonic, iC₄: 0–1.45 (0.31 ± 0.25) ± 0.000009 ± 0.000074 - 0.0029
- Methylsuccinic, iC₅: 0–3.13 (0.52 ± 0.48) ± 0.000012 ± 0.00014 ± 0.0019
- Methylglutaric, iC₆: 0–0.96 (0.08 ± 0.09) ± 0.000006* ± 0.00002 ± 0.0075

**Multi functional saturated dicarboxylic acids**

- Hydroxy succinic, hC₄: 0–15.2 (0.26 ± 1.12) ± 0.000111* ± 0.00032 - 0.038
- Ketomalonic, kC₃: 0–5.40 (0.38 ± 0.54) ± 0.00002 ± 0.00015 + 0.0052
- Ketopimelic, kC₅: 0–3.67 (0.50 ± 0.57) ± 0.000007 ± 0.00017 - 0.0014

**Unsaturated aliphatic dicarboxylic acids**

- Maleic, M: 0–2 (0.43 ± 0.40) ± 0.00005* ± 0.00001 ± 0.011
- Fumaric, F: 0–2.27 (0.41 ± 0.32) ± 0.00002* ± 0.000092 ± 0.005
- Methylmalic, mM: 0–7.66 (0.22 ± 0.44) ± 0.000007* ± 0.000012 ± 0.031

**Aromatic dicarboxylic acids**

- Phthalic, Ph: 0.01–12.8 (1.31 ± 1.49) ± 0.000009 ± 0.00043 - 0.0068
- Isophthalic, iP: 0–11.1 (0.14 ± 0.49) ± 0.00001 ± 0.00014 + 0.0071
- Terephthalic, tPh: 0–7.18 (0.39 ± 0.52) ± 0.00002 ± 0.00015 + 0.0051

**Total diacids**

2.93–555 (99.2 ± 86.4) ± 0.003 ± 0.0025 + 0.003

**ω-Oxocarboxylic acids (ng m⁻²)**

- Glyoxylic, ωC₂: 0.09–29 (4.91 ± 5.13) ± 0.00007 ± 0.00015 + 0.001
- 3-Oxopropanoic, ωC₃: 0–22.4 (0.26 ± 0.94) ± 0.00006* ± 0.00027 + 0.023
- 4-Oxobutanoic, ωC₄: 0–5.56 (0.39 ± 0.47) ± 0.00004* ± 0.00013 + 0.010
- 5-Oxopentanoic, ωC₅: 0–0.65 (0.09 ± 0.08) ± 0.00001* ± 0.000024 + 0.01
- 7-Oxoheptanoic, ωC₇: 0.02–11.6 (1.13 ± 1.27) ± 0.000006 ± 0.00037 + 0.0005
- 8-Oxooctanoic, ωC₈: 0–6.32 (0.72 ± 0.89) ± 0.00002 ± 0.00026 + 0.0027
- 9-Oxononanoic, ωC₉: 0–5.72 (0.59 ± 0.68) ± 0.00006* ± 0.00002 ± 0.010
- Total ω-oxoacids: 0.21–52.7 (8.10 ± 8.20) ± 0.0003 ± 0.00024 + 0.0037

**Ketosacids (ng m⁻²)**

- Pyruvic, Pyr: 0–6.16 (0.79 ± 0.90) ± 0.00002* ± 0.000025 ± 0.0025
- α-Dicarboxylic acids (ng m⁻²)

- Glyoxal, Gly: 0–4.69 (0.80 ± 0.78) ± 0.00005 ± 0.000036 - 0.006
- Methylglyoxal, MeGly: 0–23.9 (1.73 ± 2.28) ± 0.00014* ± 0.00067 + 0.008
- Total α-dicarboxylic acids: 0.04–26.4 (2.34 ± 2.74) ± 0.0003* ± 0.0008 + 0.012

**Ratios**

- F/M: 0–20.8 (1.83 ± 2.47) ± 0.0003* ± 0.000072 ± 0.016
- C₂/C₃: 1.46–33.1 (6.55 ± 2.86) ± 0.00034* ± 0.00083 ± 0.0045
- C₄/C₅: 0.80–154 (17.0 ± 12.3) ± 0.001* ± 0.00036 ± 0.0005
- C₆/C₇: 0.15–27.7 (2.76 ± 1.93) ± 0.00002 ± 0.000057 ± 0.0007
- Ph/C₅H: 0.01–77.0 (3.47 ± 5.10) ± 0.000006 ± 0.00015 ± 0.0002
- C₃竞争优势(C₃竞争优势): 0.38–0.94 (0.77 ± 0.06) ± 0.000008* ± 0.000017 ± 0.0010
- C₄/C₅: 5.48–131 (22.2 ± 14.7) ± 0.0013* ± 0.0004 + 0.0058
- C₆/C₇: 20.3–4050 (135 ± 332) ± 0.0022 ± 0.01 + 0.0022
- C₇/C₈: 0–758 (90.8 ± 86.3) ± 0.0067 ± 0.0025 + 0.007
- Gly/MeGly: 0–6.71 (0.98 ± 0.81) ± 0.0008* ± 0.000037 ± 0.008

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**Table 1.** Regression statistics (range, mean ± SD, and slope) of water-soluble dicarboxylic acids and related compounds (n = 607) in remote marine TSP aerosols collected at Chichijima Island during the period of 2001 to 2013. The symbol, *, indicates the trends are significant at a 95% (p < 0.05) confidence level.
In this study, we found a significant increasing trend (p < 0.05) in the concentrations of MeGly (+0.008% yr⁻¹), Pyr (+0.025% yr⁻¹) (Fig. 2d and f) and C₂/MeGly ratios (+0.007% yr⁻¹), suggesting the formation of C₂ from the oxidation of MeGly and Pyr via an intermediate compound, i.e., ωC₂ (MeGly → Pyr → acetic acid → ωC₂ → C₂) in the aqueous-phase. These results suggest that enhanced concentrations of the diacids are probably caused by an increase of biogenic isoprene-derived precursors (i.e., MeGly and Pyr), followed by the subsequent photochemical oxidation during long-range atmospheric transport over the WNP. A concurrent negative trend in Gly/MeGly (−0.008% yr⁻¹; p < 0.05) (Fig. 2h) also suggests an increase of biogenic precursors over the WNP. This point is further supported by our previous study of Boreddy and Kawamura, which reported that a significant increase (p < 0.05) in the concentrations of methanesulfonic acid (MSA; a tracer for biogenic sources) occurred during 2001–2012 over the WNP.

The observed trends of diacids may not only reflected by changes in emissions/air masses, but also influenced by changes in oxidant levels. To better understand the variations in the trends of oxidant levels over the WNP, we downloaded the monthly mean levels of total columnar ozone (DU) and daily tropospheric columnar NO₂ (cm⁻²) for the periods from 2002 to 2013 and from 2005 to 2013, respectively, from the NASA website (https://giovanni.gsfc.nasa.gov/) (Figure S3). From Figure S3, it is clear that both oxidant levels (O₃ and NO₂) showed significant (p < 0.05) increase in those trends (+0.0004% yr⁻¹ and +0.0008% yr⁻¹, respectively) during the study period, indicating that increases in oxidation processes over the WNP lead to increases in the formation of diacids during long-range atmospheric transport.

Figure 2. Trends in (a–f) temporal variations of water-soluble dicarboxylic acids (ng m⁻³) and related compounds (ng m⁻³) and (g,h) F/M and Gly/MeGly mass ratios in TSP aerosols collected at Chichijima Island during 2001 to 2013. Linear regression trends are given inset and applied over the whole observation period. The symbol, *, indicates the trends are significant at a 95% (p < 0.05) confidence level. The solid red line represents the trend line.
Based on the above results, we conclude that the enhanced concentrations of diacids over the WNP may be caused due to the increased oxidations of biogenic precursor compounds during long-range atmospheric transport. In contrast, anthropogenic precursors (e.g., Gly) have decreased (or constant) during the study period. It should be noted that all these trends are explained for the whole period (2001–2013); however, seasonal trends may give different results over the WNP because the formations of diacids are very sensitive to the sources of an air mass and the meteorological parameters as reported in Tables S4–S7.

As seen from Tables S4–S7, it is noteworthy that concentrations of total diacids in all seasons showed increases in their trends, except for summer, which showed a decreasing trend (−0.002% yr⁻¹; p > 0.05) (Table S6). The declined concentrations of diacids and related compounds in summer are probably due to the pristine air masses, which suggest the negligible local anthropogenic emissions as well as long-range continental outflow over the WNP. It is also seen from Tables S4–S7 that the trend in concentrations of C₉ showed a decrease in all seasons, except for summer. In summer, the trend of C₉ showed an increase (+0.001% yr⁻¹) during 2001–2013, indicating an importance of oxidation of biogenic unsaturated fatty acids over the WNP, particularly in summer.

**Monthly and seasonal variations.** Figure 3 presents box and whisker plots of monthly variations of diacids, ω-oxocarboxylic acids, pyruvic acid, α-dicarbonyls, and diagnostic mass ratios at Chichijima Island for the period of 2001 to 2013. Almost all the organic compounds showed clear monthly/seasonal variations with higher
Changes in emission strength and formations.  Gradual increases of total diacids, \( \omega \)-oxoacids and \( \alpha \)-dicarbonyls during late autumn to early spring are attributable to the combined effect of anthropogenic/biogenic volatile organic compounds (VOCs) emitted from East Asia followed by subsequent oxidation during long-range atmospheric transport. In our previous study, higher concentrations of non-sea salt sulfate (\( \text{ss-SO}_4^{2-} \)), nitrate (\( \text{NO}_3^- \)), non-sea salt calcium (\( \text{ss-Ca}^{2+} \)) and MSA\(^-\) were found in winter/spring over the same site under continental influence during the study period\(^{48}\). Being consistent with monthly variations of inorganic ion concentrations, \( \text{C}_2 \) to \( \text{C}_4 \) diacids show similar trends with a gradual increase from late autumn to early spring and decrease towards summer months. The seasonal variations of \( \text{C}_6 \), \( \text{Ph} \), and Gly are characterized by winter maxima and summer minima (Fig. 3h,j and m). Similarly, concentrations of Pyr and MgGly maximized in winter/spring and minimized in summer (Fig. 3l,n). These results demonstrate that, during late autumn to early spring, East Asian emissions of organic acids and their precursors, followed by long-range atmospheric transport, are important factors in controlling the distributions of diacids and related compounds in the WNP. High speed westerly winds also play an important role in causing the highest concentrations of diacids in spring over the WNP (Figure S1b).

On the other hand, lower concentrations of diacids and related compounds in summer suggest a minor contribution either from local emissions over the sampling site or marine emission of diacids and their precursors in the WNP. Mochida, et al.\(^{44}\) documented that local anthropogenic emissions for diacids are insignificant at Chichijima, based on the lower concentration ratios of benzo/[a]pyrene (BaP) to (\( \text{C}_2\text{--C}_11 \)). Therefore, it is reasonable to believe that the observed concentrations of diacids and their precursors during summer may be associated with marine biological sources and are attributable to the oxidation of unsaturated fatty acids\(^{75}\). In this context, the concentrations of \( \text{C}_9 \) (aqueous phase photo-oxidation of biogenic unsaturated fatty acids) show a gradual increase from late spring to late autumn with a maximum in June and thereafter show a gradual decrease towards winter and spring months (Fig. 3i). These results suggest that oxidations of biogenic unsaturated fatty acids are important sources of diacids over the WNP in summer/autumn, although their contribution is relatively small compared to those in winter and spring.

Influence of meteorological parameters.  Meteorological parameters such as wind speed, solar radiation and cloud cover are crucial for understanding the emission strengths of organic compounds and their oxidation processes in the atmosphere. As inferred from Figure S1, solar radiation was maximized in summer and minimized in winter/spring, indicating a strong photochemical oxidation during summer over the WNP. This result was further discussed in terms of seasonal variations in the mass concentration ratios of diacids. Being consistent with solar radiation, F/M, \( \text{C}_2/\text{C}_4 \), and \( \text{C}_2/\text{MeGly} \) ratios increased gradually from late spring to summer and stayed high until late autumn, and then decreased towards winter (Fig. 3o,q,r). These seasonal changes suggest an enhanced photochemical oxidation, superimposed with changing regional biology and meteorology in summer and important sources of the diacids over the WNP. On the other hand, wind speed and cloud cover were higher during spring and/or winter (Figure S1), suggesting the enhanced processing of precursor compounds associated with Asian outflows during atmospheric long-range transport that leads to higher concentrations of diacids during spring and winter over the WNP. Similar seasonal variations have been found in the mass concentration ratios of \( \text{C}_6/\text{C}_8 \) and \( \text{Ph}/\text{C}_4 \) with higher values in winter and/or spring (Fig. 3s and t). \( \text{C}_2/\text{C}_9 \) and \( \text{C}_7/\text{MeGly} \) ratios showed a gradual increase from early spring to autumn with a peak in early autumn, i.e., September (Fig. 3v and x). These ratios, then decreased towards winter. However, ratios of \( \text{C}_6/\text{Gly} \) did not show any clear seasonal trend during the study period, although ratios are slightly higher in summer and autumn (Fig. 3w). Although precipitation occurs throughout the year over the WNP, it was maximized in summer (Figure S1d); therefore, it is also an important sink for the diacids in addition to photochemical decompositions of oxalate-iron complexes, particularly in summer.

Source apportionment.  To quantitatively estimate the contribution of different sources to \( \text{C}_6 \) over the WNP, we performed a positive matrix factorization (PMF) analysis for the different seasons and the whole period (2001–2013) as shown in Fig. 4. PMF (version 5.1) is an effective source apportionment receptor model developed by the United States Environmental Protection Agency (U. S. EPA) and is often used in determining the sources of atmospheric aerosols\(^{76,77}\). A complete description of PMF analysis is discussed elsewhere\(^{78-80}\). The concentrations
of C$_2$–C$_6$, C$_9$, Ph, ωC$_2$, Gly, MeGly, and the tracers of water-soluble ions (MSA$^-$, Cl$^-$, nss-SO$_4^{2-}$, Na$^+$, NH$_4^+$, nss-K$^+$) were used as inputs in the PMF analysis. A total of 607 samples were used for this analysis. We identified 6 source profiles such as biogenic photochemical (indicated by blue color), mixed photochemical (green), anthropogenic 1 (purple), anthropogenic 2 (weak purple), marine biogenic (cyan), and biomass burning (red). The derived variations (%) of the species are shown in Fig. 4a–f. The contributions of all sources to the individual diacids for the different seasons as well as for the whole period are shown in Fig. 4g. The detailed descriptions of each PMF resolved-sources are described below for the different seasons over the WNP.

Based on the high variation (%) of loading of MSA$^-$, C$_2$, C$_3$, ωC$_2$, MeGly and weak or no loading of C$_6$, Ph, and Gly, we identified source 1 as biogenic photochemical 1. Source 2 was identified as mixed photochemical, which was confirmed by a significant loading of all species, which are majorly associated with photochemical oxidation of longer to shorter chain diacids. Source 3 was identified as anthropogenic 1 as evidenced by high loading of nss-SO$_4^{2-}$ and NH$_4^+$. Source 4 was attributed to anthropogenic 2, because it may be associated with biomass burning-derived VOCs as evidenced by high loading of Gly. Source 5 was considered as marine biogenic unsaturated fatty acids, as evidenced by high loading of Na$^+$, Cl$^-$, and C$_9$. Source 6 was attributed to biomass burning, which was majorly associated with primary emissions, as indicated by high loading of nss-K$^+$.

Figure 4b shows the contributions of all sources to C$_2$ for the different seasons and the whole period. The biogenic photochemical process is a prominent source of C$_2$, whose contribution to C$_2$ is highest in summer (47%), followed by winter (40%) and lowest in autumn (27%). The next prominent source is a mixed photochemical process, whose contribution is more abundant during winter (36%), summer (34%) and autumn (29%) and lowest in spring (28%). The contribution of anthropogenic source 1 is higher in autumn (14%) and winter (11%), whereas those of anthropogenic source 2 are higher in spring (24%), autumn (20%) and lower in summer (less than 9%) and winter (7%). Contributions of biomass burning to C$_2$ are highest in winter (3%) or autumn (2%) and lowest in summer (<1%), while the contributions of marine biogenic unsaturated fatty acids to C$_2$ are highest in summer (10%) followed by autumn (8%) and lowest in spring (3%). Overall, for the whole study period (Fig. 4b5), the contribution of biogenic photochemical process (42%) was the dominant source of C$_2$ followed...
by anthropogenic sources (1 plus 2 contribute ~32%) and mixed photochemical sources (20%). We found that marine biogenic unsaturated fatty acids are important sources for the formation of diacids over the WNP during winter and summer, respectively.

Conclusions and implications. The 13-year observations of water-soluble diacids and related compounds in marine aerosols in the WNP provided the following findings.

The molecular distributions of diacids showed a predominance of C$_2$ followed by C$_3$ and C$_4$. Seasonal variations of diacids and their precursor compounds showed maxima in winter to spring and minima in summer, except for C$_9$, which was maximized in summer. Annual concentrations of total diacids, ω-oxoacids, pyruvic acid and α-dicarboxyls showed continuous increases toward more recent years.

A decrease in anthropogenic emissions is inferred from a decrease in the trends of anthropogenic tracer compounds such as phthalic acid (Ph), adipic acid (C$_6$) and glyoxal (Gly), while an increase in biogenic emissions is confirmed from an increase in the concentrations of biogenic tracers including pyruvic acid (Pyr) and methyglyoxal (MeGly) during 2001 to 2013. On the other hand, satellite-derived oxidation levels (total columnar O$_3$ and tropospheric columnar NO$_3$) showed significant increases during the study period. These results demonstrate that the increased concentrations of diacids over the WNP are probably due to not only the increased biogenic emissions from East Asia but also increased oxidation processes during atmospheric long-range transport, while anthropogenic precursors of the diacids are decreased or constant during the study period over the WNP. These results further support our previous study, which reported the declined and increased concentrations of nss-SO$_4^{2-}$ and MSA$^-$, respectively, over the WNP during 2001–2012. We also found increased concentrations of C$_9$ and C$_{12}$ in summer, suggesting that marine biogenic unsaturated fatty acids are becoming important sources of diacids over the WNP, particularly, in summer.

These inferences are further supported by PMF analysis, which showed a biogenic photochemical contribution (42%) was a predominant source for C$_2$. This is the first study to explain the impact of heterogeneity in air masses on long-term trends of organic acids over the WNP. Therefore, the assessment of future climate effects of East Asian aerosols over the WNP will need continued observations because of the rapid changes in the emission strength of aerosols and their precursors over East Asia. The results of this study should be important for climate modelers, who are interested in radiative forcing calculations over the WNP.

Methodology
Collection of aerosol samples. Aerosol samples were collected on a quartz filter (20 × 25 cm, Pallflex 2500QAT-UP) from 2001 to 2013 using a high volume sampler (HVS) with a flow rate of 1 m$^3$ min$^{-1}$ at Chichijima Island (27°04′N, 142°13′E) in the WNP (Fig. 1). Before sampling, filters were pre-combusted at 450°C for three hours. The HVS was set up 5 m above ground level at the Satellite Tracking Centre of Japanese Aerospace Exploration Agency (JAXA, elevation 254 m a.s.l.) in Chichijima Island$^{43}$. 4–6 day integrated samples were collected during the study period. After the sampling, filters were put in a pre-baked (450°C for 6 hrs) glass bottle with a Teflon lined screw cap and stored at −20°C prior to the analysis of diacids. A total of 607 aerosol and 61 field blank samples were used in this study. A field blank sample was collected every ten aerosol samples by placing a clean filter in the cartridge of the HVS for 10 sec without the pump running.

Determination of diacids and related compounds. The collected filter samples were analyzed for diacids and related compounds using the improved method of Kawamura$^{41}$ and Kawamura and Ikushima$^1$. Briefly, a portion of each filter sample was extracted three times with 5 ml of organic-free ultrapure water (resistivity of >18 MΩ cm$^{-1}$, Sartorius arium 611 UV) under ultrasonication. The extracts were filtered through a Pasteur pipette packed with quartz wool to remove the filter debris and insoluble materials and placed in a 50 ml pear-shaped flask. The water extracts were pH-adjusted to 8.5–9.0 using a 0.05 M potassium hydroxide (KOH) solution and then concentrated to almost dryness using a rotary evaporator under vacuum. A 14% boron trifluoride in n-butanol solution was added to the extracts and then heated at 100°C for 1 hour to derivatize the carboxyl and aldehyde groups. The derived dibutyl esters and dibutoxy acetals were extracted with n-hexane and washed three times with ultrapure water (for the removal of polar compounds, including hydrogen fluoride (HF) and boronic acid (H$_3$BO$_3$) derived from the boron trifluoride) and concentrated using a rotary evaporator in a vacuum and nitrogen (N$_2$) blow down system. After being dried, a known amount of n-hexane was added to the ester fraction and derivatives were determined using a gas chromatography with a flame ionization detector (GC/FID; Hewlett-Packard, HP6890). Identification of the GC peaks was confirmed by comparing the GC retention times with those of authentic standards and confirmed by mass spectral examination using a GC/mass spectrometry (GC/MS; Thermoquest, Trace MS).

In order to check the recovery, 5μL of authentic diacids were spiked on the pre-combusted (450°C for 6 hrs) quartz filters and analyzed like a real sample using the above-mentioned procedure. The recoveries were 85% for oxalic acid, 90% for malonic acid and more than 90% for succinic, glutaric, and adipic acids. The analytical errors in the duplicate analyses are less than 10%. The concentrations of all diacids and related compounds reported in this study have been corrected using the field blanks. The blank levels were less than 5% for the major species measured in the real samples.

Several studies show that positive and negative artifacts can be significant during filter sampling and organic analysis of aerosols$^{42, 83}$. They generally arise from adsorption of gaseous species to the substrate surface and from particle losses on the walls and volatilization of semi-volatile species, respectively. In order to evaluate the potential artifacts, including adsorption or evaporation of collected particles to the gas phase, we performed simultaneous measurements of the diacids in marine aerosols using a HVS and a denuder/filter/denuder system$^{45}$. It is well established that the artifacts due to adsorption or evaporation are to be minor for the denuder system. We found good agreement between the two different techniques, suggesting that the HVS technique is valid for
collection of diacids in marine aerosols\(^1\). This is true because, in general, evaporation loss of particulate organic species is minor compared to gases adsorbed on the filter surfaces\(^1\). On the other hand, adsorption of gases is limited because diacids are predominantly observed in particle phase\(^1\). However, it is noteworthy that organic compounds originating from continental regions are much more aged during atmospheric transport toward Chichijima Island; therefore the artifacts may be minor for the diacids. More details about the potential sampling artifacts are described in elsewhere\(^1\).

**Statistical analysis.** To explore the inter-annual difference in the concentrations of the diacids and related compounds, the statistical regression analysis of variations (ANOVA)\(^2\) was performed by comparing all the data points during the study period. Differences with \(p < 0.05\) were considered to be statistically significant and indicated by a star (*) in the trend analyses as shown in Table 1. This statistical approach is simple, robust and easy to interpret\(^3\). For example, the sign of the diacid trend depends on the value of the slope of the regression analysis. In this kind of interpretation, when the slope is greater than zero, the trend is positive (increases) whereas the trend is negative (decreases) when the slope is less than zero. When the slope is equal to zero, there is no trend in the diacid concentrations. Uncertainties in the trends are reported as standard errors in the slope of regression lines and variations in the trends (% yr\(^{-1}\)) are also reported for each organic compound in Table 1.

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Author Contributions
All authors contributed to the data interpretation and reviewed the manuscript. K.K. designed and led the research work. E.T. performed the data analysis. S.K.R.B. wrote the manuscript with significant contributions from K.K.

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