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Total organic carbon and the contribution from speciated organics in cloud water: airborne data analysis from the CAMP²Ex field campaign

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Received: 13 May 2021 – Discussion started: 13 May 2021

Revised: 11 August 2021 – Accepted: 30 August 2021 – Published: 23 September 2021

Abstract. This work focuses on total organic carbon (TOC) and contributing species in cloud water over Southeast Asia using a rare airborne dataset collected during NASA's Cloud, Aerosol and Monsoon Processes Philippines Experiment (CAMP²Ex), in which a wide variety of maritime clouds were studied, including cumulus congestus, altocumulus, altostratus, and cumulus. Knowledge of TOC masses and their contributing species is needed for improved modeling of cloud processing of organics and to understand how aerosols and gases impact and are impacted by clouds. This work relies on 159 samples collected with an axial cyclone cloud-water collector at altitudes of 0.2–6.8 km that had sufficient volume for both TOC and speciated organic composition analysis. Species included monocarboxylic acids (glycolate, acetate, formate, and pyruvate), dicarboxylic acids (glutarate, adipate, succinate, maleate, and oxalate), methanesulfonic acid (MSA), and dimethylamine (DMA). TOC values range

between 0.018 and 13.66 ppm C with a mean of 0.902 ppm C. The highest TOC values are observed below 2 km with a general reduction aloft. An exception is samples impacted by biomass burning for which TOC remains enhanced at altitudes as high as 6.5 km (7.048 ppm C). Estimated total organic matter derived from TOC contributes a mean of 30.7 % to total measured mass (inorganics + organics). Speciated organics contribute (on a carbon mass basis) an average of 30.0 % to TOC in the study region and account for an average of 10.3 % to total measured mass.

The order of the average contribution of species to TOC, in decreasing contribution of carbon mass, is as follows (± 1 standard deviation): acetate (14.7 ± 20.5 %), formate (5.4 ± 9.3 %), oxalate (2.8 ± 4.3 %), DMA (1.7 ± 6.3 %), succinate (1.6 ± 2.4 %), pyruvate (1.3 ± 4.5 %), glycolate (1.3 ± 3.7 %), adipate (1.0 ± 3.6 %), MSA (0.1 ± 0.1 %), glutarate (0.1 ± 0.2 %), and maleate ($< 0.1 \pm 0.1$ %). Ap-

proximately 70 % of TOC remains unaccounted for, highlighting the complex nature of organics in the study region; in samples collected in biomass burning plumes, up to 95.6 % of TOC mass is unaccounted for based on the species detected. Consistent with other regions, monocarboxylic acids dominate the speciated organic mass (~ 75 %) and are about 4 times more abundant than dicarboxylic acids.

Samples are categorized into four cases based on back-trajectory history, revealing source-independent similarity between the bulk contributions of monocarboxylic and dicarboxylic acids to TOC (16.03 %–23.66 % and 3.70 %–8.75 %, respectively). Furthermore, acetate, formate, succinate, glutarate, pyruvate, oxalate, and MSA are especially enhanced during biomass burning periods, which is attributed to peat emissions transported from Sumatra and Borneo. Lastly, dust (Ca^{2+}) and sea salt (Na^+/Cl^-) tracers exhibit strong correlations with speciated organics, supporting how coarse aerosol surfaces interact with these water-soluble organics.

1 Introduction

The last 2 decades have witnessed an acceleration in research to unravel the nature of the organic fraction of airborne particles, including speciation (Hallquist et al., 2009; Kanakidou et al., 2005), with implications for how particles impact air quality, public health, and the planet's energy balance. However, there has been much less progress on organic research into cloud droplets, owing largely to the inaccessibility of clouds compared with particles that can be measured more easily near the surface. Analyzing organic matter in cloud water will lead to a better understanding of secondary aerosol formation and the nature of cloud condensation nuclei (CCN) that form droplets. The interaction of aerosol particles and clouds constitutes the largest uncertainty in estimating total anthropogenic radiative forcing (IPCC, 2013), which motivates the use of cloud composition as a tool to learn about these processes (MacDonald et al., 2020). Characterizing cloud water composition is insightful for atmospheric chemical processes such as the removal of gases that would otherwise participate in gas-phase reactions and for aqueous reactions that yield products without an efficient gas-phase source (e.g., dicarboxylic acids) (Ervens et al., 2013). While modeling of sulfate production in clouds is fairly mature (Barth et al., 2000; Faloona, 2009; Kreidenweis et al., 2003; Liu et al., 2021), the formation and evolution of organics in cloud water is much more poorly constrained (Ervens, 2015).

Advancing this research requires *in situ* measurements of cloud water composition. Among the most common methods of characterizing the organic fraction of cloud water samples is total organic carbon (TOC) analysis. Irrespective of whether studies have been focused on cloud water or fog water, most work has shown the following: (i) TOC is enhanced

in air masses with higher anthropogenic influence (Collett et al., 1998; Deguillaume et al., 2014; Herckes et al., 2013; Raja et al., 2009); (ii) ~ 40 %–85 % of TOC is attributed to unidentified species (Benedict et al., 2012; Boris et al., 2016, 2018; Herckes et al., 2002; Raja et al., 2008); (iii) organic acids usually account for 15 % of TOC (Deguillaume et al., 2014; Gioda et al., 2011; Straub et al., 2007); (iv) monocarboxylic acids are more abundant than dicarboxylic acids (Löflund et al., 2002); and (v) acetic and formic acids are the most dominant organic acids contributing to TOC (Collett et al., 2008; Gioda et al., 2011). Most of the aforementioned studies focused on fog, motivating a closer look at cloud water, as solute concentrations depend on the type of aqueous medium (Fig. 1). More specifically, TOC concentrations are reported to be higher in fog water relative to rain water (Kim et al., 2020), whereas cloud water solute concentrations exceed those in rain water (Decesari et al., 2005; Gioda et al., 2008).

Southeast Asia is an ideal laboratory to investigate the nature of TOC and its constituents, as it is impacted by a multitude of emission sources in an environment with persistent cloud cover from a variety of cloud types (e.g., shallow cumulus and cumulus congestus clouds) (Reid et al., 2013). The complex meteorology of the region makes it very difficult to model (Wang et al., 2013; Xian et al., 2013) but also simultaneously provides a remarkable opportunity to learn more about how aerosols impact (and are impacted by) tropical cloud systems. A knowledge gap exists, as there have been no studies of cloud composition in this region based on airborne measurements. Analysis of fog water at Baengnyeong Island in the eastern Yellow Sea revealed that organic acids accounted for 36 %–69 % of TOC (Boris et al., 2016). The Acid Deposition Monitoring Network in East Asia (<https://www.eanet.asia/>, last access: 10 August 2021) provides data on wet deposition at surface sites such as at the Manila Observatory (Ma et al., 2021), Metropolitan Manila (Metro Manila), Philippines, and is limited to inorganic ions. Previous studies such as the Seven South East Asian Studies (7SEAS) (Reid et al., 2013) and the Cloud, Aerosol and Monsoon Processes Philippines Experiment (CAMP²Ex) weatHER and CompoSition Monitoring (CHECSM) were carried out in this region; however, these campaigns were ground- and ship-based, and focused mainly on aerosol particles and not cloud composition (Hilario et al., 2020b; Reid et al., 2015, 2016). Moreover, it should be noted that there have also been a handful of high-elevation studies carried out in Southeast Asia examining fog and cloud water organic acids (i.e., Decesari et al., 2005; Li et al., 2017; Mochizuki et al., 2020).

Recent studies in Metro Manila, Philippines, provide the following results of relevance to this work: (i) a third to a half of the total aerosol particle mass is often unaccounted for after considering water-soluble species (inorganic and organic acid ions and elements) and black carbon (Cruz et al., 2019; Stahl et al., 2020); (ii) organic acids account for less

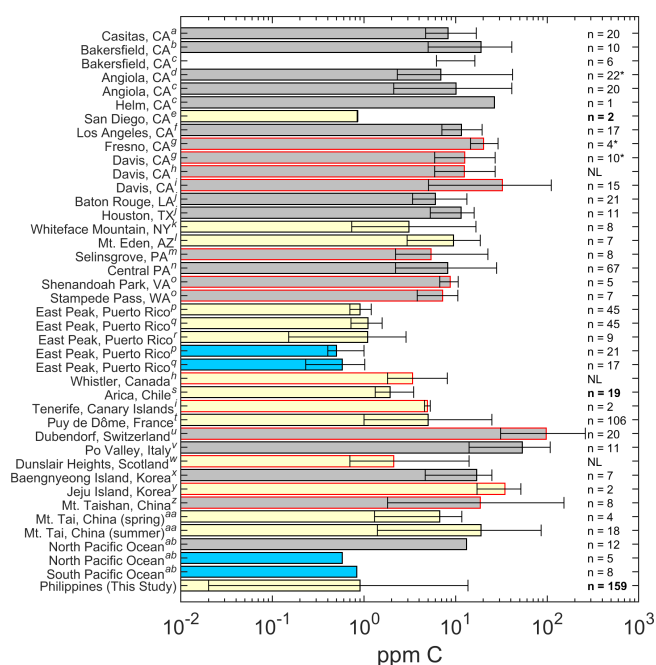


Figure 1. TOC (or dissolved organic carbon, DOC, if TOC values were unavailable) concentrations reported for past studies in relation to this work organized by continent. Bars represent the average values, and the error bars represent the minimum and maximum values. The absence of a solid bar means that no average was available. No error bars means that there was no range given, and “*” indicates that the median value was reported rather than an average. Gray, yellow, and blue bars represent studies looking at fog, clouds, and rain, respectively. Bars that are outlined in black are studies that used TOC, and bars outlined in red are studies that used DOC. The n values represent the number of samples used in the study, and NL means that the number of samples was not listed. Bolded n values denote airborne samples. This figure is similar to Fig. 2 in Herckes et al. (2013) with additional information presented and organized by continent. The superscript letters used in the figure denote the following: a – Boris et al. (2018), b – Collett et al. (1998), c – Herckes et al. (2002), d – Herckes et al. (2007), e – Straub et al. (2007), f – Erel et al. (1993), g – Ehrenhauser et al. (2012), h – Ervens et al. (2013), i – Zhang and Anastasio (2001), j – Raja et al. (2008), k – Cook et al. (2017), l – Hutchings et al. (2008), m – Straub et al. (2012), n – Straub (2017), o – Anastasio et al. (1994), p – Gioda et al. (2011), q – Gioda et al. (2008), r – Reyes-Rodríguez et al. (2009), s – Benedict et al. (2012), t – Deguillaume et al. (2014), u – Capel et al. (1990), v – Gelencser et al. (2000), w – Hadi et al. (1995), x – Boris et al. (2016), y – Decesari et al. (2005), z – Wang et al. (2011), aa – Shen (2011), and ab – Kim et al. (2020).

than 1 % of total aerosol mass, with oxalate being the most abundant acid (Stahl et al., 2020); (iii) organic acid concentrations are more enhanced during biomass burning periods (Hilario et al., 2020a), especially succinate and oxalate (Braun et al., 2020; Stahl et al., 2020); and (iv) wet deposition samples clearly show the influence of biomass burning tracer species on cloud composition (Ma et al., 2021).

Based on these points, we test two hypotheses: (i) the relative contribution of organic acids to TOC will exceed what was observed in the surface layer over Metro Manila owing to more aged air masses aloft compared with the surface layer in Metro Manila, which is exposed to fresher emissions; and (ii) clouds impacted by biomass burning emissions will exhibit chemical profiles shifted to higher TOC concentrations and with a greater portion of that TOC accounted for by organic acids. To address these hypotheses in addition to characterizing the organic fraction of cloud water, we utilized a rich set of cloud water samples collected around the Philippines during CAMP²Ex between August and October in 2019. The subsequent results and discussion focus on TOC concentrations in addition to the relative contribution and interrelationships between a suite of organic species (organic acids, methanesulfonic acid, and dimethylamine) both spatially and as a function of altitude and air mass source origin. A unique aspect of this dataset is the large sample number with both TOC and speciated organic acid information from an airborne platform.

2 Methods

2.1 Study overview

A total of 159 cloud water samples were collected by the NASA P-3B Orion aircraft across 19 research flights (RF; 23 August–5 October 2019) during CAMP²Ex and were measured for ions, pH, and TOC. Flights were based out of Clark International Airport (15.189° N, 120.547° E) and extended to regions around the island of Luzon (Fig. 2). Cloud water samples were collected over a wide range of altitudes ranging from 0.2 to 6.8 km.

2.2 Cloud water collection and handling

Samples were collected using an axial cyclone cloud-water collector (AC3), (Crosbie et al., 2018), which efficiently (> 60 % collection efficiency) collects cloud droplets with diameters > 20 μm. The size dependence of the collection efficiency may influence the measured properties of the bulk cloud water in cases where there is a strong size dependence in the droplet composition. Sample water evaporation was identified to affect low-liquid-water-content environments and may increase aqueous concentrations. For this study, the pipe position was set to position 10, as described in Crosbie et al. (2018), and mounted to the fuselage pylon approximately 300 mm from the skin. The AC3 has a shutter attached to a servo motor, allowing the collector to be closed when not in a cloud to prevent contamination. Samples were collected for between 10 s and 10 min depending on cloud availability and liquid water content (i.e., shorter times possible with higher liquid water content). Cloud water was collected in prewashed 15 mL plastic conical vials. Due to thorough prewashing of the plastic conical vials, leaching

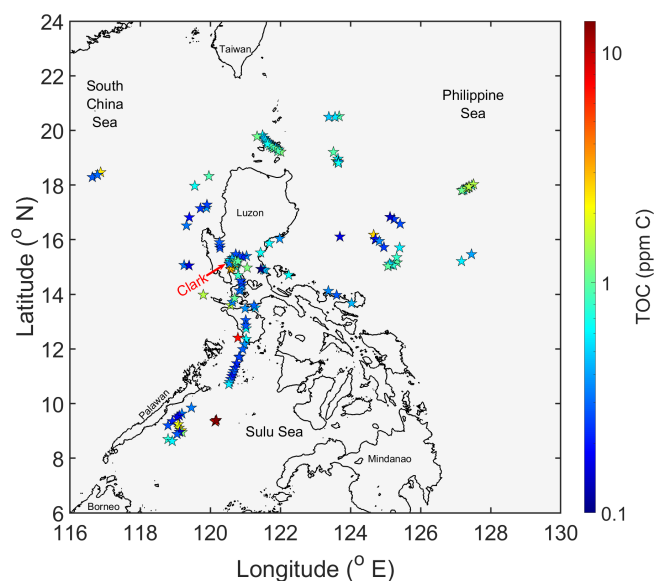


Figure 2. Map of the sample region. The stars represent the mid-point of the cloud water samples where total organic carbon (TOC) was measured, and they are colored by TOC on a logarithmic scale.

of organics into samples was negligible. Additional laboratory tests also indicated that there was no appreciable evidence that organics were adsorbing to the walls of the conical vials. Before each flight, the collector was flushed with ~ 1 L of ultra-purified Milli-Q water ($18.2 \text{ M}\Omega\cdot\text{cm}$) prior to obtaining two blank samples. Blanks, which were similarly flushed prior to collection, were also collected post-flight. During flight, samples were collected and stored in a cooler containing a sufficient number of ice packs to reduce possible decomposition. After flights, samples were immediately taken to an on-site laboratory where sample volumes were recorded and analyzed for ionic composition, TOC, and pH. A background was subtracted from the samples based on the bottom 10th percentile of all blanks collected during the campaign (both pre- and post-flight). The 10th percentile of the blanks was used instead of the mean, as it is a compromise between removing the influence of background contamination and conserving data points. Excess samples were stored in a refrigerator for future analyses that are outside the scope of this study.

2.3 Cloud water analysis

2.3.1 Ion chromatography

Cloud water was speciated using ion chromatography (IC; Dionex ICS-2100) immediately after each flight to reduce the possibility of sample degradation. Measured anionic species of interest were glycolate, acetate, formate, methanesulfonic acid, pyruvate, glutarate, adipate, succinate, maleate, oxalate, Cl^- , NO_2^- , Br^- , NO_3^- , and SO_4^{2-} . Measured cations were Na^+ , NH_4^+ , K^+ , dimethylamine (DMA), Mg^{2+} , and Ca^{2+} .

A 23 min instrument method with a 2 min equilibration period was used for both anion and cation columns, yielding a 25 min sampling period per sample. The instrument flow rate was 0.4 mL min^{-1} . The anions were measured using a Dionex IonPac AS11-HC $2 \text{ mm} \times 250 \text{ mm}$ column, a Dionex AERS 500e suppressor, and potassium hydroxide as the eluent. The cations were measured using a Dionex IonPac CS12A $2 \times 250 \text{ mm}$ column, a Dionex CERS 500e suppressor, and using methanesulfonic acid (MSA) as the eluent. The instrument methods used for analysis are as follows: (i) for anions, the eluent concentration started at 1 mM, ramped up to 4 mM between 0 and 10 min, ramped up to 6 mM between 10 and 11 min, and finally ramped up to 7 mM between 11 and 23 min using a suppressor current of 8 mA; (ii) for cations, the eluent concentration started at 5 mM and remained isocratic between 0 and 10 min, ramped up to 18 mM between 10 and 12 min, and finally remained isocratic at 18 mM between 12 and 23 min using a suppressor current of 22 mA. The limits of detection (LODs) for these species can be found in Table 1 and were calculated using $3S_a b^{-1}$, where S_a is the standard deviation of the response, and b is the slope of the calibration curve for that species.

2.3.2 Total organic carbon and pH

Total organic carbon (TOC) was measured using a Sievers 800 Turbo TOC analyzer. Sample aliquots were diluted to obtain the minimum volume needed by the instrument. The TOC analyzer was operated in turbo mode, and TOC values were averaged over a stable concentration period. Milli-Q water was used as an internal reference, and calibrations were performed before and after each batch of samples was analyzed (i.e., one batch approximately every three to four flights) using a range of different concentrations from an oxalate standard solution. A volume of approximately 10 mL was used for each measurement, and Milli-Q water was used intermittently to flush the instrument between each sample.

The pH of the cloud water samples was measured using an Orion Star™ A211 pH meter with an Orion™ 8103BNUWP ROSS Ultra™ pH electrode (precision of 0.01). A two-point calibration ($\text{pH} = 4$ and $\text{pH} = 7$) was performed at the beginning of the analysis of a particular flight's set of samples.

2.3.3 Units

While many studies report concentrations in terms of air-equivalent concentrations, we instead use the native liquid-phase concentrations. Aqueous concentrations of TOC and individual molecular components are reported in units of parts per billion by mass (ppb). TOC concentrations are specific to the mass of carbon atoms only, whereas molecules measured by IC correspond to the specific mass of the species (unless noted otherwise). TOC was converted to total organic matter (TOM) via multiplication by 1.8 (Zhang et al., 2005).

Table 1. Mass concentration limits of detection (LODs), minimum, maximum, mean, 1 standard deviation, and median values (ppb; left), in addition to the mass fraction (%; right) for the 159 CAMP²Ex cloud water samples with TOC data; note that mass fraction values depend on the C mass of each organic species shown. Total measured mass is defined as the sum of TOM, Na⁺, NH₄⁺, K⁺, Mg²⁺, Ca²⁺, Cl⁻, NO₂⁻, Br⁻, NO₃⁻, and SO₄²⁻. The abbreviations used in the table are as follows: MCAs – monocarboxylic acids, DCAs – dicarboxylic acids, MSA – methanesulfonic acid, DMA – dimethylamine, MO – measured organics, TOM – total organic matter, and DL – detection limit.

	LOD	Concentration (ppb)					Mass fraction (%)				
		Min	Max	Median	Mean	SD	Min	Max	Median	Mean	SD
Glycolate	98.76	< DL	224.8	10.65	13.49	20.34	0.0	35.0	0.6	1.3	3.7
Acetate	6.376	< DL	3926	159.4	251.4	409.9	0.0	100.0	10.5	14.7	20.5
Formate	19.77	2.095	3819	66.58	188.5	432.5	0.2	100.0	3.8	5.4	9.3
Pyruvate	5.452	< DL	296.9	5.359	24.35	41.28	0.0	56.1	0.5	1.3	4.5
MCAs	–	13.40	8042	253.4	477.8	857.8	0.6	100.0	16.9	22.6	33.9
Glutarate	43.70	< DL	258.7	< DL	6.824	27.32	0.0	1.0	0.0	0.1	0.2
Adipate	39.21	< DL	71.45	2.977	5.331	8.306	0.0	43.7	0.4	1.0	3.6
Succinate	38.64	< DL	1372	< DL	55.23	137.7	0.0	9.3	0.0	1.6	2.4
Maleate	14.81	< DL	14.73	< DL	0.6880	2.310	0.0	0.8	0.0	0.0	0.1
Oxalate	55.23	< DL	1135	38.64	95.65	148.2	0.0	43.9	1.7	2.8	4.3
DCAs	–	1.479	2766	61.40	163.7	295.3	0.1	69.8	3.3	5.5	7.5
MSA	88.01	< DL	24.79	3.9	5.107	5.313	0.0	0.9	0.1	0.1	0.1
DMA	56.97	< DL	183.8	< DL	11.16	32.41	0.0	45.3	0.0	1.7	6.3
MO	–	29.46	10820	334.3	657.7	1125	1.5	100.0	23.8	30.0	41.2
TOC	0.05	18.00	13 660	546	902	1435	↑ Relative to TOC (%)				
Inorganic/TOM	–	0.05	90.29	3.31	5.82	8.56	↓ Relative to total measured concentrations (%)				
pH	–	3.79	5.93	5.19	5.04	0.51					
MO	–	–	–	–	–	–	0.8	57.6	7.2	10.3	9.2
TOM	–	32.40	24 590	983	1624	2584	1.1	95.1	23.2	30.7	24.5
Inorganic	–	25.99	117 900	3894	8651	13 645	4.9	98.9	76.8	69.3	24.5
Na	16.62	< DL	29 280	609	1650	3192	0.0	26.6	9.5	10.0	7.8
NH ₄	176.8	< DL	8099	427	804	1010	0.0	68.1	7.2	11.2	13.2
K	142.4	< DL	1211	21.40	75.35	144	0.0	21.8	0.5	0.8	2.0
Mg	46.20	< DL	3701	57.87	182	379	0.0	4.0	1.0	1.1	0.9
Ca	74.81	< DL	1951	118	201	277	0.0	25.2	1.6	3.5	4.6
Cl	76.59	< DL	38 200	908	2451	4438	0.0	42.7	15.3	16.0	11.7
NO ₂	46.24	< DL	16.31	< DL	1.551	3.304	0.0	0.4	0.0	0.0	0.1
Br	7.817	< DL	44.05	1.398	4.081	7.120	0.0	0.2	0.0	0.0	0.0
NO ₃	17.33	< DL	26 560	572	1488	2925	0.0	43.4	10.4	12.6	8.2
SO ₄	414.7	2.318	15 680	868	1795	2495	0.4	34.9	14.1	14.0	8.5

The choice to focus on aqueous-equivalent rather than air-equivalent concentrations was made for various reasons. First, our analysis focuses heavily on relative amounts of species that were unaffected by multiplying native aqueous units by cloud liquid water content. Second, the definition of liquid water content applied by studies can vary widely based on the lower and upper bound of what is considered a droplet. Third, relationships between solute concentrations in cloud water and liquid water content, anticipated from nucleation scavenging, are ineffective when gases like acetic and formic acids absorb directly into droplets rather than having been part of the initial CCN activating into droplets (Khare et al., 1999; Marinoni et al., 2004). Lastly, many studies of cloud water composition that our results can be contrasted

with also use liquid units. The primary liquid units reported for cloud water concentrations are parts per million by volume (ppm) and ppb. However, it should be noted that species concentrations in cloud water can be high simply due to the liquid water content being low, or inversely, the concentrations can be low due to being diluted by high liquid water content.

2.4 Aerosol composition

To complement the cloud water composition results, we use aerosol composition results from a high-resolution time-of-flight aerosol mass spectrometer (AMS; Aerodyne, Inc.), which reports non-refractory composition for the submi-

rometer range (DeCarlo et al., 2006). As summarized by Hilario et al. (2021), the AMS deployed in CAMP²Ex functioned in 1 Hz Fast-MS mode with data averaged to a 30 s time resolution with the lower limit of detection (units of $\mu\text{g m}^{-3}$) as follows for the measured species: organic (0.169), NH_4^+ (0.169), SO_4^{2-} (0.039), NO_3^- (0.035), and Cl^- (0.036). Negative mass concentrations were recorded owing to the difference method used with the limits of detection. These negative values were included in the analyses to avoid positive biases but were interpreted as zero concentrations. We also use data specifically for the mass spectral marker representative of acid-like oxygenated organic species (m/z 44 = COO^+) (Aiken et al., 2008) and its mass relative to total organic mass (f_{44}). AMS data were omitted from the analysis if the total mass of all detected species was $< 0.5 \mu\text{g m}^{-3}$. By convention for airborne sampling, AMS data are reported at standard temperature and pressure (STP; 273 K, 1013 hPa).

AMS data were reported separately for cloud-free and cloudy conditions owing to the use of a counterflow virtual impactor (CVI) inlet (Brechtel Manufacturing Inc.) (Shingler et al., 2012) in clouds to isolate and dry droplets, leaving the residual particles for sampling by the AMS. Cloud-free data involve sampling with a separate inlet designed by the University of Hawaii (McNaughton et al., 2007). For cloud-free AMS results, data were selected 60 s before and after each cloud water sample's start and end time, respectively, when the aircraft was not in cloud. AMS–CVI data were reported for data collected within the period of cloud water collection. It should be noted that cloud-free AMS data are missing for some cloud water samples when the CVI was still in use for the 60 s before and after a sample's start and end time, respectively.

2.5 HYSPLIT

Air mass origination was determined using 5 d back trajectories from the National Oceanic and Atmospheric Administration (NOAA) Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Rolph et al., 2017; Stein et al., 2015). Trajectories were generated at 1 min temporal resolution with meteorological inputs from the Global Forecast System (GFS) reanalysis with a horizontal resolution of $0.25^\circ \times 0.25^\circ$ using the “model vertical velocity” method.

2.6 NAAPS

The Navy Aerosol Analysis and Prediction System (NAAPS) global aerosol model was implemented to assist in identifying biomass burning cases (Lynch et al., 2016) (<https://www.nrlmry.navy.mil/aerosol/>, last access: 10 August 2021). NAAPS relies on global meteorological fields from the Navy Global Environmental Model (NAVGEM) (Hogan and Brody, 1993; Hogan and Rosmond, 1991) that analyzes and forecasts a $1^\circ \times 1^\circ$ grid with 6 h intervals and 24 vertical lev-

els. In terms of identifying biomass burning cases, surface smoke concentrations were examined.

3 Cumulative results

3.1 Concentration statistics

TOC values ranged from 0.018 to 13.66 ppm C, with median and mean concentrations of 0.546 and 0.902 ppm C, respectively (Table 1). Samples in this study exhibited nearly the lowest mean TOC value of all cloud water studies surveyed in Fig. 1, with the other lowest values being over the Pacific Ocean west of San Diego, California (0.85 ppm C), (Straub et al., 2007) and East Peak, Puerto Rico (0.90 ppm C), (Gioda et al., 2008, 2011; Reyes-Rodríguez et al., 2009). The CAMP²Ex dataset exhibited the lowest minimum TOC value of all shown studies. For context, the highest mean and maximum TOC masses in cloud water studies were 34.5 and 51.7 ppm C, respectively, at Jeju Island, Korea, while the peak dissolved organic carbon (DOC) mass in cloud water was 85.6 ppm C at Mt. Tai, China. For comparisons to published cloud water measurements, DOC and TOC are assumed to be sufficiently similar in nature to directly compare values. Differences in TOC between our study and other work can partly be attributed to the different types of clouds studied in the CAMP²Ex region (e.g., cumulus congestus, cumulus, altocumulus, and altostratus) and the higher collection altitudes being conducive to enhanced liquid water contents and droplet sizes compared with stratocumulus clouds in regions like the northeastern (Straub et al., 2007) and southeastern Pacific Ocean (Benedict et al., 2012). Previous studies have primarily sampled stratocumulus or stratus clouds (Fig. 1). Also, some of our samples may have included rain water, which naturally has lower TOC concentrations than cloud water due to dilution (Fig. 1). To illustrate the importance of this dilution effect, an average of the mean values from the studies listed in Fig. 1 shows the following (ppm C): fog = 17.8, cloud = 6.4, and rain = 0.6. We further note that direct comparisons of our results to others need to account for the fact that water collectors have different transmission efficiency behaviors as a function of droplet size, as well as compositional differences across the droplet size spectrum (i.e., Boris et al., 2016; Collett et al., 2008; Herckes et al., 2013).

The order of species is as follows in terms of decreasing average contribution of C mass relative to total TOC (± 1 standard deviation): acetate ($14.7 \pm 20.5\%$), formate ($5.4 \pm 9.3\%$), oxalate ($2.8 \pm 4.3\%$), DMA ($1.7 \pm 6.3\%$), succinate ($1.6 \pm 2.4\%$), pyruvate ($1.3 \pm 4.5\%$), glycolate ($1.3 \pm 3.7\%$), adipate ($1.0 \pm 3.6\%$), MSA ($0.1 \pm 0.1\%$), glutarate ($0.1 \pm 0.2\%$), and maleate ($< 0.1 \pm 0.1\%$). An average of 70.0 % of TOC mass went unaccounted for. The predominant sources and production pathways of these species are briefly explained here. Precursor emission sources for ac-

etate and formate include plants, soil, vehicles, and biomass burning, with key production routes including the oxidation of isoprene, the ozonolysis of olefins, and peroxy radical reactions (Khare et al., 1999, and references therein). Pyruvate is considered the most abundant aqueous reaction product of methylglyoxal, generated by the oxidation of gas-phase anthropogenic volatile organic compounds (Boris et al., 2014; Carlton et al., 2006; Lim et al., 2013; Stefan et al., 1996; Tan et al., 2010). Glycolate has been linked to aqueous processing of acetate and is a precursor of glyoxylate (Boris et al., 2014), which is formed via the oxidation of glycolaldehyde by hydroxide radicals (Thomas et al., 2016). Oxalate is the most abundant dicarboxylic acid across different environments (Cruz et al., 2019; Stahl et al., 2020; Yang et al., 2014; Ziemba et al., 2011) and can be emitted directly by biogenic sources (Kawamura and Kaplan, 1987), combustion exhaust (Kawamura and Kaplan, 1987; Kawamura and Yasui, 2005), and biomass burning (Narukawa et al., 1999; Yang et al., 2014); however, it is also formed through the oxidation and degradation of longer-chain organic acids and acts as a notable tracer for cloud processing (Ervens et al., 2004; Sorooshian et al., 2006). Saturated organics like glutarate, adipate, and succinate are linked to fresh emissions, mainly from the ozonolysis of cyclic alkenes (such as from vehicular emissions), in the study region (Hatakeyama et al., 1985; Stahl et al., 2020). Maleate can be secondarily formed from the photooxidation of benzene (Rogge et al., 1993) or from the primary emissions of combustion engines (Kawamura and Kaplan, 1987). Alkyl amines (i.e., DMA) have numerous sources such as biomass burning, vehicular emissions, industrial activity, animal husbandry, waste treatment, and the ocean (Youn et al., 2015). Finally, MSA is formed via photooxidation reactions involving dimethylsulfide (DMS) from oceanic emissions (Berresheim, 1987; Saltzman et al., 1983) or dimethyl sulfoxide (DMSO) from anthropogenic emissions (Yuan et al., 2004), in addition to being linked to agricultural emissions and biomass burning (Sorooshian et al., 2015).

Measured organic species were further grouped into categories: monocarboxylic acids (MCAs; glycolate, acetate, formate, and pyruvate), dicarboxylic acids (DCAs; glutarate, adipate, succinate, maleate, and oxalate), and measured organics (MO; the sum of MCAs, DCAs, MSA, and DMA). Total MCA concentrations accounted on average for $\sim 75\%$ of MO and were approximately 4 times larger than those of DCAs. MO values ranged from 29.46 to 10 820 ppb, accounting for an average of 30.0% (median 23.8%) of TOC when masses were converted to just the C masses of the measured species (Table 1). Examples of other undetected organics include tricarboxylic acids, aromatics, alcohols, sugars, carbohydrates, and aldehydes. Previous studies reported that undetected species account for $\sim 45\%$ (Boris et al., 2016) and 82.9% (Boris et al., 2018) of organics. Interestingly, the ionic charge balance for the 159 samples shows an anion deficit (Fig. S1 in the Supplement), with a slope of 0.95

(i.e., anion charge on y axis). This strong charge balance suggests that detected organic species were balanced by cations detected via IC analysis. Species contributing to the anion deficit likely include a mix of unspiciated organic and inorganic anions.

TOC was converted to total organic matter (TOM) by multiplying it by 1.8 (Zhang et al., 2005), as in other cloud water studies (Boris et al., 2016, 2018), in order to compare it to total measured mass (i.e., the sum of TOM, Na^+ , NH_4^+ , K^+ , Mg^{2+} , Ca^{2+} , Cl^- , NO_2^- , Br^- , NO_3^- , and SO_4^{2-}). We caution that using a fixed 1.8 conversion value yields uncertainty, as samples were collected in a range of air masses, but 1.8 is a value that is fairly intermediate with respect to those reported in the literature: 1.6 ± 0.2 for urban aerosols (Turpin and Lim, 2001), 2.07 ± 0.05 in nonurban areas (Yao et al., 2016), and values for biomass burning organic aerosols ranging from 1.56 to 2.0 (Aiken et al., 2008; Turpin and Lim, 2001) based on fuel type and combustion conditions (Aiken et al., 2008). Higher values are expected for more oxidized organics. Estimated TOM accounted for a median and mean of 23.2% and 30.7% of total measured mass, respectively, with the maximum for a single sample being 95.1%. The median and mean ratios of MO to TOM were 38.1% and 46.4%, respectively. Furthermore, the median and mean ratios of MO to total measured mass were 7.2% and 10.3%, respectively, with a maximum of 57.6%. On average, chloride, sulfate, and nitrate were the most abundant species ($\geq 12.6\%$), with the median and mean ratio of total inorganic mass to TOM being 3.3 and 5.8, respectively. The pH of the cloud water with TOC measurements ranged from 3.79 to 5.93 and averaged 5.04 ± 0.51 . The lowest pH values all occurred over the ocean.

Our calculated percentages of MO relative to total measured mass are in contrast with results from a surface site in Metro Manila (Stahl et al., 2020), where most of the same organic species (adipate, succinate, maleate, oxalate, and MSA) accounted for $\sim 1.3\%$ of the total aerosol mass, excluding black carbon. Therefore, the first hypothesis of this study – that the contributions of measured organic species account for a greater portion of total measured mass in cloud water compared with surface particulate matter – holds true.

Gravimetry was used to measure total mass in the surface measurements, whereas total measured mass was more restrictive in cloud water in terms of being based on measurable species, thereby qualifying our percentages as an upper bound. However, the measured ions in cloud water should contribute relatively more to total measured mass in cloud water, owing to their hygroscopic nature (e.g., sea salt) and the greater ease with which they become associated with cloud water compared with more hydrophobic species (Chang et al., 2017; Dalirian et al., 2018; Pringle et al., 2010) like black carbon that contribute significantly to total aerosol mass in the boundary layer of Metro Manila (Cruz et al., 2019). For example, black carbon accounted

for 78.1 % and 51.8 % of the total mass between 0.10–0.18 and 0.18–0.32 μm in Metro Manila surface aerosol particles (Cruz et al., 2019), respectively, which are size ranges that are highly relevant to droplet activation. Air masses aloft in the CAMP²Ex region, especially those processed by clouds, are likely more aged and oxidized than fresh organic emissions (e.g., automobiles, industry, and burning) in the surface layer over Metro Manila, which is the most populated urban area within the CAMP²Ex flight domain. Recent work has shown that cloud processing of isoprene oxidation products (a key fraction of organic precursor vapors involved with organic aerosol generation) is the main source of secondary organic aerosol (SOA) in the mid-troposphere (4–6 km) (Lamkaddam et al., 2021). This motivates the examination of vertical TOC and organic species characteristics in more detail, which is discussed next.

3.2 Vertical profiles

The vertical profiles of TOC mass were of interest, as they relate to the general vertical distribution of organic matter in the troposphere. Measurements off the coast of Japan approximately 2 decades ago during the ACE-Asia campaign revealed unexpectedly high organic aerosol concentrations in the free troposphere due to presumed SOA formation (Heald et al., 2005). During that campaign, organic aerosol concentrations in the boundary layer and free troposphere, and their relative contribution to total non-refractory aerosol mass (organic, SO_4^{2-} , NO_3^- , and NH_4^+), were amongst the highest of the various global regions examined (Heald et al., 2011). Therefore, it is of interest to examine these types of vertical profiles farther south in the CAMP²Ex region where data are more scarce, with the unique aspect of this work being the focus on cloud water composition.

The highest TOC masses were observed in the bottom 2 km, with a general reduction above that altitude (Fig. 3). The decrease in the TOC concentration with respect to altitude could be attributed to more dilution in larger droplet sizes; the results of cloud microphysical data will be the focus of forthcoming work. Four data points influenced by biomass burning were singled out (red markers Fig. 3a) as they had systematically higher TOC masses than other points. Those points will be discussed in more detail in Sect. 4, and it is noteworthy that clouds were impacted by biomass burning across a wide range of altitudes up to almost 7 km.

Focusing on the non-biomass burning (non-BB) data, there was considerable variation in TOC in the bottom 2 km, with concentrations as low as 0.144 ppm C and as high as 3.362 ppm C. Interestingly, cloud water collected above 5 km still tended to show enhanced TOC masses, reaching up to 1.530 ppm C (6.1 km) among the non-BB points. The composition contributing to TOC was similar with altitude in non-BB and biomass burning (BB) conditions, with $\sim 75\%$ of TOC mass unaccounted for by the measured species and

MCAs dominating the measured organic mass (Fig. 3b). The exception to that was the high-altitude BB point where 95.6 % of TOC was unassigned. Figure 3c–d show that there was some qualitative agreement in the vertical profile of AMS organic and m/z 44 mass concentrations for data collected immediately adjacent to the cloud water samples in cloud-free air; more specifically, the highest concentrations of AMS organic mass, m/z 44 mass, and TOC were in the bottom 2 km. However, some interesting differences exist, as they related to specific air mass types, as will be discussed in Sect. 4. Some differences could be rooted in how AMS data represent submicrometer particles whereas cloud water data encompass a wider range of particle sizes that activated into cloud droplets (including supermicrometer dust and sea salt particles) and also gases partitioning to cloud water.

Vertical profiles of ratios representative of the relative amount of oxidized organics are shown in Fig. 4. The MO:TOC ratio was quite variable with altitude, ranging from 0.16 to 0.32 based on the locally averaged curve shown; individual sample values ranged from 0.01 to 0.92. Vertically resolved ratio values for f_{44} in cloud-free air and in cloud (downstream CVI) ranged on average from 0 to 0.35 and from 0.13 to 0.35, respectively. While mass concentrations decreased with altitude (Fig. 3), ratios relevant to the degree of organic aerosol oxidation and makeup of the organic component of cloud water did not exhibit a clear change with altitude.

4 Case studies

Four subsets of samples are examined here to probe how the organic nature of cloud water varies for distinct air masses. Sources of the air masses are visually shown in Fig. 5 based on 5 d HYSPLIT back trajectories: (i) “North” (RF11, $n = 20$) collected off the northern coast of Luzon with influence from East Asia, the Korean Peninsula, and Japan; (ii) “East” (RF13, $n = 11$) collected off the eastern coast of Luzon with back trajectories traced to southern China with subsequent passage across Luzon before arriving in the sample collection area; (iii) “Biomass Burning” (RF09, $n = 4$) collected to the southwest of Luzon above the Sulu Sea with influence from biomass burning plumes from Borneo and Sumatra primarily consisting of peat as the fuel type (Field and Shen, 2008; Levine, 1999; Page et al., 2002; Stockwell et al., 2016; Xian et al., 2013); and (iv) “Clark” (RF04, RF06, RF07, RF09, RF10, and RF11, $n = 25$) collected around the operational area over Luzon, approximately ~ 90 km northwest of Metro Manila, with back trajectories extending to the west and southwest of Luzon.

Biomass burning samples were identified based on the following criteria: flight scientist notes, elevated surface smoke concentrations and aerosol optical depth (AOD) from the NAAPS model, and the remarkable enhancement in chemical concentrations in cloud water. TOC, K^+ , SO_4^{2-} , and NH_4^+

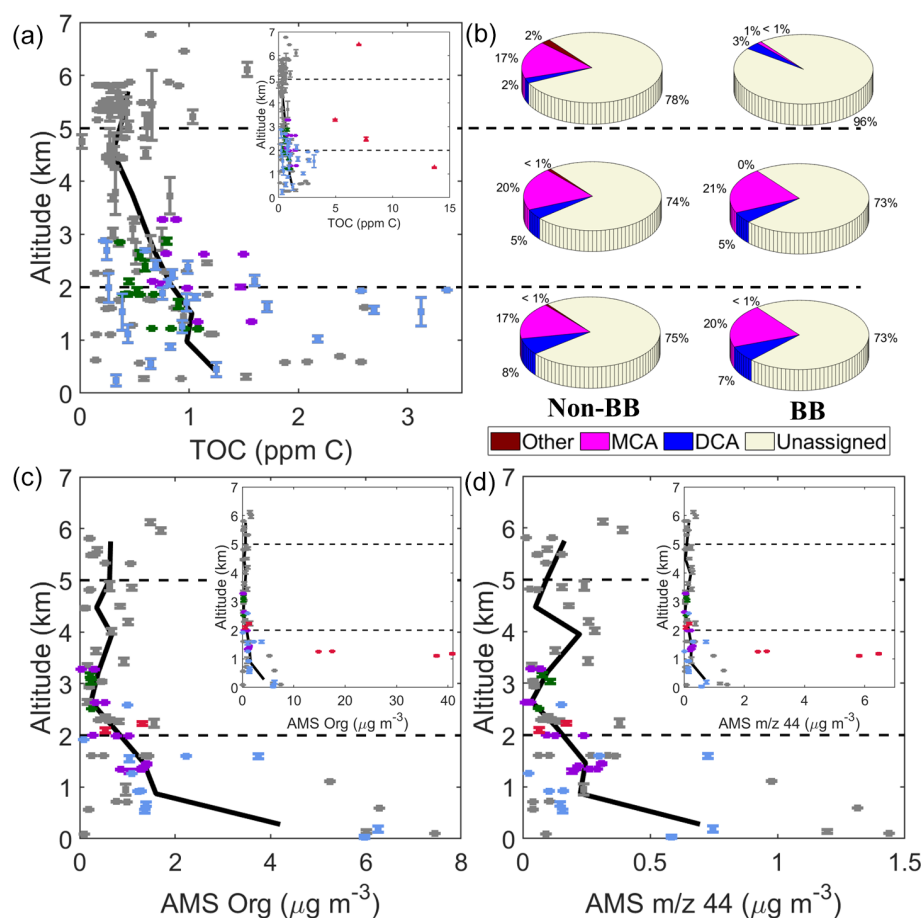


Figure 3. (a) Vertical profile of TOC concentrations ($n = 159$ samples) with the smaller inset including the four samples with enhanced TOC owing to biomass burning (BB) influence. (b) Mass fractions of different subsets of species contributing to TOC at high (> 5 km), mid (2–5 km), and low (< 2 km) altitude, with the beige area representing undetected species. Vertical profile of AMS (c) organic and (d) m/z 44 mass concentrations corresponding to spatially and temporally adjacent cloud-free periods of the collected cloud water samples. Colors in panels (a), (b), and (d) represent the case study points in Sect. 4: North (green), East (purple), Biomass Burning (red), Clark (blue), and non-case points (gray). The solid black lines in panels (a), (b), and (d) represent locally weighted average values. The error bars represent 1 standard deviation of the altitude variance.

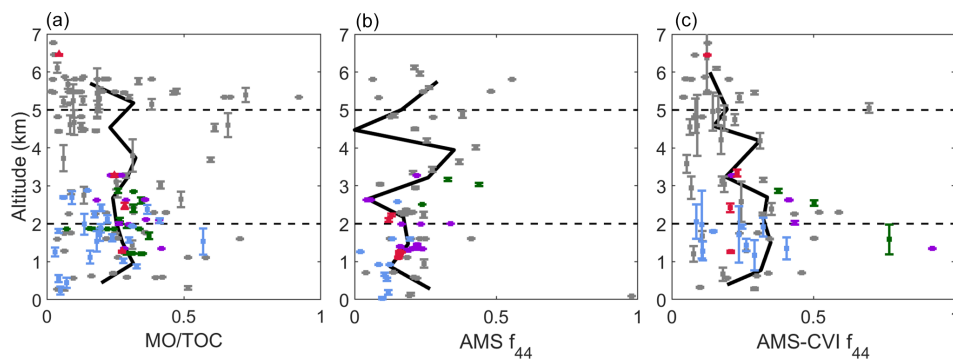


Figure 4. Vertical profile of (a) the ratio of C mass from measured organics (MO) to TOC for cloud water samples, (b) AMS f_{44} in cloud-free air and (c) AMS-CVI f_{44} in cloudy air. AMS data in panel (b) correspond to cloud-free periods that were spatially and temporally adjacent to the collected cloud water samples, while the data in panel (c) are within the period of cloud water collection times in cloud. Colors in panels (a), (b), and (d) represent the same case study points as in Fig. 3: North (green), East (purple), Biomass Burning (red), Clark (blue), and non-case points (gray). The black lines in panels (a), (b), and (d) represent locally weighted average values.

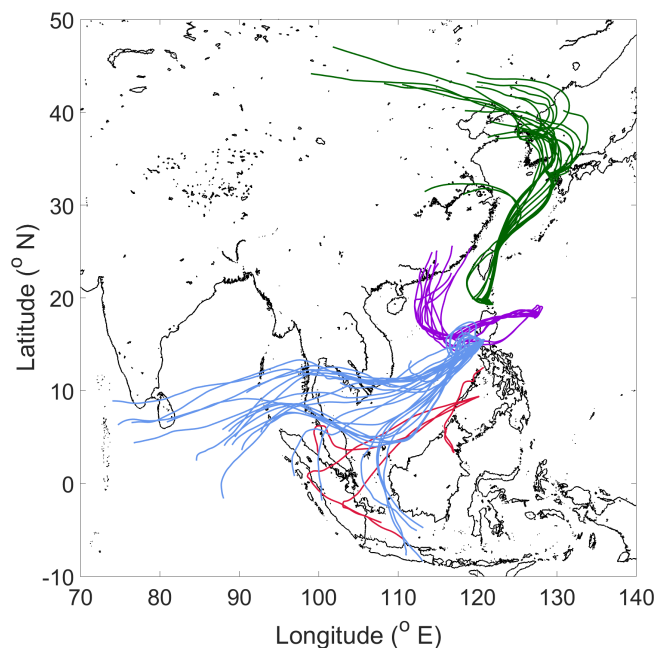


Figure 5. Spatial summary of 120 h back trajectories for each sample included in respective case study sample sets: North (green; $n = 20$), East (purple; $n = 11$), Biomass Burning (red; $n = 4$), and Clark (blue; $n = 25$).

in particular were enhanced in these samples with concentrations exceeding 4 ppm C, 0.13 ppm, 2.3 ppm, and 2.5 ppm, respectively.

The vertical profile results shown previously (Figs. 3, 4) display markers corresponding to these four case studies. With the exception of one BB sample collected at 6.5 km, samples in the four cases were obtained below 3.3 km.

4.1 North

This category of samples was unique in that the mean MO (527.5 ± 301.6 ppb) and TOC (636.1 ± 230.4 ppb C) concentrations were the lowest of all four cases (Table 2). The largest three organic contributors to TOC (± 1 standard deviation) were acetate (177.8 ± 72.96 ppb C; $11.5 \pm 4.0\%$), oxalate (148.7 ± 81.47 ppb C; $6.0 \pm 1.3\%$), and formate (83.16 ± 79.65 ppb C; $3.0 \pm 2.2\%$). Maleate and DMA were not detected for this case, and 73.3 % of TOC went unaccounted for. Samples in this category were collected between 1.2 and 2.9 km, without any pronounced organic chemical trends with altitude (Figs. 3, 4).

This case exhibited a few distinct features worth noting. First, it had the highest sea salt presence based on the highest case-wide concentrations of Na^+ (3238 ± 2861 ppb), Cl^- (5277 ± 4333 ppb), Mg^{2+} (347.1 ± 328.3 ppb), and Br^- (15.56 ± 8.036 ppb), the latter of which is a trace component of sea salt (Seinfeld and Pandis, 2016). MSA originates partly from marine emissions of DMS, but its concentra-

tion was among the lowest of all species for all four cases with a mass contribution to total TOC (based on C mass) of only $0.17 \pm 0.05\%$ in the North category (Table 3). In their analysis of aerosol data in the surface layer of Metro Manila, Stahl et al. (2020) showed lower overall organic acid aerosol concentrations in the northeast monsoon season where northeasterly air masses originated predominantly from East Asia; Stahl et al. (2020) also showed that those air masses were characterized by an enhancement in organic acid masses in the supermicrometer size range owing to adsorption to coarse particle types such as sea salt and dust, although with a preference for dust (Mochida et al., 2003; Rinaldi et al., 2011; Sullivan and Prather, 2007; Turekian et al., 2003). As there was no direct evidence of dust in this case because the $\text{Ca}^{2+} : \text{Na}^+$ ratio was on average (0.04) nearly the same as sea salt (0.038) (Seinfeld and Pandis, 2016), organic acids could have interacted with sea salt. There were strong correlations between sea salt constituents, TOC, and almost all detected organics (Table S1 in the Supplement).

The second notable feature of this case was limited air mass aging characteristics based on speciated ratios. The acetate : formate ratio is often used to indicate the relative influence of fresh emissions (higher ratios) as compared to secondary production (lower ratios) (Talbot et al., 1988; Wang et al., 2007). In at least one study, fresh emissions were linked to cloud water ratios above 1.5 and aged samples having values below 1 (Coggon et al., 2014). The mean acetate : formate ratio for this air mass category was 4.21 ± 3.26 , which was the highest of all four categories in Table 2, suggestive of fresh emissions and low aging. This was consistent with the $\text{Cl}^- : \text{Na}^+$ ratio (1.70 ± 0.13) being close to sea water (1.81); our use of this ratio in the study assumes that these species originate primarily from sea salt. Lower $\text{Cl}^- : \text{Na}^+$ values in the study region coincide with sea salt reactions with acids such as sulfuric, nitric, and organic acids (AzadiAghdam et al., 2019). This was one of the two cases that had adipate present, with this category exhibiting the highest mean concentration (5.146 ± 6.266 ppb). This suggests that there was influence from cyclic organics possibly originating from combustion sources, among others, during transport to the sample region. Adipate exhibited negative correlations with almost all other organic species in this case (r from -0.48 to -0.72), suggestive of limited aging to form shorter-chain carboxylic acids via photochemical reactions (Table S1 in the Supplement). With the exception of adipate, interrelationships between the other organics detected in this case exhibited positive and significant correlations with one another, which is suggestive of common precursors and/or production mechanisms. Therefore, the results of the North case point to influences from marine emissions and limited aging signatures based on speciated ratios.

Table 2. Speciated concentrations of organics (ppb) for each case study: the first group of rows is monocarboxylic acids (MCAs); the second group of rows is dicarboxylic acids (DCAs); the third group of rows is other organics, including total measured organics (MO) and total organic carbon (TOC); the fourth group of rows is inorganic ions; and the fifth group is select ratios. *n* stands for number of samples, Ace denotes acetate, and For represents formate.

	North (<i>n</i> = 20)					East (<i>n</i> = 11)					Biomass Burning (<i>n</i> = 4)					Clark (<i>n</i> = 25)				
	Min	Max	Median	Mean	SD	Min	Max	Median	Mean	SD	Min	Max	Median	Mean	SD	Min	Max	Median	Mean	SD
Glycolate	5.963	37.18	15.80	17.59	9.15	5.339	30.00	12.21	13.68	7.249	<DL	46.86	7.20	15.31	22.10	<DL	53.42	6.900	11.68	14.61
Acetate	1.935	288.8	184.9	177.8	72.96	301.7	423.5	358.6	359.0	40.71	47.85	3926	1704	1845	1668	<DL	1105	185.2	296.7	325.8
Formate	10.28	232.2	62.66	83.16	79.65	61.02	492.8	248.3	258.2	122.2	151.0	3819	2370	2178	1589	2.422	1041	152.0	266.1	316.8
Pyruvate	2.143	126.5	35.91	42.98	38.98	7.502	78.24	24.65	32.25	21.01	<DL	296.9	103.4	125.9	126.1	1.072	161.8	16.08	30.31	35.84
MCAs	25.32	632.2	299.9	321.5	183.6	431.3	923.7	673.2	663.2	142.7	245.7	8042	4184	4164	3336	31.93	2066	369.2	604.7	641.9
Glutarate	<DL	10.18	<DL	1.527	2.758	<DL	10.86	4.074	5.122	3.672	62.46	258.7	140.2	150.4	82.20	<DL	62.46	1.358	9.423	16.85
Adipate	<DL	17.44	<DL	5.146	6.266	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	37.43	<DL	3.777	7.888
Succinate	<DL	136.2	28.09	42.30	47.84	15.45	176.9	63.90	74.11	57.27	24.58	1372	416	557.0	575.6	<DL	498.5	18.96	67.74	123.7
Maleate	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	11.61	5.360	5.583	6.456	<DL	14.73	<DL	2.714	4.170
Oxalate	37.53	330.4	124.8	148.7	81.47	52.51	311.2	123.3	153.6	81.06	303.8	1135	520.1	619.7	360.1	5.547	448.9	43.63	88.33	103.9
DCAs	67.84	467.1	149.3	197.6	125.2	80.60	493.4	194.1	232.9	136.4	735.8	2766	914.6	1333	968.9	7.673	1010	72.10	172.0	238.0
MSA	1.550	14.72	7.748	8.290	3.160	3.10	17.82	10.07	10.57	4.400	<DL	24.79	3.874	8.135	11.69	<DL	10.85	3.874	4.184	3.620
DMA	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	61.25	<DL	6.454	15.89
MO	99.36	1089	483.8	527.5	301.6	567.0	1364	855.6	906.6	269.4	1017	10820	5094	5505	4187	54.05	3054	433	787.3	837.3
TOC	364.0	1085	555.0	636.1	230.4	663.0	1570	985.0	1051	330.6	4974	13660	7366	8342	3730	220.0	3362	849	1181	920.2
Na ⁺	693.2	11 870	2273	3238	2861	617.7	6546	1970	2569	1738	832.6	4425	2160	2394	1624	12.28	5870	624.9	1105	1403
NH ₄ ⁺	180.6	1955	644.4	847.0	515.5	512.7	2379	1307	1432	587.4	2517	8099	3685	4496	2483	45.29	2880	946.8	1009	814.6
K ⁺	13.83	404.9	49.69	88.89	99.63	18.32	493.0	66.14	122.9	136.0	132.2	724.3	272.3	350.3	258.4	2.462	264.0	31.82	63.07	75.51
Mg ²⁺	41.42	1338	236.3	347.1	328.3	62.35	668.1	209.4	273.2	182.7	83.62	500.5	242.3	267.2	191.2	<DL	631.2	64.73	117.2	152.6
Ca ²⁺	<DL	764.9	96.85	173.8	218.9	49.47	778.2	166.5	269.2	219.1	105.7	533.5	236.0	277.8	209.7	38.83	903.5	176.8	230.3	183.6
Cl ⁻	1445	18 520	3772	5277	4333	900.2	8357	2760	3510	2196	1126	5989	2553	3055	2124	45.43	9083	990.6	1716	2107
NO ₂ ⁻	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	<DL	5.598	1.339	2.069	2.670	<DL	16.06	<DL	3.427	5.126
Br ⁻	3.496	35.66	12.59	15.56	8.036	2.098	6.992	4.195	4.132	1.482	1.398	6.293	2.448	3.147	2.174	<DL	13.29	1.398	2.545	3.204
NO ₃ ⁻	477.2	5265	1197	1810	1506	1084	8724	2902	3772	2296	1880	7045	3344	3903	2277	64.84	2759	691.3	930.8	736.1
SO ₄ ²⁻	1305	12 120	3281	4503	2865	1212	5296	2819	3223	1385	2313	9993	4177	5165	3343	23.39	4406	1157	1416	1157
pH	3.92	4.92	4.48	4.40	4.40	4.27	4.92	4.51	4.51	0.20	3.96	4.65	4.35	4.33	0.29	4.66	5.76	5.25	5.29	0.33
Ace/For	0.19	9.66	2.65	4.21	3.26	0.75	5.67	1.52	1.93	1.51	0.32	1.03	0.70	0.69	0.30	0	3.86	0.98	1.12	0.84
Cl ⁻ /Na ⁺	1.52	2.08	1.69	1.70	1.13	1.28	1.51	1.40	1.40	0.06	1.07	1.43	1.35	1.30	0.16	1.38	3.70	1.69	1.84	0.56
Ca ²⁺ /Na ⁺	0	0.08	0.04	0.04	0.02	0.05	0.14	0.10	0.10	0.03	0.08	0.13	0.12	0.11	0.02	0.05	6.17	0.32	0.99	1.46
K ⁺ /Na ⁺	0.02	0.03	0.02	0.02	0.01	0.03	0.08	0.04	0.04	0.01	0.10	0.18	0.16	0.15	0.04	0.01	3.93	0.05	0.25	0.78
MO/TOC	0.07	0.37	0.29	0.27	0.08	0.18	0.42	0.29	0.31	0.07	0.04	0.28	0.26	0.21	0.11	0.03	0.57	0.19	0.20	0.13

Table 3. Average organic composition for each case study where the first, second, and third groups of rows show the percentage contributions (%) of individual components to monocarboxylic acids (MCAs), dicarboxylic acids (DCAs), and total organic carbon (TOC), respectively.

Group	Species (%)	North (<i>n</i> = 20)		East (<i>n</i> = 11)		BB (<i>n</i> = 4)		Clark (<i>n</i> = 25)	
		Mean	SD	Mean	SD	Mean	SD	Mean	SD
MCAs	Glycolate	7.20	9.20	1.84	0.81	5.09	9.87	17.65	29.05
	Acetate	64.03	17.74	64.20	10.85	45.86	14.07	46.35	23.98
	Formate	16.54	9.83	28.62	10.35	46.40	7.24	29.09	11.72
	Pyruvate	12.23	6.90	5.33	2.87	2.65	1.87	6.91	4.90
DCAs	Glutarate	0.65	1.00	2.91	1.41	17.15	9.28	4.02	5.02
	Adipate	8.04	9.47	0	0	0	0	16.05	21.48
	Succinate	20.82	20.08	38.52	12.15	41.95	25.27	26.53	25.39
	Maleate	0	0	0	0	0.75	0.88	3.20	5.93
	Oxalate	70.49	12.29	58.57	11.52	40.16	16.50	50.20	17.42
TOC	MSA	0.17	0.05	0.13	0.04	0.01	0.02	0.06	0.07
	DMA	0	0	0	0	0	0	0.43	1.17
	MCAs	17.79	6.17	23.66	5.99	16.03	10.13	16.28	11.91
	DCAs	8.75	2.65	6.82	2.94	5.21	1.60	3.70	2.67
	MO	26.72	7.86	30.61	7.35	21.25	11.32	20.46	13.34
	Undetected	73.28	7.86	69.39	7.35	78.75	11.32	79.54	13.34

4.2 East

The dominant organic contributors to TOC (1051 ± 330.6 ppb C) in the East case were the same as in the North case with the difference being the order after acetate (± 1 standard deviation): acetate (359.0 ± 40.71 ppb; 14.9 ± 3.1 %), formate (258.2 ± 122.2 ppb; 7.2 ± 3.8 %), and oxalate (153.6 ± 81.06 ppb; 3.8 ± 1.2 %). The percentage of TOC unaccounted for by the speciated measurements (69.4 %) was the lowest out of all of the cases. This case resembled the North case in that there was marine influence, although the differences were a more pronounced dust influence and greater evidence of aging based on chemical ratios. Marine signatures come from the second highest concentrations of Na^+ , Cl^- , and Mg^{2+} after North, with high correlations between these species (Table S2).

Unlike the previous case, the $\text{Ca}^{2+} : \text{Na}^+$ ratio (0.10) was elevated from that of typical sea salt (0.038). Wang et al. (2018) showed that East Asian dust can get lofted up during dust storms, which could contribute to the transport to the Philippines. Previous studies have shown that organic acids adsorb more readily to dust compared with sea salt due to dust's more alkaline nature (Stahl et al., 2020; Sullivan and Prather, 2007). While Ca^{2+} was correlated with 6 of the 11 organic species for this case ($r = 0.70$ – 0.96 ; Table S2), the magnitude of the correlations was very similar to those between either Na^+ or Cl^- and the speciated organics. TOC also exhibited similar correlations with Na^+ , Cl^- , and Ca^{2+} ($r = 0.83$ – 0.87). Therefore, it is too difficult with the given data to assert whether (if at all) the organic acids had a preference towards either salt or dust aerosol particles; of note, however, is that oxalate exhibited the strongest correla-

tion with either Na^+ , Cl^- , or Ca^{2+} ($r = 0.96$ – 0.99) among all species and also TOC. Additionally, Park et al. (2004) showed enhanced Ca^{2+} and NO_3^- in the coarse mode owing to continental Asian dust. In the East case, speciated organics were fairly well correlated with NO_3^- ($r = 0.68$ – 0.99), which has been associated with adsorption onto coarse aerosol types like dust and sea salt (e.g., Maudlin et al., 2015; Stahl et al., 2020). Nitrate was especially well correlated with Na^+ , Cl^- , and Ca^{2+} ($r = 0.98$ – 1.00), which exceeded correlations of other common inorganic ions such as SO_4^{2-} and NH_4^+ .

The vertical profiles clearly show the systematically higher TOC masses relative to the North case across roughly the same altitude range (1.3–3.3 km); however, the AMS organic and m/z 44 values (although sparse) were more comparable, which again can simply be due to the differences in what is being measured, with AMS not accounting for the supermicrometer particles types (i.e., dust and sea salt) that were likely more influential in the cloud water in the East case. However, the importance of droplet uptake of water-soluble organic gases should also be considered, as they can influence TOC mass.

Evidence of greater aging compared with the North case comes from a few ratios of interest. The $\text{Cl}^- : \text{Na}^+$ ratio for this case (1.40 ± 0.06) was lower than for the North case, suggestive of more sea salt reactivity aided by presumed aging. Furthermore, the acetate : formate ratio (1.93 ± 1.51) was less than half the value from the North case. More broadly, the overall contribution of MCAs and DCAs to TOC were very similar between the North and East cases and also the next two cases: MCA : TOC = 16.03 %–23.66 %, and DCA : TOC = 3.70 %–8.75 % (Table 3). In contrast with the North case, this category of samples had weaker interrela-

tionships between organic species, which was presumed to be due to the mixture of sources impacting this case including dust, marine particles, and likely other anthropogenic and biogenic sources over land.

4.3 Biomass Burning

The Biomass Burning (BB) category samples exhibited the highest concentrations of TOC (8342 ± 3730 ppb C) and almost every organic, with the dominant contributors to TOC (± 1 standard deviation) being formate (2178 ± 1589 ppb; 7.0 ± 4.5 %), acetate (1845 ± 1668 ppb; 8.4 ± 5.6 %), and succinate (557.0 ± 575.6 ppb; 2.4 ± 1.7 %). As acetate and formate were so abundant, the relative enhancement of MCA mass was much larger than DCA mass compared with the three other cases examined (Table 2). While the correlation matrix for this case was quite sparse in terms of significant values, owing partly to the use of only a few points ($n = 4$), TOC and K^+ were highly correlated ($r = 0.99$), which demonstrates the strong linkage between TOC and biomass burning emissions (Table S3) as also shown by others (Cook et al., 2017). For context, Desyaterik et al. (2013) reported cloud water TOC masses of 100.6 ppm C in a biomass burning air mass at Mt. Tai in eastern China that were 8 times higher than typical values in the absence of agricultural burning. Cook et al. (2017) observed significantly higher cloud water TOC masses during wildfire periods (16.6 ppm C) at Whiteface Mountain, New York, than during biogenic (2.16 ppm C) or urban (2.11 ppm C) periods.

In our BB samples, mean values of succinate (557.0 ± 575.6 ppb), glutarate (150.4 ± 82.20 ppb), and pyruvate (125.9 ± 126.1 ppb) were significantly elevated above the other cases. Stahl et al. (2020) recently showed that succinate, oxalate, and MSA were especially enhanced in aerosol samples collected in the study region during BB periods in the 2018 southwest monsoon season. Study-wide peak concentrations of succinate (1372 ppb), oxalate (1135 ppb), and MSA (24.79 ppb) were found in this case, reinforcing those findings (Stahl et al., 2020). Unlike the previous two cases, maleate was detected in BB samples (5.583 ± 6.456 ppb). Although maleate is associated with combustion sources (Kawamura and Kaplan, 1987; Rogge et al., 1993), such as from extensive ship traffic around the sampling area, other studies have shown enhancements of maleate in BB air masses (i.e., Mardi et al., 2019; Tsai et al., 2013). The percentage of mass contributing to TOC that was unaccounted for was 78.7 %, with the highest sample at 6.5 km having 95.6 % undetected, which was surprisingly large based on the prevalence of organic acids in biomass burning emissions (Reid et al., 1998). Therefore, the second hypothesis posed in this study is partly true in that the BB case exhibited much higher TOC values; however, these samples did not exhibit a greater contribution from organic acids to TOC, as the North and East cases actually had a greater contribution from such species. This highlights

the need for more attention to be paid to organic chemical speciation in clouds impacted by biomass burning emissions, as such a large portion of the TOC mass went unaccounted for in this study.

While absolute concentrations of most organics were greatly enhanced in BB, the relative contributions of individual organics within the MCA and DCA subsets of species also varied. Most notably in the MCA category, formate was greatly enhanced with a mass contribution to total MCA mass of 46.40 % versus 16.54 %– 29.09 % for other cases. In the DCA population of species, glutarate and succinate accounted for higher mass fractions in the BB case (17.15 % and 41.95 %, respectively) than in the other three cases (0.65 %– 4.02 % and 20.82 %– 38.52 %, respectively).

The $Cl^- : Na^+$ ratio was 1.30 ± 0.06 and suggestive of Cl^- depletion, which has been observed in other regions experiencing biomass burning and has been linked to high concentrations of inorganic and organic acids (Braun et al., 2017, and references therein). This is supported by how the values of MO , SO_4^{2-} , and NO_3^- were the highest in this case (Table 2). The acetate : formate ratio was 0.69 ± 0.30 , but it is unclear how effective this and other ratios are as aging indicators when biomass burning is present, especially as fuel type varies between regions. Talbot et al. (1988) and Wang et al. (2007) both report that the acetate : formate ratio is substantially larger in biomass burning samples, which is contradictory to the ratios that are reported for this case (0.32 to 1.03). This could be due to the fuel type or aging of the biomass burning plume; however, this is speculative and should be examined more extensively.

4.4 Clark

Samples in this category were collected during ascents after takeoff and descents during approaches to the airfield, which allowed for sample collection closer to the surface than the other categories (altitude range of 0.2 – 2.9 km). Clark International Airport is located within the Clark Freeport Zone, which is part of both the Pampanga and Tarlac provinces and consists of five cities and municipalities: Angeles city, Mabalacat city, Porac, Capas, and Bamban. This gives the Clark area a population of approximately $996\,000$ with a population density of ~ 3100 km $^{-2}$, which is low in comparison to the most populated city in the Philippines, Quezon City in Metro Manila, with 2.94 million people and a population density of $\sim 17\,000$ km $^{-2}$ (PSA, 2016). In addition to Metro Manila just to the southeast (~ 90 km), Clark lies between Mt. Pinatubo to the west and Mt. Arayat to the east, which are active and potentially active volcanoes, respectively.

The average TOC for this case (1181 ± 920.2 ppb C) was most similar to the East case and exhibited the most variability relative to the mean TOC value of all four cases, which we attribute to numerous sources impacting these samples, including local and regional emissions, time of day variability, local spatial variability, and number of flights.

This case exhibited the highest percentage of TOC mass that was unaccounted for by speciated organics (79.5 %), with the three largest measured contributors (± 1 standard deviation) consisting of acetate (296.7 ± 325.8 ppb; 9.6 ± 9.5 %), formate (266.1 ± 316.8 ppb; 4.8 ± 3.3 %), and oxalate (88.33 ± 103.9 ppb; 1.7 ± 1.0 %). A few notable features are mentioned that are specific to this case. This was the only case that had DMA present (6.454 ± 15.89 ppb), albeit with a low mass contribution to total TOC (0.43 ± 1.17 %). This case exhibited the highest mass fractions of maleate (3.20 ± 5.93 %) and adipate (16.05 ± 21.48 %) relative to DCA mass, suggestive of greater anthropogenic emission influence and processed aromatic compounds. DMA was only correlated with maleate ($r = 0.67$) among the organic species, which is suggestive of a similar source (Table S4). Stahl et al. (2020) showed increased aerosol concentrations of freshly emitted organics (i.e., phthalate and maleate) owing to the vast sources of combustion engines to the southeast of the Clark area. Clark is situated near a major highway that could also contribute to the high combustion sources, although commercial aircraft emissions could also play a significant role.

Because succinate peaked in concentration for this case (498.50 ppb) and back trajectories originated from Borneo and Sumatra, there may have been some influence from biomass burning (Fig. 5). The $K^+ : Na^+$ ratio was elevated (0.25) above that of sea salt (0.036) (Seinfeld and Pandis, 2016) and was even higher than the Biomass Burning case (0.15), suggestive of local and/or regional biomass burning influence. This case exhibited the highest mean $Ca^{2+} : Na^+$ ratio (0.99) and was well above the sea salt value (0.038), which we presume could be largely linked to resuspended and/or transported dust. For Metro Manila, Cruz et al. (2019) showed that resuspended dust, especially linked to vehicular traffic, is an important source of dust in the study region. Stahl et al. (2020) showed that adipate is most influenced by crustal sources in the study region and was unique among the studied organics in this work in that it exhibited a prominent peak in the supermicrometer range based on surface aerosol measurements in Metro Manila. Consistent with that work, Ca^{2+} was only correlated with adipate in the Clark samples ($r = 0.71$) among the studied organics (Table S4), adding support for how organic acids like adipate can partition to dust with the novelty here being that the signature was observed in cloud water.

5 Conclusions

This work analyzed 159 cloud water samples collected over a 2-month period as part of the CAMP²Ex airborne campaign around the Philippines. TOC and a total of 11 organic compounds comprised of 4 MCAs (glycolate, acetate, formate, and pyruvate), 5 DCAs (glutarate, adipate, succinate, maleate, and oxalate), MSA, and DMA were measured. The

measured organics were then compared to TOC to determine the percentage of organic species measured compared with the total organic composition. Notable results are summarized below including responses to the two hypotheses proposed at the end of Sect. 1.

TOC masses ranged widely between 0.018 and 13.66 ppmC from 0.2 to 6.8 km, with a mean value of 0.902 ppmC. The contribution (in C mass) of the 11 measured species to total TOC was on average 30 %. Using a conversion factor of 1.8 for organic matter relative to organic carbon, the mean amount of total organic matter (TOM) accounted for by our measured 11 species was 46.4 %. Furthermore, the mean contribution of TOM and speciated organics to total mass (inorganics + organics) was 30.7 % (maximum of 95.1 %) and 10.3 % (maximum of 57.6 %), respectively. The mean ratio of inorganic to TOM was 5.8. The study's first hypothesis that the measured organic species account for a higher mass fraction relative to total mass compared with surface layer aerosol measurements over Luzon, excluding black carbon (~ 1.3 %) (Stahl et al., 2020) holds true. This is likely owing to more processed air masses aloft and the reduced influence of black carbon that is so abundant in areas like Metro Manila (Cruz et al., 2019; Hilario et al., 2020a). The uptake of water-soluble gases can also attribute to greater organic mass contributions.

In terms of the chemical profile of the speciated organics, the order in decreasing contribution of C mass relative to TOC was as follows (± 1 standard deviation): acetate (14.7 ± 20.5 %), formate (5.4 ± 9.3 %), oxalate (2.8 ± 4.3 %), DMA (1.7 ± 6.3 %), succinate (1.6 ± 2.4 %), pyruvate (1.3 ± 4.5 %), glycolate (1.3 ± 3.7 %), adipate (1.0 ± 3.6 %), MSA (0.1 ± 0.1 %), glutarate (0.1 ± 0.2 %), and maleate ($< 0.1 \pm 0.1$ %). Approximately 70.0 % of TOC went unaccounted for, highlighting the complexity and difficulty of organic speciation in the study region, with this value being fairly similar to other regions (Benedict et al., 2012; Boris et al., 2016, 2018; Herckes et al., 2002; Raja et al., 2008). Monocarboxylic acids dominated the speciated organic mass (~ 75 %) and were about 4 times more abundant than dicarboxylic acids, which is suggestive of a higher abundance of gaseous species and precursors. It should also be noted that MCAs have a higher volatility than DCAs, which could contribute to the higher organic mass. Additionally, the MCAs measured in this study were predominately short-chain organics that have naturally higher volatilities (Chebbi and Carlier, 1996; Wang et al., 2007).

Vertical profiles of TOC revealed higher concentrations in the bottom 2 km with a reduction above that. Samples impacted by biomass burning emissions were substantially enhanced in TOC and most speciated organic masses, ranging in altitude from as low as 1.3 km to as high as 6.5 km. While vertical profiles of AMS organic and m/z 44 mass concentrations qualitatively resembled that of TOC with reductions above 2 km, the vertical behavior of chemical ratios relevant to the composition of the cloud (ratio of C mass

from measured organics to TOC) and aerosol organics (f_{44}) did not reveal any clear trend. For both non-BB and BB samples, monocarboxylic acids uniformly dominated C mass with $\sim 75\%$ of TOC mass unaccounted for across the range of altitudes studied.

The second hypothesis in this study proved to be partly true, as clouds impacted by biomass burning exhibited markedly higher TOC values (4.974–13.66 ppm C) and masses of most of the other species detected compared with the other three categories of samples in Sect. 4 (North, East, and Clark). However, the part of the hypothesis about speciated organic acids contributing more to BB samples did not hold true, as total measured organics accounted on average for 21.25% of TOC, which was lower than two of the other categories of samples (North, 26.72%, and East, 30.61%). Interestingly, the highest BB sample (6.5 km) had 95.6% of the C mass unaccounted for by speciated organics. This highlights the need for increased attention to be paid to organic speciation in clouds impacted by biomass burning.

Four categories of samples with different air mass history characteristics were compared, revealing a few notable features: (i) while speciated concentrations and TOC masses varied considerably between the four cases, the contributions of MCAs and DCAs (based on C mass) to TOC were remarkably similar with little variation (MCA : TOC = 16.03%–23.66% and DCA : TOC = 3.70%–8.75%); (ii) dust and sea salt tracer species were strongly correlated with most of the speciated organics for the North and East cases, suggestive of interactions between such species and coarse aerosol surfaces, as supported by past work (Stahl et al., 2020; Sullivan and Prather, 2007); (iii) for samples with limited aging (North case) based on selected chemical ratio values, adipate was more abundant and was negatively correlated with smaller carboxylic acids; (iv) BB samples exhibited the highest TOC concentrations (8342 ± 3730 ppb C) as well as significantly elevated levels of individual organics such as acetate, formate, succinate, glutarate, pyruvate, oxalate, and MSA; and (v) the Clark case had a higher variability in TOC (1181 ± 920.2 ppb C) than the North and East cases, presumably owing to a greater mix of influential sources such as fresh anthropogenic emissions (e.g., enhanced maleate) as well as the transport of biomass burning plumes from Borneo and Sumatra (e.g., enhanced succinate), dust, and spatial and temporal variances across different flights. Related to dust, Ca^{2+} was only correlated with adipate in the Clark samples, which is consistent with a recent study in Metro Manila (Stahl et al., 2020) showing that adipate uniquely exhibits a prominent supermicrometer peak among organic acids, attributed to interactions with dust.

Data availability. All data used can be found on the NASA data repository at <https://doi.org/10.5067/Suborbital/CAMP2EX2018/DATA001> (last access: 10 August 2021). Specifically, the two datasets used in this

paper can be found at https://doi.org/10.5067/Airborne/CAMP2Ex_Cloud_AircraftInSitu_P3_Data_1 (NASA/LARC/SD/ASDC, 2020a) and https://doi.org/10.5067/Airborne/CAMP2Ex_Aerosol_AircraftInSitu_P3_Data_1 (NASA/LARC/SD/ASDC, 2020b).

Supplement. The supplement related to this article is available online at: <https://doi.org/10.5194/acp-21-14109-2021-supplement>.

Author contributions. EC, RAB, CS, ABM, and AS designed the experiment. All coauthors carried out various aspects of the data collection. EC, CS, and AS conducted analysis and interpretation of the data. CS and AS prepared the manuscript with contributions from the coauthors.

Competing interests. The contact author has declared that neither they nor their coauthors have any competing interests.

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Special issue statement. This article is part of the special issue "Cloud, Aerosol and Monsoon Processes Philippines Experiment (CAMP2Ex) (ACP/AMT inter-journal SI)". It is not associated with a conference.

Acknowledgements. The authors acknowledge support from NASA (grant no. 80NSSC18K0148) in the framework of the NASA CAMP2Ex project. Rachel A. Braun acknowledges support from the ARCS Foundation, Melliza Templonuevo Cruz was supported by the Philippine Department of Science and Technology's ASTHRD program, and Alexander B. MacDonald acknowledges support from the Mexican National Council for Science and Technology (CONACYT).

Financial support. This research has been supported by the National Aeronautics and Space Administration (grant no. 80NSSC18K0148).

Review statement. This paper was edited by Manvendra K. Dubey and reviewed by two anonymous referees.

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