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## Dimethylamine in cloud water: a case study over the northwest Atlantic Ocean†

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This study analyzes characteristics of an important alkyl amine species, dimethylamine (DMA), in cloud water over the northwest Atlantic. Data were gathered from the winter and summer 2020 deployments of the Aerosol Cloud meTeorology Interactions oVer the western ATLantic Experiment (ACTIVATE) on board the HU-25 Falcon. Thirty-eight out of 98 samples exhibited DMA above detection limits, with the overwhelming majority in the winter season (33, 52% of winter samples) compared to summer (5, 14% of summer samples). Higher levels of DMA were observed in the winter, especially during cold air outbreaks (CAOs), which was also the case for  $\text{NO}_3^-$ ,  $\text{NH}_4^+$ , and non sea salt- $\text{SO}_4^{2-}$ . This is in part due to a combination of low temperatures and offshore flow enhanced with continental pollutants such as from agriculture, industry, urban activity, and biomass burning. Unlike the inorganic acidic anions, oxalate was significantly correlated to DMA in summer and the winter in both CAO and non-CAO conditions, with a presumed reason being biomass burning supported by the consistent correlation between DMA and  $\text{nss-K}^+$  in each season. ACTIVATE data are compared to a cloud water dataset from the northeast Pacific, with the latter exhibiting much higher DMA levels due possibly to more abundant ocean biological emissions. The seasonal differences and enhancement in DMA during CAO conditions relative to non-CAO winter days motivates continued research into the partitioning behavior of DMA and its sources as amines play an important role in carbon and nitrogen cycles in the marine environment.

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### Environmental significance

Amines are ubiquitous organic bases that play a crucial role in several atmospheric chemical processes, such as new particle formation, formation of low-volatility amine salts in aerosol particles, and aqueous chemistry. Their presence in particles can impact hygroscopic properties and lead to adverse health effects. While research into the nature of these species in the gas and aerosol phases has intensified in recent decades, much less is known about their nature in cloud water, which offers a large aqueous reservoir for dissolution of gaseous amines while also containing amines through activation of cloud condensation nuclei containing amines. This study exploits a unique airborne dataset over the northwest Atlantic across different seasons to examine the concentration characteristics of one of the most abundant amines, dimethylamine. The results show that this species is much more common in the winter relative to the summer and that conditions associated with cold air outbreaks lead to especially higher concentrations. This work provides additional support for how biomass burning may be an important source of amines in cloud water.

## 1. Introduction

Characterizing cloud water composition is insightful in providing clues about air mass types impacting clouds.<sup>1–4</sup> Solute composition in cloud water impacts pH and thus processes dependent on acidity,<sup>5</sup> such as aqueous phase reactions.<sup>6,7</sup> Cloud water species can also impact gas partitioning<sup>8</sup> and affect ecosystems due to wet deposition.<sup>9,10</sup> A potentially important yet understudied portion of species in cloud water includes organonitrogen species such as amines.

Amines are ubiquitous organic bases that play a crucial role in several atmospheric chemical processes.<sup>11</sup> They participate in

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new particle formation<sup>12–14</sup> and growth processes,<sup>15,16</sup> in addition to formation of low-volatility amine salts in aerosol particles.<sup>17–20</sup> Their impact extends from altering hygroscopic properties<sup>21–23</sup> to adverse health effects.<sup>24,25</sup> While research into the gas-aerosol partitioning of amines, reactivity with existing inorganic salts,<sup>26,27</sup> and their properties in the aerosol phase has intensified in recent decades,<sup>28,29</sup> much less is known about their nature in cloud water, which offers a large aqueous reservoir for dissolution of gaseous amines in addition to the activation of cloud condensation nuclei (CCN) containing amines.<sup>30–32</sup>

Dimethylamine (DMA) is one of the more easily detected alkyl amines in aerosol and cloud water samples due to its solubility and abundance relative to many other amine species.<sup>28,32–35</sup> DMA has a range of sources, including emissions from industry,<sup>35</sup> biomass burning,<sup>36,37</sup> vehicle emissions,<sup>38</sup> animal husbandry,<sup>39,40</sup> and the ocean.<sup>41</sup> Past work focused on DMA and other alkyl amines in the gas and aerosol phases reported concentrations in different regions and examined interrelationships with the major competing base species (ammonium,  $\text{NH}_4^+$ ) and acidic species that it can form salts with such as inorganic<sup>17,33,42–45</sup> and organic acids.<sup>15,16,46,47</sup> To our knowledge, one study focused on DMA specifically in cloud water with airborne measurements.<sup>32</sup> This study shows that DMA concentrations are enhanced closest to the California coast (relative to farther offshore) potentially due to marine biogenic emissions and that its concentrations increased significantly when impacted by biomass burning due to presumed dissolution of DMA into cloud water. Stahl *et al.*<sup>48</sup> analyzed cloud water collected during the Clouds, Aerosols and Monsoon Processes – Philippines Experiment (CAMP<sup>2</sup>Ex) airborne campaign and reported that DMA, on average, accounted for 1.7% of total organic carbon during flights around the Philippines. Crosbie *et al.*<sup>49</sup> analyzed three case flights of developed maritime convection from that same campaign and reported no measurable DMA despite two cases being moderately polluted and the third being influenced by smoke. Others have examined DMA in cloud water using surface-based measurements, such as at mountain Schmücke in central Germany, showing enhanced DMA levels owing to its high solubility.<sup>50</sup> DMA has been detected in cloud droplet residual particles, further confirming a role for it in clouds.<sup>51</sup> Knowledge gaps pertaining to DMA include a full understanding of its partitioning behavior between gas, solid, and aqueous media and the dependence of these processes on factors such as pH, humidity, and temperature.<sup>11,28</sup>

We leverage a unique cloud water dataset collected in recent deployments of the Aerosol Cloud meTeorology Interactions oVer the western ATlantic Experiment (ACTIVATE)<sup>52</sup> to study the seasonal variation of DMA levels in cloud water, relationships between DMA and other species in cloud water, and possible influential sources for DMA. Results from ACTIVATE are then directly compared to the Youn *et al.*<sup>32</sup> study that investigated DMA in cloud water off the California coast. We focus on DMA as that was the only alkyl amine detected in ACTIVATE cloud water samples.

## 2. Data

### 2.1 ACTIVATE campaign description

The ACTIVATE campaign comprised two aircraft (HU-25 Falcon and King Air) flying in vertical coordination over the northwest Atlantic to investigate aerosol-cloud-meteorology interactions. *In situ* data were collected by the Falcon, which flew below, in, and just above boundary layer clouds. The King Air flew at  $\sim 9$  km and conducted remote sensing of the atmospheric column underneath the aircraft and launched dropsondes. Six deployments were conducted between 2020 and 2022, with separate winter and summer deployments each year. This study focuses on cloud water samples collected by the Falcon during the first two deployments in 2020: winter (14 February – 12 March 2020) and summer (13 August – 30 September 2020). We further divided the winter flights into two categories to isolate data on cold air outbreak (CAO) conditions, as recent work shows unique conditions in those flights affecting aerosol particles and clouds.<sup>53–56</sup> We used the same methods in recent ACTIVATE studies<sup>53,57</sup> to distinguish CAO days using a combination of Visible Infrared Imaging Radiometer Suite (VIIRS) imagery (NASA Worldview), flight reports, weather forecast data, and King Air dropsonde data.

The Falcon flight path typically constituted a series of legs below, in, and above clouds.<sup>58</sup> The most relevant legs to this study are the “below cloud base (BCB)” legs from which we used meteorological data, and the “above cloud base (ACB)” and “below cloud top (BCT)” legs where cloud water samples were gathered. Cloud water samples might have also been collected during slant ascents and descents between the legs whenever the aircraft was in cloud.

### 2.2 Falcon data

**2.2.1 Cloud water.** The Falcon collected cloud water samples using the Axial Cyclone Cloud-water Collector AC3.<sup>59</sup> The AC3 inertially separates droplets from the air stream with a collection efficiency  $>60\%$  above diameters of  $20\ \mu\text{m}$ . The median duration of collection per sample was 5.0 minutes (25th/75th percentile = 2.5/9.4 minutes). Samples collected in vials were analyzed post-flight *via* ion chromatography (IC) using documented methods.<sup>60,61</sup> Each sample ran for 30 minutes, which included a 5 minute equilibrium time between samples at a flow rate of  $0.4\ \text{mL}\ \text{min}^{-1}$ . Anion analysis relied on a multi-step gradient method and a Dionex IonPac ASH11-HC ( $2\ \text{mm} \times 250\ \text{mm}$ ) column, AERS 500e suppressor, and potassium hydroxide (KOH) as the eluent. For cation analysis, an isocratic method was used with methanesulfonic acid (MSA) as the eluent, a Dionex IonPac CS12A ( $2\ \text{mm} \times 250\ \text{mm}$ ) column, and a CERS 500e suppressor. The subset of detected species relevant to this study includes DMA, sodium ( $\text{Na}^+$ ), methanesulfonate (MSA), ammonium ( $\text{NH}_4^+$ ), nitrate ( $\text{NO}_3^-$ ), oxalate, and non-sea salt forms of sulfate ( $\text{nss-SO}_4^{2-}$ ), calcium ( $\text{nss-Ca}^{2+}$ ), and potassium ( $\text{nss-K}^+$ ). The nss parameters were calculated using knowledge of  $\text{Na}^+$  and concentrations in each sample of either  $\text{SO}_4^{2-}$ ,  $\text{Ca}^{2+}$ , or  $\text{K}^+$  and the ratio of  $\text{Na}^+$  to each of those three species in natural sea water<sup>62</sup> with the



assumption that all  $\text{Na}^+$  is from sea salt. The limits of detection can be found in Table S1.† Air-equivalent solute concentrations are reported following the same method described in Gonzales *et al.*<sup>61</sup> Briefly, aqueous concentrations from the IC are multiplied by the mean liquid water content (LWC) in clouds (*i.e.*, when  $\text{LWC} \geq 0.02 \text{ g m}^{-3}$ ) during the period of sample collection divided by the density of liquid water. LWC data are used from the Fast Cloud Droplet Probe (FCDP; 3  $\mu\text{m}$ –50  $\mu\text{m}$ ; Stratton Park Engineering Company Incorporated (SPEC Inc.)).<sup>63,64</sup>

Concentrations of a suite of elements were measured using an inductively coupled plasma mass spectrometer (ICP-MS; Agilent 7800 Series). Samples were analyzed in general purpose plasma mode (helium) with an integration time of 0.5 seconds for all elements. A semi-quantitative analysis was performed using a two-point calibration (calibration blank and 10 ppb semiquantitative standard). Internal standards of indium (I) and bismuth (Bi) were added to all standards and samples *via* on-line addition at a final concentration of 50 ppb. Elements discussed in this work include: Al, V, Fe, Ni, Cu, Zn, As, Cd, Pb. We additionally measured pH of the cloud water samples using an Orion Star™ A211 pH meter coupled to an Orion™ 8103BNUWP ROSS Ultra™ pH electrode (precision of 0.01). Two-point calibrations (pHs 4 and 7) were conducted for the pH measurements.

Of the 98 cloud water samples gathered during the 40 research flights in 2020, 38 exhibited DMA concentrations above the detection limit. The median cloud water sample volumes collected were 3.9 mL and 1.9 mL for the winter and summer, respectively. Although this might seem like a limited number of samples, there is significant difficulty in collecting such samples and DMA is generally at low concentration as compared to species more readily measured by IC such as  $\text{Na}^+$  and  $\text{SO}_4^{2-}$ . The low DMA concentrations do not diminish the importance of characterizing this species as it is a marker representative of alkyl amines that are poorly understood in the context of aerosol–cloud interactions and general characterization of cloud water. There is significant scarcity in cloud water DMA concentrations reported in the literature motivating the importance of learning about this sample set.

**2.2.2 Other Falcon data.** Meteorological data used in this study include static air temperature (Rosemount 102 sensor) and water vapor mixing ratio (WVMR) and relative humidity (RH) derived from the Diode Laser Hygrometer (DLH).<sup>65</sup> Carbon monoxide (CO) data were obtained from a Picarro G2401 gas concentration analyzer.<sup>66</sup> Submicrometer aerosol composition data were measured by an Aerodyne High-Resolution Time-of-Flight Aerosol Mass Spectrometer (AMS)<sup>67</sup> with organic mass concentrations of most relevance to this study. Those data were collected in a 1 Hz fast-MS mode and averaged up to 25 second resolution. All other instrument data from this section are used at 1 second resolution.

### 2.3 Supporting datasets

Air mass transport pathways leading up to the point of cloud water samples were identified using the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPPLIT)<sup>68,69</sup> model. Global

Data Assimilation System (GDAS) meteorological data ( $0.5^\circ \times 0.5^\circ$ ) were used to acquire 72 and 240 hour back trajectories with 6 hour time resolution per trajectory. Speciated optical depth data over North America and the northwest Atlantic are used from the Navy Aerosol Analysis and Prediction System (NAAPS).<sup>70,71</sup> We use NAAPS data at  $1^\circ \times 1^\circ$  spatial resolution and 6 hour temporal resolution with a focus on smoke optical depths.

To aid in aerosol type identification during one of the case flights presented in this work, we used data from the nadir-viewing high spectral resolution lidar (NASA Langley Research Center HSRL-2) installed on the King Air. We used these data in combination with *in situ* and model datasets to infer the presence of biomass burning aerosols above the planetary boundary layer. Readers are referred to Burton *et al.*<sup>72</sup> for information about the HSRL-2 operational details.

To examine the potential role of ocean biogenic emissions as a source of DMA in cloud water,<sup>18</sup> we obtained Level-3 (8 day average, 4 km resolution) MODIS-Aqua data for chlorophyll *a* ( $\text{mg m}^{-3}$ ) and the particulate backscattering coefficient ( $b_{\text{bp}}$ ; 443 nm;  $\text{m}^{-1}$ ) that encompassed the winter and summer 2020 ACTIVATE flight domains. The chlorophyll *a* product represents the concentration of the photosynthetic pigment in the upper ocean derived from passive remote sensing of ocean color.<sup>73</sup> The particulate backscattering coefficient describes the radiant intensity scattered into the backwards direction by particles in the water column, which can be used as a proxy for the particulate organic carbon concentration in waters where scattering is dominated by organic particles (*e.g.*, open ocean regions with low concentrations of inorganic particles).<sup>74</sup> For two case studies presented in this work where high-level clouds prevented passive ocean color remote sensing retrievals, HSRL-2 retrievals of  $b_{\text{bp}}$  (532 nm; 10 m depth) measured from the King Air were used to supplement the seasonal patterns highlighted in the passive imagery.<sup>75</sup> We caution that chlorophyll *a* and  $b_{\text{bp}}$  are not perfect proxies for upper ocean biomass or the amount of organic carbon enrichment in marine-emitted aerosol particles;<sup>76,77</sup> however, in the face of these limitations there have been links shown between amine concentrations over the ocean and at least chlorophyll *a*.<sup>32,33,78</sup>

Lastly, to put the results from ACTIVATE in perspective, data are used from another airborne campaign with documented DMA concentrations in cloud water,<sup>32</sup> which was the Nucleation in California Experiment (NiCE; July–August 2013) based in Marina, California with 23 flights over the northeast Pacific.<sup>79–81</sup> The Center for Interdisciplinary Remotely-Piloted Aircraft Studies (CIRPAS) Twin Otter payload in that campaign is summarized elsewhere,<sup>82</sup> and of most relevance here is the modified Mohnen slotted-rod collector<sup>83</sup> which obtained 119 samples that were analyzed with the same IC system as the ACTIVATE samples.<sup>32</sup> Past work has intercompared the AC3 and slotted-rod collector on the CIRPAS Twin Otter and showed that campaign-wide mean concentrations of trace species exhibited good agreement between the two collectors during the 2016 Fog and Stratocumulus Evolution (FASE) campaign over the northeast Pacific.<sup>59</sup>



### 3. Results and discussion

#### 3.1 DMA seasonal concentrations

We take advantage of ACTIVATE's dataset for two different seasons to study interseasonal DMA concentration trends. Others have shown seasonal differences in amine behavior, such as how higher atmospheric water and particle acidity in the summer (*versus* fall) can enhance aminium salt formation in particles.<sup>46</sup> Cloud water concentrations of DMA arise mainly from this species being associated with the CCN that nucleate into cloud drops or from dissolution of gaseous amines into cloud water. Furthermore, processes such as hydrolysis in the aqueous phase can convert amines into amides and *vice versa*.<sup>84</sup>

Past work over the northeast Pacific suggested that DMA levels were significantly larger in cloud water compared to aerosol measurements with a particle-into-liquid sampler around and in clouds, with the speculated reason being volatilization of the DMA upon droplet evaporation,<sup>32</sup> much like what has been observed for nitrate.<sup>81</sup> Amines were detected in higher relative amounts in cloud residues relative to outside of clouds in a different study with the reasoning being their high solubility.<sup>50</sup>

The number of samples with detectable DMA was distributed as follows relative to the total number of cloud water samples collected per category as a percentage (locations in Fig. 1): winter non-CAO = 16 out of 24 (66.7%); winter CAO = 17 out of 39 (43.6%); summer = 5 out of 35 (14.3%). The median/mean/

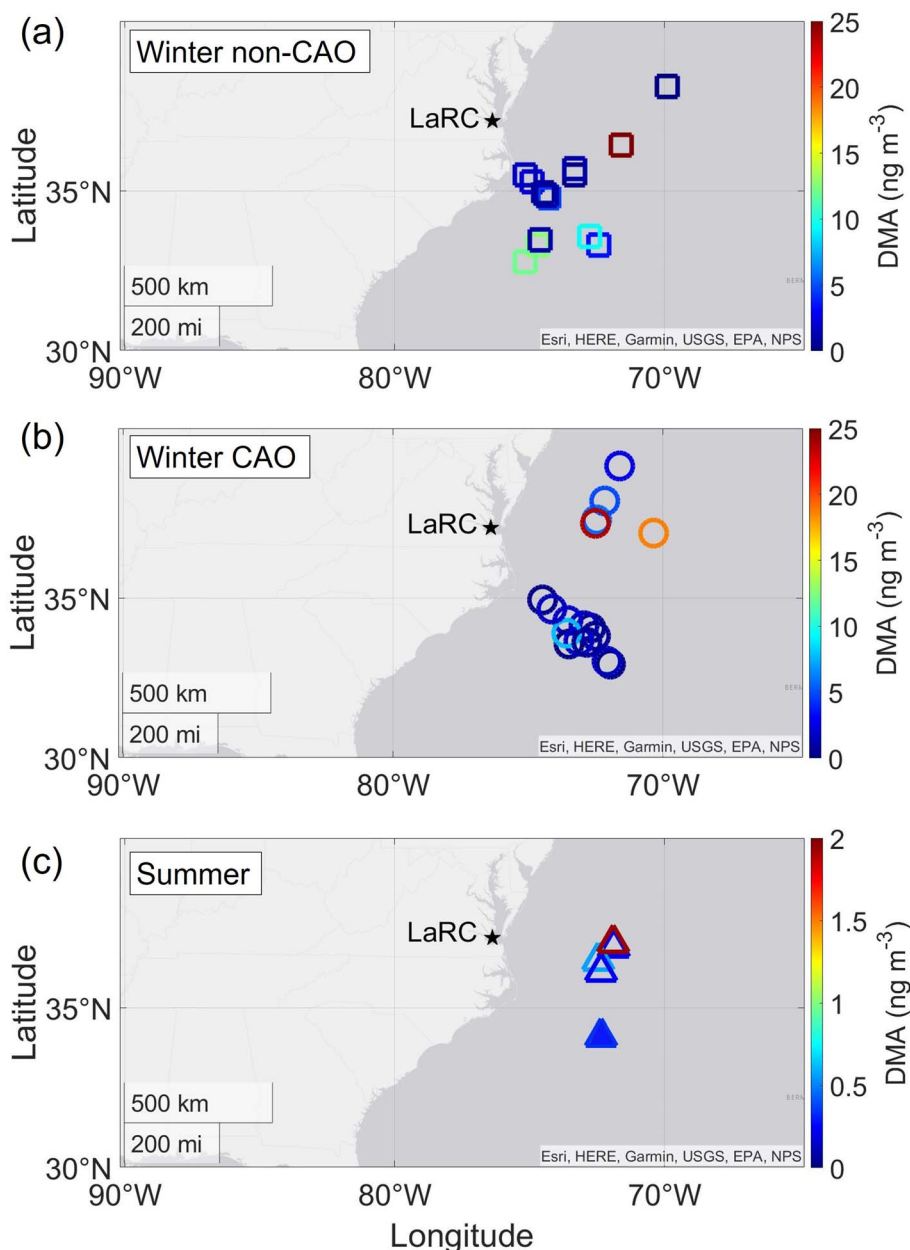


Fig. 1 Cloud water DMA air-equivalent concentration ( $\text{ng m}^{-3}$ ) spatial maps for (a) winter non-CAO, (b) winter CAO, and (c) summer periods in 2020. The filled marker in panel (c) represents a sample influenced by biomass burning.



**Table 1** Median values and sample numbers for the following parameters categorized into winter non-CAO, winter CAO, and summer conditions: cloud water pH and air-equivalent solute concentrations, cloud water sample altitudes, the cloud water DMA :  $\text{NH}_4^+$  molar ratio, relevant meteorological variable values from below cloud base (BCB) legs, and ocean biological parameter values from MODIS-Aqua

	Winter non-CAO		Winter CAO		Summer	
	Median	<i>n</i>	Median	<i>n</i>	Median	<i>n</i>
Sample altitude (m)	1397	16	1403	17	1668	5
Cloud water pH	4.54	11	5.21	16	5.91	6
$\text{NH}_4^+$ ( $\text{ng m}^{-3}$ )	146	24	318	39	29	29
DMA ( $\text{ng m}^{-3}$ )	1.61	16	4.97	17	0.25	5
$\text{NO}_3^-$ ( $\text{ng m}^{-3}$ )	1813	24	2960	39	507	33
nss- $\text{SO}_4^{2-}$ ( $\text{ng m}^{-3}$ )	147	24	164	39	25	35
DMA : $\text{NH}_4^+$ (molar basis)	0.003	16	0.002	17	0.001	5
BCB carbon monoxide (ppm)	0.14	4015	0.14	4215	0.12	7041
BCB air temperature ( $^\circ\text{C}$ )	4.52	6521	1.00	7258	19.75	9737
BCB relative humidity (%)	86	6521	84	7258	88	9737
BCB water vapor mixing ratio (ppmv)	7834	6521	6151	7258	20 938	9737
Chlorophyll <i>a</i> ( $\text{mg m}^{-3}$ )	0.84	5	0.84	5	0.49	7
Particulate backscattering coefficient ( $b_{\text{bp}}$ ; 443 nm; $\text{m}^{-1}$ )	0.003	5	0.003	5	0.002	7

standard deviation/25th percentile/75th percentile levels ( $\text{ng m}^{-3}$ ) of DMA in each category were as follows: winter non-CAO = 1.61/3.91/6.11/0.64/3.45; winter CAO = 4.97/7.39/7.86/1.51/7.79; summer = 0.25/0.63/0.69/0.25/0.34. These results indicate markedly less DMA abundance in cloud water in the summer. During the winter, DMA concentrations were considerably higher in CAO conditions (when comparing samples with detectable DMA), with over a factor of 3 enhancement in DMA's median concentration relative to non-CAO conditions, which exceeded enhancements of all other species when comparing those two categories (Table 1).

For context, median cloud water DMA concentrations off the California coast<sup>32</sup> were  $5.8 \text{ ng m}^{-3}$  ( $n = 42$ ) and  $10.1 \text{ ng m}^{-3}$  ( $n = 11$ ) for samples without and with biomass burning influence, respectively. In terms of aerosol concentrations over the North Atlantic (West Irish Coast), median submicrometer levels during periods of low ( $n = 7$ ) and high ( $n = 5$ ) biological activity were 1 and  $10 \text{ ng m}^{-3}$ , respectively.<sup>18</sup> Median DMA levels in the submicrometer range (0.14–0.42  $\mu\text{m}$ ) at Cape Verde were as follows: May 2007 =  $0.21 \text{ ng m}^{-3}$  ( $n = 11$ ); June 2007 =  $0.21 \text{ ng m}^{-3}$  ( $n = 10$ ); December 2007 =  $0.54 \text{ ng m}^{-3}$  ( $n = 24$ ). Reported DMA concentrations over the northwest Atlantic during ACTIVATE flights and those archived from the eastern North Atlantic are considerably lower than those over the northeast Pacific.

### 3.2 Seasonal variability in related variables

Table 1 examines median values of different variables for comparison with DMA. The decreasing order of median DMA concentrations from winter CAO to winter non-CAO and summer is similar to the following three species among the various ions and elements examined:  $\text{NH}_4^+$ ,  $\text{NO}_3^-$ , nss- $\text{SO}_4^{2-}$ . Factors affecting DMA concentration variability include air mass source, temperature, acidity, and RH.<sup>20,33,40,45,46</sup> Although causal evidence is not provided, we compare seasonal trends in these factors *versus* trends in DMA concentrations. It is first important to note that there was not a significant difference ( $p >$

0.05; Mann–Whitney test) between the altitude that cloud water samples were collected for each of the three categories based on Table 1 values.

**3.2.1 Cloud water pH.** Cloud water was least acidic during the summer (5.91) as compared to winter (CAO = 5.21; non-CAO = 4.54), with the median values being statistically different ( $p < 0.05$ ; Mann–Whitney test). Rehbein *et al.*<sup>85</sup> showed that trimethylamine partitions better onto pre-existing particles during periods of acidic cloud processing. Total Henry's law constants of amine bases increase substantially with decreasing pH promoting greater uptake of amines from the gas phase.<sup>28</sup> The trend of more acidic cloud water and higher DMA levels in the winter periods in the study region is consistent with elevated DMA levels at higher acidity in the literature.

**3.2.2 Temperature and water vapor.** We next examined meteorological parameters measured in below cloud base (BCB) legs to assess why summertime concentrations of DMA,  $\text{NH}_4^+$ ,  $\text{NO}_3^-$ , and nss- $\text{SO}_4^{2-}$  are lowest and why in winter, their concentrations are highest in CAO conditions (when detected). A statistically significant difference was observed between the medians ( $p < 0.05$ ; Mann–Whitney test) of all three categories for all the variables in Table 1, with the only exceptions being: summer *versus* winter non-CAO for CO; sample altitude; DMA :  $\text{NH}_4^+$  molar ratio between the two winter seasons; chlorophyll *a* and particulate backscattering coefficient between summer and winter due to insufficient sample size.

Higher air temperature and water vapor mixing ratio were observed during summer over the study area, while the lowest values were in winter CAO conditions. As one source of DMA in cloud water is activation of CCN containing DMA, it important to note that there is a thermodynamic preference amines (similar to  $\text{NH}_4^+$ ,  $\text{NO}_3^-$ ) have in the aerosol phase for lower temperatures owing to their semi-volatile nature.<sup>36</sup> Moreover, lower temperatures help gaseous amines partition more effectively into the aqueous phase.<sup>28</sup> While RH showed little difference between the three temporal categories owing to a large



extent to its dependence on temperature, the water vapor mixing ratio was considerably lower in winter. It is unclear how important that factor was *versus* the lower ambient temperatures to lead to higher cloud water DMA concentrations. In contrast to ACTIVATE and other regions like the southeast U.S.,<sup>36</sup> data from the semi-arid southwestern U.S. revealed higher particulate DMA levels with higher temperature and water vapor mixing ratio with the speculated cause being more sulfate available for ammonium salt formation, more biogenic volatile organic compound emissions, and more aerosol-phase water to promote equilibrium partitioning.<sup>86</sup> Certainly, the range of results among regions warrants continued research to understand the sensitivity of DMA partitioning behavior to both temperature and various proxies of humidity such as water

vapor mixing ratio and RH. Temperature and humidity variables could potentially co-vary with other factors that may be influential for DMA concentrations, which is hard to decipher with the current dataset.

**3.2.3 Atmospheric circulation and carbon monoxide.** Fig. 2 shows 72 hour HYSPLIT air mass trajectories leading up to where cloud water samples were detected. The range of DMA sources have been noted already, but here we list relevant sources in the specific study region. There is great potential for marine biogenic emissions as the flight data were over the northwest Atlantic. But there are also agricultural sources such as dairy farms and poultry operations in Pennsylvania, Delaware, Maryland, and Virginia that are major emitters of  $\text{NH}_3$ <sup>87</sup> and presumably could be as well for amines. There are extensive

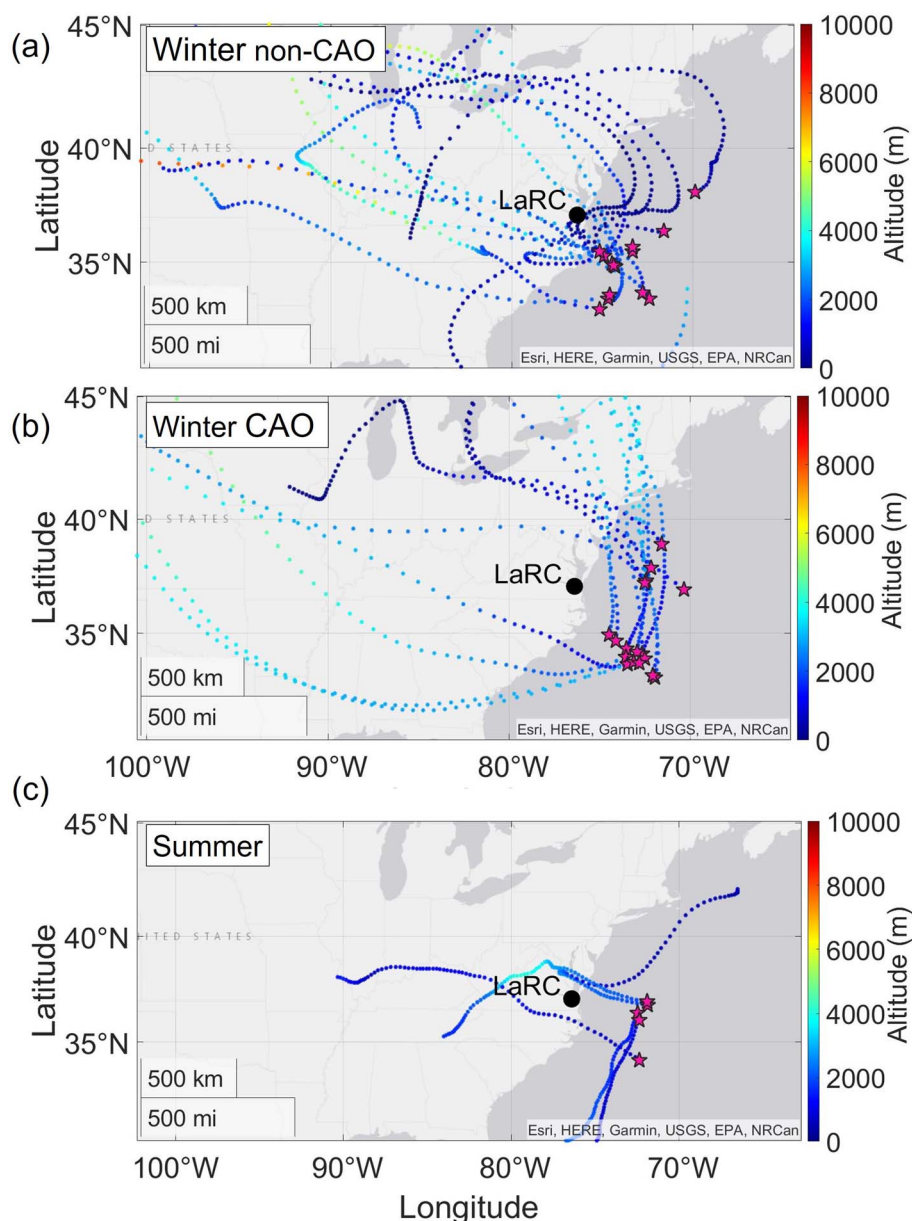


Fig. 2 HYSPLIT 72 hour back trajectories corresponding to the cloud water samples marked by stars for (a) winter non-CAO, (b) winter CAO, and (c) summer. Color bar shows the altitude for each back trajectory every 6 hours.



industrial operations along the U.S. east coast along with major populated areas leading to a wide range of anthropogenic emissions.<sup>56</sup> Biomass burning is common throughout the year, ranging from residential wood burning along the U.S. east coast, especially in winter<sup>88</sup> to transported smoke plumes from wildfires and agricultural burning.<sup>89</sup>

As expected, trajectories were mainly from over the U.S. east coast for the two winter categories without much discernible difference. These air masses were thus significantly impacted by continental emissions, including those from anthropogenic sources, agricultural activity, and likely biomass burning.<sup>88,90</sup> In summer, the trajectories were different (albeit based on five samples) and characteristic of the summertime atmospheric circulation with boundary layer winds moving north along the coast;<sup>91</sup> thus, these air masses are presumed to have had less continental influence than the winter categories. A higher median altitude was calculated for the total 72 hour duration of back trajectories in winter CAO conditions (2951 m) as compared to winter non-CAO (1888 m), with the values being statistically different ( $p < 0.05$ ; Mann-Whitney test). Higher altitudes coming from the north during the winter CAO are consistent with colder air, which as noted in Section 3.2.2 helps promote more partitioning of DMA into cloud water.

Carbon monoxide is a tracer for anthropogenic emissions. It has been shown to be higher in winter 2020 ACTIVATE flights

compared to summer and thus possibly helpful in explaining why there is more prevalent nucleation (*e.g.*, more precursor concentrations such as  $\text{SO}_2$ ).<sup>53</sup> Table 1 reinforces that CO is higher in winter, which is consistent with air mass trajectories being aligned with offshore flow. There was not a major difference in median CO levels between the two winter categories ( $\sim 0.14$  ppm), even though statistically, they were different (winter non-CAO > winter CAO). Therefore, more continental influence in the winter likely contributed to higher DMA levels. However, the relative importance of air mass source origin *versus* other factors such as temperature cannot be teased out with this dataset.

A few noteworthy features exist when comparing Fig. 1 and 2. In winter non-CAO conditions, the samples with the highest ( $>10$  ng  $\text{m}^{-3}$ ) are linked to trajectories that traveled over the southeast U.S., which has extensive agricultural burning during the winter months, such as with burning related to the sugarcane pre-harvesting season.<sup>92–95</sup> The samples with notably high DMA ( $>15$  ng  $\text{m}^{-3}$ ) in winter CAO conditions near  $37^\circ\text{N}$  were closest to land based on trajectory paths (mainly northerly) and thus least impacted by processes reducing pollutant concentrations during transport such as dilution and wet scavenging that commonly occurs offshore during CAO events which have high cloud fraction.<sup>96,97</sup> As will be shown later, one DMA sample in the summer (Fig. 1c) was linked to transported biomass burning.

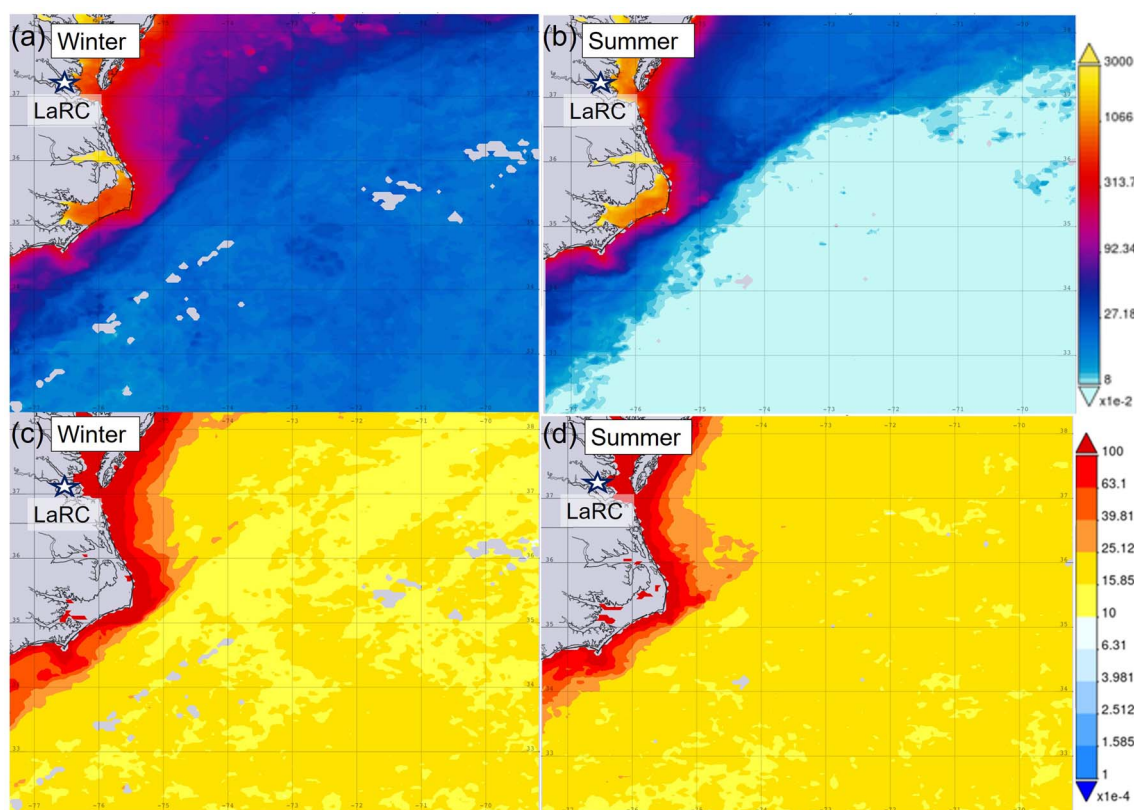


Fig. 3 Averaged MODIS-Aqua satellite images for the ACTIVATE region (<https://www.oceancolor.gsfc.nasa.gov/>) coinciding with (a and c) winter (14 February 2020–12 March 2020) and (b and d) summer (13 August 2020–30 September 2020) deployment periods. Panels (a) and (b) show results for chlorophyll *a* ( $\text{mg m}^{-3}$ ) in winter and summer, respectively. Particulate backscattering coefficient ( $b_{\text{bp}}$ ; 443 nm;  $\text{m}^{-1}$ ) seasonal spatial maps are shown in panels (c) and (d).



**3.2.4 Ocean biology.** There is mounting evidence that DMA production is influenced by ocean biological activity over the eastern North Atlantic<sup>18,33</sup> and other regions such as the northeast Pacific Ocean.<sup>32</sup> Periods of high ocean biological activity are linked to increased levels of organic nitrogen over the western North Pacific.<sup>98</sup> Table 1 shows the domain-wide mean values of chlorophyll *a* and particulate backscattering coefficient (ocean areas between 32–38°N, 69–77°W) based on the visual illustration in Fig. 3. There are higher chlorophyll *a* and particulate backscattering coefficient values near the coast. There were no major changes in either of the two parameters during the winter and summer 2020 campaigns, preventing the opportunity to compare periods of low and high biomass as in other studies.<sup>18</sup> A cautionary note again in light of this discussion is that there is an important distinction between biomass and biological activity as high standing stock does not necessarily translate into high emissions rates. However, we find value in still examining DMA trends as a function of ocean biomass levels.

While it is difficult to spatially link such values in these maps to actual cloud water samples owing to the range of limitations in making such a comparison, we focus on seasonal differences in ocean parameters and cloud water DMA. Both chlorophyll *a* and the particulate backscattering coefficient were larger in the winter, similar to DMA. Others have emphasized the limitation of these ocean biological parameters in terms of relationships with marine aerosol and cloud water organic enrichment,<sup>76,77</sup> but seasonal differences in Table 1 are at least supportive of a relationship that requires more detailed research. What remains difficult to disentangle is the relative importance of ocean biogenic emissions *versus* continental sources and other factors in governing cloud water DMA levels.

The study of Youn *et al.*,<sup>32</sup> which reported considerably higher cloud water DMA concentrations, was characterized by higher chlorophyll *a* values ( $\sim 2\text{--}6\text{ mg m}^{-3}$ ; see their Fig. 7) in the domain of the 72 hour back trajectories impacting their sampled air masses. Although impossible to prove here, those higher values are at least in support of a possible link between higher biogenic emissions and more cloud water DMA in that marine region compared to the northwest Atlantic. Comparing chlorophyll *a* values between studies is challenging as several factors impact how marine emissions affect clouds including the spatial domain over which these values are obtained.

### 3.3 Species interrelationships

Table S2† reports the interrelationships between various species of interest, broken down into the three temporal categories discussed already. The associated scatterplots are in Fig. S1.† We report Pearson's correlation coefficient (*r*) values only if they are statistically significant at a 95% confidence level ( $p < 0.05$ ). The absolute *r* value categorizes the strength of the relationship as follows:  $r = 0\text{--}0.2$ : weak;  $r = 0.21\text{--}0.4$ : moderately weak;  $r = 0.41\text{--}0.6$ : moderate;  $r = 0.61\text{--}0.8$ : moderately strong; and  $r \geq 0.81$ : strongly correlated.<sup>99</sup> A variety of species are included in Table S2† to represent different types of sources with the caveat that many of these species have multiple

sources:<sup>62,79,100–104</sup> sea spray ( $\text{Na}^+$ ), crustal matter (Al, Fe, nss- $\text{Ca}^{2+}$ ), marine biogenic emissions (MSA), combustion (As, Pb, Zn, Cu, V, Ni, nss- $\text{K}^+$ ), waste processing (Cd), and secondary aerosol formation processes ( $\text{NH}_4^+$ ,  $\text{NO}_3^-$ , nss- $\text{SO}_4^{2-}$ , oxalate). Relationships in Table S2† are insightful in providing hints about potential emissions sources and/or suggesting which acidic gases (*e.g.*, sulfuric, nitric, organic acids) participate in salt formation with DMA.

The winter temporal categories have one common feature regarding DMA and that is the significant correlation between DMA and  $\text{NH}_4^+$  ( $r = 0.97$ ). This is likely due to ammonia and amines being co-emitted from various sources such as from industry,<sup>35</sup> biomass burning,<sup>36</sup> animal husbandry,<sup>40</sup> and the ocean.<sup>41</sup> In addition,  $\text{NH}_4^+$  is a major base participating in salt formation with acids much like DMA does,<sup>18,105</sup> which has been supported by similar mass size distributions between DMA and  $\text{NH}_4^+$  in various regions.<sup>32,45</sup> Other similarities between the winter categories for DMA include a moderate correlation with oxalate ( $r = 0.58$ ), moderately strong and strong correlations with nss- $\text{Ca}^{2+}$ , nss- $\text{K}^+$ , and Al ( $r$ : 0.79–1.00).

Both nss- $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$  are major inorganic anions participating in salt formation with amines, as evident from similar mass size distributions with DMA.<sup>32,45,106</sup> It is somewhat surprising that DMA is only correlated to both acids in winter CAO conditions, with the correlation being slightly stronger for  $\text{NO}_3^-$  ( $r = 0.94$ ) than nss- $\text{SO}_4^{2-}$  ( $r = 0.79$ ). Oxalate is examined as there has been increased research investigating properties of alkylammonium carboxylate particles<sup>107</sup> pointing to appreciable hygroscopicity depending on the amine and carboxylic acid participating in salt formation together.<sup>22</sup> Oxalate was correlated to DMA in both winter seasons as already noted providing at least some support for a common formation mechanism and/or emissions source(s). Although not the only source, biomass burning is common in the wintertime along the U.S. east coast<sup>88</sup> and is an important source for DMA, oxalate, and several other species in Table S2† such as MSA, nss- $\text{K}^+$ , and Al.<sup>32,36,106</sup> We caution that some of the difference between winter categories can be due to the reduced sample sizes as is visually illustrated in Fig. S1† as a minority of points often can strongly influence a correlation coefficient. An important result is that the winter CAO conditions yield not only higher overall levels of cloud water DMA,  $\text{NH}_4^+$ ,  $\text{NO}_3^-$ , and nss- $\text{SO}_4^{2-}$ , but also more statistically significant correlations with species in Table S2† presumably due to more polluted air masses from the continent enriched with most of the various species studied here.

As will be shown in a case study flight, the summertime correlation results are driven mainly by one sample that was influenced by biomass burning. This is the most plausible explanation for why DMA is strongly correlated with nss- $\text{K}^+$ , oxalate, nss- $\text{SO}_4^{2-}$ , and  $\text{Na}^+$  ( $r$ : 0.88–0.97), as these species have been shown to be enhanced in concentration during periods of biomass burning.<sup>104,106,108</sup> Other strong correlations with DMA in summer include Cu, Zn, Cd, and Pb ( $r = 1.00$ ), which suggests these species were also enhanced in the single sample with biomass burning influence.

MSA is of relevance to this work as it stems partly from marine biogenic emissions<sup>109</sup> and because nucleation in the



marine atmosphere can occur between mixtures of water, amines, and MSA.<sup>110</sup> However, MSA was only significantly correlated to DMA in winter non-CAO conditions and strongly correlated with  $\text{NH}_4^+$  and  $\text{NO}_3^-$  in that same category ( $r = 0.96$  for both). This suggests it is not co-emitted with DMA to the extent that it is a strong predictor for DMA enhancement. Similar conclusions apply to the sea salt tracer species  $\text{Na}^+$ , which did not show a consistently strong correlation with DMA in all three temporal categories.

We compared the correlation results to those from the NiCE cloud water dataset<sup>32</sup> over the northeast Pacific (Table S3†). The NiCE results show that DMA was strongly correlated with  $\text{NH}_4^+$  ( $r = 0.83$ ), like ACTIVATE, but also with oxalate ( $r = 0.83$ ). In fact, oxalate was more correlated with DMA than the major inorganic anions  $\text{NO}_3^-$  ( $r = 0.63$ ) and  $\text{nss-SO}_4^{2-}$  ( $r = 0.38$ ). This reinforces the importance of continued research into alkylammonium carboxylate particles.<sup>22</sup> Like ACTIVATE, there was no strong correlation throughout the NiCE campaign between DMA and either  $\text{Na}^+$  or MSA, indicating that these species commonly derived from different ocean emission processes are not good predictors for the presence of DMA in clouds.

Of interest is the relative abundance on a molar basis of DMA versus  $\text{NH}_4^+$ , as the latter is the primary base in salt formation. Other work has suggested that there is competitive uptake between DMA and  $\text{NH}_3$  on acidic particles in the marine atmosphere dependent on the DMA :  $\text{NH}_3$  molar ratio.<sup>111</sup> One study showed that the ratio of particulate dimethylammonium :  $\text{NH}_4^+$  in liquid particles (compared to dry particles) exceeded that of DMA :  $\text{NH}_3$ , despite excess  $\text{NH}_3$  in the gas phase, demonstrating the important role of the aqueous phase;<sup>112</sup> they defined dimethylammonium as  $n\text{NH}_2(\text{CH}_3)_2^+ + n\text{NH}(\text{CH}_3)_2$ . Table 1 shows that the median DMA :  $\text{NH}_4^+$  molar ratio was higher in the two winter categories (0.002 = CAO, 0.003 = non CAO) than the summer (0.001). The DMA :  $\text{NH}_4^+$  molar ratio is considerably higher in NiCE (0.018;  $n = 77$ ) than ACTIVATE (0.002;  $n = 37$ ) in the absence of biomass burning. When compared to those values, there was very little difference in the respective campaign-specific ratios in the presence of biomass burning for NiCE (0.019;  $n = 10$ ) and ACTIVATE (0.001;  $n = 1$ ). Note that this excluded classification of ACTIVATE data into the biomass burning category if there was residential wood burning, such as in the winter,<sup>88</sup> which is harder to identify as compared to wildfire plumes. Molar ratios reported from aerosol measurements in southern Arizona and the NiCE region by the central California coast ranged from 0 to 0.04,<sup>32</sup> whereas values in the fine mode ranged from 0.005–0.2 for rural and continental air by Toronto.<sup>45</sup>

Fig. 4 shows the relationship between  $\text{NH}_4^+$  and DMA. The positive correlation is slightly higher for NiCE ( $R^2 = 0.69$ ) relative to ACTIVATE ( $R^2 = 0.54$ ). The separation between the data points per campaign points to some unique differences between the two regions. It is unclear if the separating factors are more related to pollution sources, transport behavior and photochemical aging differences between  $\text{NH}_3/\text{NH}_4^+$  and DMA, or meteorology, warranting analysis of data from other regions.

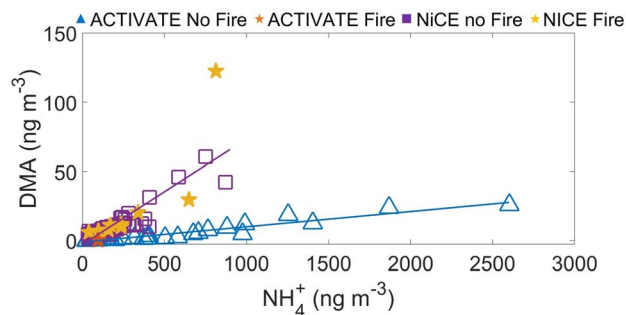


Fig. 4 Relationship between DMA and  $\text{NH}_4^+$  mass for ACTIVATE ( $R^2 = 0.54$ ; blue) and NiCE ( $R^2 = 0.69$ ; purple). Star markers represent those samples in each campaign that were influenced by biomass burning from summertime wildfires. Equations of the best fit lines for each campaign were as follows: NiCE =  $0.08x - 2.89$ ; ACTIVATE =  $0.01x - 1.02$ .

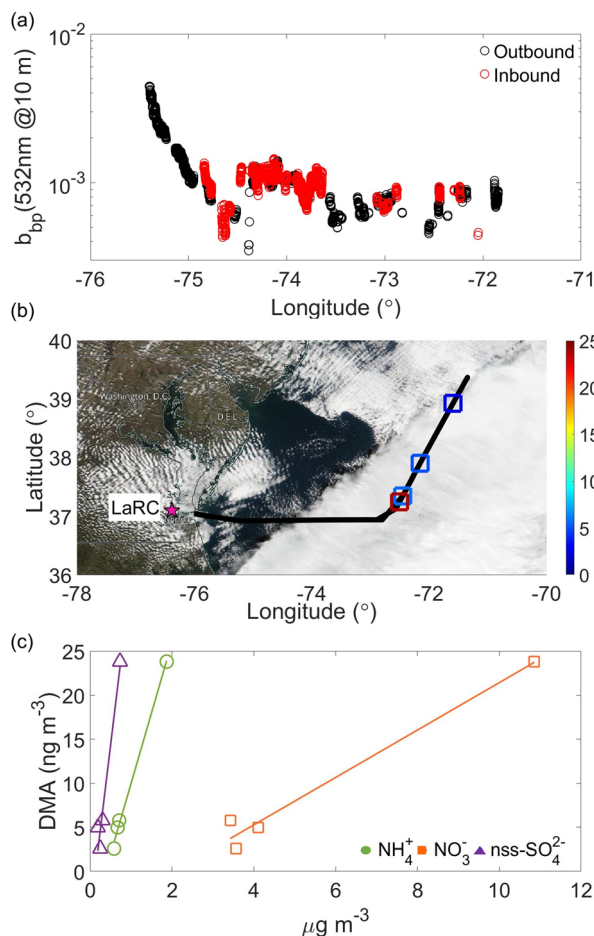
### 3.4 Case studies

**3.4.1 Winter flights.** Two case flights are summarized here that were selected based on having numerous cloud water samples. The first flight (Fig. 5a–c) on 14 February 2020 was in CAO conditions, unlike the second flight (Fig. 6a–c) on 6 March 2020. The flight on 14 February was in post-frontal conditions with high cloud fraction during most of the flight, with sampling in warm and mixed-phase clouds. The wind direction was northerly/northwesterly along most of the flight track, consistent with the organization of the cloud street features characteristic of CAO conditions over the northwest Atlantic. DMA concentrations in the four collected cloud water samples ranged from 5.63 to 24.78  $\text{ng m}^{-3}$ , which is on the high end of the concentrations during ACTIVATE flights (Table 1). These samples were collected at altitudes ranging between 837 and 1400 m.

DMA exhibited strong correlations with  $\text{NH}_4^+$ ,  $\text{nss-SO}_4^{2-}$ ,  $\text{NO}_3^-$ , and  $\text{nss-K}^+$  ( $R^2 \geq 0.89$ ). The strong relationship with  $\text{nss-K}^+$  in the wintertime could be due to residential wood burning emissions.<sup>88</sup> The spatial distribution of concentrations shows one sample with especially high concentrations near the turn point of the aircraft. The results of this flight support continental emissions being an important driver of the high concentrations owing to the origins of these species from anthropogenic precursors. We speculate that low temperatures characteristic of CAOs may help facilitate the formation of ammonium- and aminium-based salts that can activate into drops, in addition to dissolution of gaseous amines into cloud water.<sup>28</sup> It is hard to tease out the relative importance of both mechanisms with the current dataset.

As the ocean biological proxies from MODIS-Aqua have coarse temporal resolution, here we provide context during the flight by coloring the flight track with the HSRL-2 retrieval of the particulate backscattering coefficient. Further motivation for using the airborne HSRL-2 data is that ocean satellite data is obscured by clouds, including cirrus clouds. Linking the lidar data for upper ocean characteristics to cloud data is complex and needs to consider the actual amount of ocean biogenic

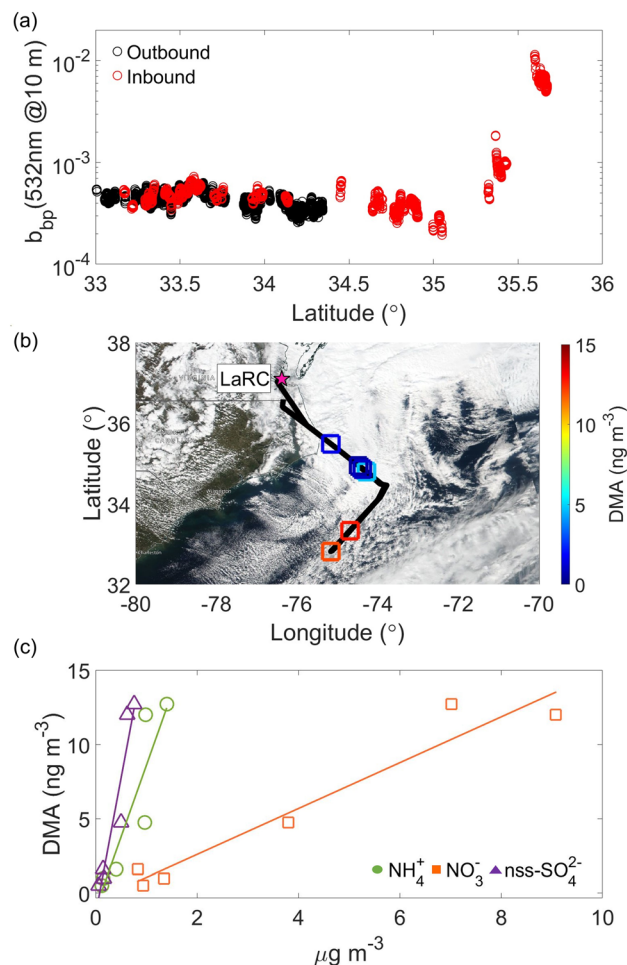




**Fig. 5** Summary of data from ACTIVATE research flight 1 on Friday 14 February 2020. (a) HSRL-2 particulate backscattering ( $b_{bp}$ ; 532 nm) coefficient near the ocean surface (depth of 10 m) as a function of longitude during the round flight. Red and black markers show data for the inbound and outbound tracks, respectively. (b) Spatial map of DMA concentrations overlaid on true-color imagery from the Visible Infrared Imaging Radiometer Suite (VIIRS; NASA Worldview). (c) Scatter plots of DMA versus inorganic species with linear best fits shown ( $R^2$ : DMA- $\text{NH}_4^+$  = 0.99, DMA- $\text{NO}_3^-$  = 0.98, DMA- $\text{nss-SO}_4^{2-}$  = 0.96).

emissions, transport path, and the possibility of wet scavenging of particles during transport offshore. However, this is a unique application of HSRL-2 data that we explore here and recommend leveraging in future studies of this nature. Fig. 5a reveals a longitudinal gradient with higher particulate backscattering coefficient values closer to the coast, suggesting that likely ocean emissions were not a strong driver of DMA in this flight based on correlations DMA exhibited with other species linked to anthropogenic activity.

The case flight on 6 March 2020 is summarized in Fig. 6 and was conducted on a day without CAO conditions. Winds were predominantly westerly with continental air mass influence. Six cloud water samples were collected at aircraft altitudes ranging between 618 and 2060 m. DMA concentrations ranged from 0.52 to 12.72  $\text{ng m}^{-3}$ , with strong correlations when compared to  $\text{NH}_4^+$ ,  $\text{NO}_3^-$ , and  $\text{nss-SO}_4^{2-}$  ( $R^2 \geq 0.82$ ), similar to the 14 February flight. This flight had a weaker correlation between



**Fig. 6** Summary of data from ACTIVATE research flight 16 on Friday 6 March 2020. (a) HSRL-2 particulate backscattering ( $b_{bp}$ ; 532 nm) coefficient near the ocean surface (depth of 10 m) as a function of latitude during the flight. Red and black markers show data for the inbound and outbound flights, respectively. (b) Spatial map of DMA concentrations overlaid on true-color imagery from the Visible Infrared Imaging Radiometer Suite (VIIRS; NASA Worldview). (c) Scatter plots of DMA versus inorganic species with linear best fits shown ( $R^2$ : DMA- $\text{NH}_4^+$  = 0.82, DMA- $\text{NO}_3^-$  = 0.94, DMA- $\text{nss-SO}_4^{2-}$  = 0.91).

DMA and  $\text{nss-K}^+$  ( $R^2 = 0.40$ ). The HSRL-2  $b_{bp}$  data show expectedly higher concentrations near the coast (Fig. 6a). It is noteworthy to highlight the HSRL-2  $b_{bp}$  coefficient data reproducibility for both flights (Fig. 5a and 6a) for the inbound (red markers) and outbound (black markers) flights.

**3.4.2 Summertime flight with fire influence.** Research flight 28 on 26 August 2020 was unique in that the Falcon sampled a cloud with biomass burning influence supported by various datasets. Summertime is typically marked by lower cloud fraction over the northwest Atlantic compared to winter-time, which is evident in Fig. 7. Winds were generally westerly on this flight. While the HSRL-2 often detected smoke during ACTIVATE flights, it was not always the case that there was clear evidence of it mixing with boundary layer clouds sampled by the Falcon. Research flight 28 was separately investigated by Mardi *et al.*<sup>89</sup> who showed the mixing of smoke plumes with boundary



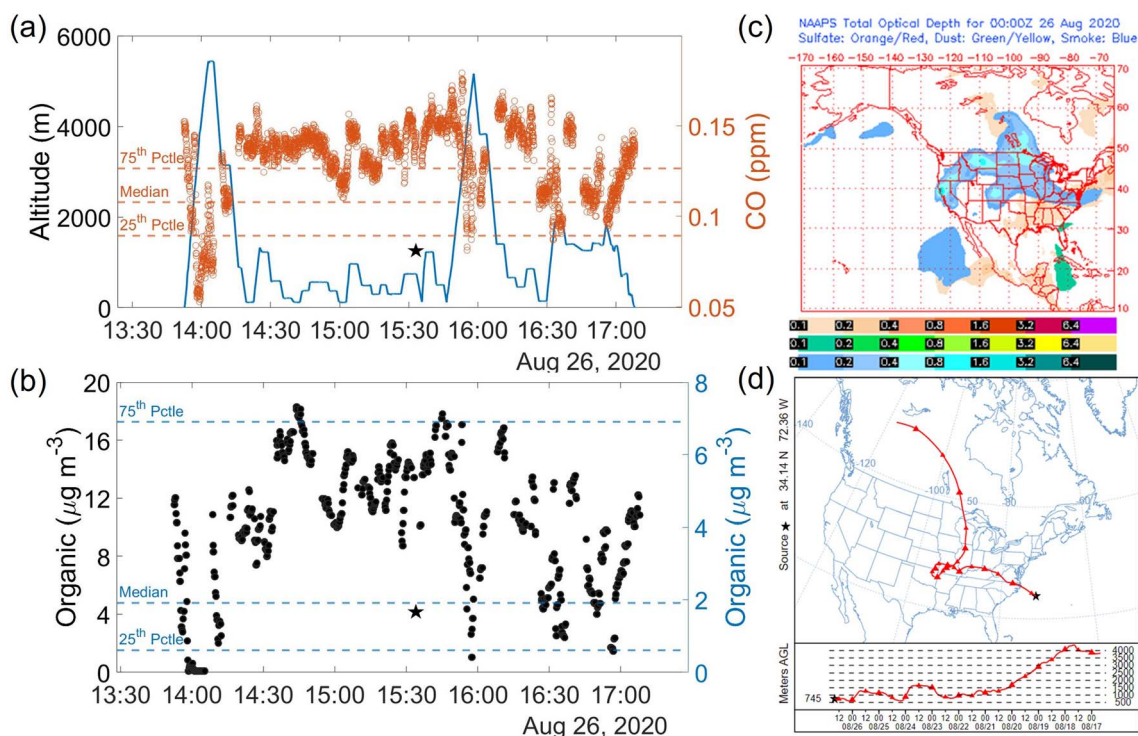


Fig. 7 (a) Time series for *in situ* measurements of CO and flight altitude over the ACTIVATE region during RF28 (26 August 2020). (b) Time series for *in situ* AMS organic concentration in  $\mu\text{g m}^{-3}$ . Dashed lines show the median, 25th percentile (pctle), and 75th pctle for the all the summer data. (c) Prognostic NAAPS map of smoke aerosol distribution. (d) HYSPLIT 240 hour back trajectories corresponding to the cloud water sample collected during RF28. The black star in (a) and (b) shows the collection time of the cloud water sample.

layer clouds. That study showed enhanced aerosol extinction coefficient (532 nm) in the marine boundary layer and even higher values in the free troposphere (>2 km). Furthermore, flight scientist reports pointed to biomass burning influence, including support from *in situ* measurements of CO and AMS organic concentrations.<sup>113,114</sup> Organic and CO concentrations along the flight track are clearly above the respective summer median values, including the time during the cloud water collection (Fig. 7a and b). NAAPS data for smoke and sulfate (Fig. 7c) and HYSPLIT 72 hour back trajectory data strongly suggest that the sampled smoke was influenced in part by air transported from western North America where there were active wildfires (Fig. 7d).

The concentration of DMA in this single sample was  $0.38 \text{ ng m}^{-3}$ , slightly above the median in the summer of  $0.25 \text{ ng m}^{-3}$ . Regarding other chemical measurements in cloud water, this sample's oxalate to sulfate ratio was the highest of the 2020 flights (0.24). Recent work has attributed such high ratios to biomass burning influence.<sup>115</sup> Oxalate's concentration in this single sample was  $130.76 \text{ ng m}^{-3}$  in contrast to the summer median value of  $92.77 \text{ ng m}^{-3}$ , indicating that it was higher than normal.

In contrast, we also examined the other two flights in the summer with detectable cloud water DMA levels (13 and 17 August), and none of them showed any evidence of smoke. A previous study found that amines (C1–C6) are detected during biomass burning plume events, suggesting that biomass

burning can be a considerable source of amines in the southeastern U.S.<sup>36</sup> As noted already, cloud water DMA levels were significantly higher during biomass burning conditions over the northeast Pacific.<sup>32</sup>

## 4. Conclusions

A unique airborne cloud water dataset is analyzed in this work with a focus on dimethylamine (DMA), which is a common alkyl amine observed in atmospheric gases, particles, and cloud water. This study adds important DMA data to a very limited archive of such data in cloud water based on airborne measurements, with the only other such studies to our knowledge being that of the NiCE dataset from the northeast Pacific and CAMP<sup>2</sup>Ex from southeast Asia.<sup>48,49</sup>

Notable results in this work are that concentrations of this species are markedly lower over the northwest Atlantic in ACTIVATE compared to the northeast Pacific in NiCE. One possible explanation is more ocean biological emissions in the latter region. When comparing ACTIVATE winter and summer flights, there were more relative samples with detectable DMA in the winter (67% of winter non-CAO samples; 44% of winter CAO samples) as compared to the summer (14%). The concentration trend of  $\text{NO}_3^-$ ,  $\text{NH}_4^+$ , and  $\text{nss-SO}_4^{2-}$  were uniquely similar to DMA among species examined in that their levels were highest in winter CAO conditions, followed by winter non-CAO conditions, and lastly summer. Analysis of other



seasonally-dependent factors suggests that winter CAO levels are highest due to a presumed combination of low temperatures and offshore flow with anthropogenic pollutants, including biomass burning. This work adds to the growing evidence of how biomass burning is an important source of alkyl amines, such as DMA. This dataset motivates continued research into the partitioning behavior of amines between the gas, solid, and aqueous phases, in addition to examining the effect of cloud water acidity on partitioning. The winter samples in ACTIVATE were more acidic than summer, and it is uncertain how influential this was in promoting higher DMA levels.

As amines have been linked to new particle formation<sup>116–118</sup> it is important to note that winter CAO conditions (lower temperature and humidity) were most conducive to such events during ACTIVATE,<sup>53</sup> especially near clouds, with low temperatures and RH shown to be potentially contributing factors. Amines associated with nucleated particles can grow into sizes sufficiently large to activate into drops and thus be sampled as cloud water. Although outside the scope of this study, future research is encouraged to assess if these nucleation events identified around clouds in the northwest Atlantic are contributing appreciably to DMA solute concentrations in cloud water, which would have implications for other species participating in those nucleation events. Studying amines is important as they play an important role in the marine environment's carbon and nitrogen cycles.

## Data availability

ACTIVATE airborne data can be accessed from <http://doi.org/10.5067/ASDC/>

[ACTIVATE\\_Aerosol\\_AircraftInSitu\\_Falcon\\_Data\\_1](http://doi.org/10.5067/ASDC/ACTIVATE_AerosolCloud_AircraftRemoteSensing_KingAir_Data_1) and [http://doi.org/10.5067/ASDC/ACTIVATE\\_AerosolCloud\\_AircraftRemoteSensing\\_KingAir\\_Data\\_1](http://doi.org/10.5067/ASDC/ACTIVATE_AerosolCloud_AircraftRemoteSensing_KingAir_Data_1). ACTIVATE data can be accessed by registering for an Earthdata account/login: <https://urs.earthdata.nasa.gov/>. MONARC data can be accessed at: <https://doi.org/10.6084/m9.figshare.5099983.v10>.

## Author contributions

AFC and AS conceived the study and drafted the manuscript. YC, BLC, EC, HD, JPD, GSD, MF, SK, RHM, JBN, MAS, CTS, TS, KLT, CV and LDZ contributed to data collection and archival from ACTIVATE. All authors contributed to editing the final version of the manuscript.

## Conflicts of interest

The authors declare no conflict of interest.

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